The role of aneuploidy in the evolution of cancer drug resistance

Remus Stana¹, Uri Ben-David², Daniel B. Weissman³, and Yoav Ram^{1,*}

¹School of Zoology, Faculty of Life Sciences, Tel Aviv University, Tel Aviv, Israel
²Department of Human Molecular Genetics and Biochemistry, Faculty of Medicine, Tel Aviv University, Tel Aviv, Israel

³Department of Physics, Emory University, Atlanta, GA *Corresponding author: yoav@yoavram.com

February 12, 2024

Abstract

Evolutionary rescue is the process by which a population survives a sudden environmental change that initially causes the population to decline towards extinction. A prime example of evolutionary rescue is the ability of cancer to survive exposure to various treatments. One evolutionary mechanism by which a population of cancer cells can to adapt to chemotherapy is aneuploidy. Aneuploid cancer cells can have higher fitness in an environment altered by anticancer drugs, e.g., because of incomplete pathways targeted by the drugs. Indeed, aneuploidy is highly prevalent in tumors, and moreover, some anti-cancer drugs fight cancer by increasing chromosomal instability. Here, we examine how aneuploidy impacts the fate of a population of cancer cells. We use multi-type branching processes to approximate the probability that a tumor survives drug treatment as a function of the initial tumor size, and rates at which aneuploidy and other beneficial mutations occur, and the growth rates of the sensitive and resistant cells. Additionally, we investigate what effect having a fraction of tumor cells being an euploid before the onset of therapy have on the probability of evolutionary rescue. We also approximate the mean recurrence time for the tumor to revert back to its initial size. We find that aneuploidy can play an important role in the relapse of secondary tumors who have not yet been detected and thus are smaller in size.

Keywords: whole-chromosome duplication, evolutionary model, adaptive evolution, cancer, drug resistance, chromosome instability

Introduction

Aneuploidy in cancer. Each year approximately 10 million people die from cancer (Kocarnik et al., 2022), so understanding the factors that contribute to failure of interventions is of great importance. One hypothesized factor is aneuploidy, where cells are characterized by an imbalanced karyotype and chromosome copy number alterations (Schukken and Foijer, 2018), caused by chromosomal instability (CIN), the mitotic process in which cells suffer from chromosome mis-segregation that leads to aneuploidy. Importantly, aberrations in chromosome copy number have been shown to allow cancer cells to survive under stressful conditions such as drug therapy (Lukow et al., 2021, Rutledge et al., 2016). Indeed, cancer cells are often likely to be aneuploid, and aneuploidy is associated with poor patient outcomes (Ben-David and Amon, 2020, Smith and Sheltzer, 2018).

Ippolito et al. (2021) induced aneuploidy in cancer cell lines by exposing them to reversine, a small-molecule inhibitor of the mitotic kinase Mpsi1, and then to chemotherapeutic agents such as vemurafenib. Reversine-treated cells had higher proliferation rate in the environment altered by anti-cancer drugs compared to wildtype cancer cells. Similarly, Lukow et al. (2021) induced aneuploidy in cancer cells and observed that such cells have an advantage compared to wildtype cells during chemotherapy, despite having lower fitness before the onset of chemotherapy. One proposed mechanism through which aneuploidy is able to confer resistance to chemotherapeutics is by antagonizing cell division, which prevents the drugs from damaging DNA and microtubules (Replogle et al., 2020).

An important aspect of aneuploidy is the rate with which cells become aneuploid, which is several orders of magnitude higher than the beneficial mutation rate (Bakker et al., 2023). Consequently, a cell exposed to a stress such as chemotherapeutic drugs can acquire aneuploidy faster when compared to acquiring a mutation, especially when several proposed anti-cancer drugs elevate the rate of missegregation in order to fight cancer (Lee et al., 2016).

Evolutionary rescue. Populations adapted to a certain environment are vulnerable to environmental changes, which might cause extinction of the population. Examples of such environmental changes include climate change, invasive species or the onset of drug therapies. Adaptation is a race against time as the population size decreases in the new environment (Tanaka and Wahl, 2022). *Evolutionary rescue* is the process where the population acquires a trait that increases fitness in the new environment such that extinction is averted. It is mathematically equivalent to the problem of crossing of fitness valley (Weissman et al., 2009, 2010). There are three potential ways for a population to survive environmental change: migration to a new habitat similar to the one before the onset of environmental change (Cobbold and Stana, 2020, Harsch et al., 2014, Zhou, 2022); adaptation by phenotypic plasticity without genetic modification (Carja and Plotkin, 2017, 2019, Gunnarsson et al., 2020, Levien et al., 2021); and adaptation through genetic modifications, e.g., mutation (Gomulkiewicz and Holt, 1995, Orr and Unckless, 2014, Uecker and Hermisson, 2011, 2016, Uecker et al., 2014).

Gunnarsson et al. (2020) analyze a model where a tumor consisting of two populations of cancer cells, one drug resistant and the other drug sensitive, is able to evade extinction by cells switching between the two phenotypes through epigenetic mutations. They found that even when the drug resistant type is barely viable the epimutations have the effect of guaranteeing evolutionary rescue. Evolutionary rescue in one step in which an initially declining population, after a sudden environmental change, has to acquire a mutation has been studied in the context of population genetics by (Orr and Unckless, 2008, 2014). They analyzed a model where the mutant strain is present in small number at the onset of therapy and concluded that this can significantly enhance the chance that the population will survive.

Most models focus on the probability that at least one mutation rescues the population. How multiple mutations contribute to the survival of the population is less explored, but Wilson et al. (2017) have shown that evolutionary rescue is significantly enhanced by soft selective sweeps when

multiple mutations contribute. Evolutionary rescue that requires two successive mutations has been investigated using diffusion approximation by Martin et al. (2013).

Here we build on previous work on evolutionary rescue after a sudden environment change caused by the initiation of chemotherapy. We wish to understand what effect does aneuploidy have on the probability of evolutionary rescue when it acts as an intermediary between the wildtype and the mutant cancer cells. We also calculate the mean time that an initially declining tumor cell population reaches its pre-treatment size. Given that aneuploidy is present in many tumors even before the onset of therapy (Lukow et al., 2021) we also take into consideration the effect that standing genetic variation has on the tumor dynamics.

Methods

Evolutionary model. We follow the number of cancer cells that have one of three different genotypes at time t: wildtype, w_t ; aneuploid, a_t ; and mutant, m_t . These cells divide and die with rates λ_k and μ_k (for k = w, a, m). The difference between the division and death rate is $\Delta_k = \lambda_k - \mu_k$. We assume the population of cells is under a strong stress, such as drug therapy, to which the wildtype genotype is susceptible or sensitive and therefore $\Delta_w < 0$, whereas the mutant is resistant to the stress, $\Delta_m > 0$. We analyze three scenarios: in the first, an euploid cells are partially resistant, $\Delta_m > \Delta_a > 0$; in the second, an euploid cells are tolerant, $0 > \Delta_a > \Delta_w$ (see Brauner et al., 2016, for the distinction between susceptible, resistant, and tolerant); in the third, aneuploid cells are non-growing, stationary or "barely growing", that is, either slightly tolerant or slightly resistant, such that $\Delta_a \approx 0$. We assume that both chromosomal missegregation and mutation occur during the process of mitosis. Wildtype cells may divide and then missegregate to become aneuploids at rate $u\lambda_w$. Both aneuploid and wildtype cells may divide and mutate to become mutants at rates $v\lambda_a$ and $v\lambda_w$, respectively. To model standing genetic variation, we assume that before the onset of therapy, wildtype cells become aneuploid with rate $\tilde{u}\lambda_w$ (which may differ from $u\lambda_w$) and that an euploidy confers a fitness cost s in the drug-free environment (we assume that an euploidy significantly increases the death rate of wildtype cancer cells in an environment which is not affected by chemotherapeutic drugs). See Figure 1 for a schematic representation of the model and Figure 2 for sample trajectories of the different genotypes.

Stochastic simulations. Simulations are performed using the *Gillespie stochastic simulation algo*rithm (Gillespie, 1976, 1977) implemented in Python (Van Rossum and Others, 2007). The simulation monitors the number of cells of each type: wildtype, an euploid, and mutant. The wildtype population initially consists of $w_0 = N$ cells, whereas the other cell types are initially absent.

The state of the stochastic system at time t is represented by the triplet (w_t, a_t, m_t) . The following describes the events that may occur (right column), the rates at which they occur (middle column), and the effect these events have on the state (left column, see Figure 1):

```
(+1,0,0): \lambda_w w_t (1-u-v) (birth of wildtype cell),
(-1,0,0): \mu_w w_t (death of wildtype cell),
(0,+1,0):
             u\lambda_w w_t (wildtype cell divides and becomes an euploid),
(0,0,+1):
                        (wildtype cell divides and becomes mutant),
(0, +1, 0):
             \lambda_a a_t (1 - v) (birth of an euploid cell),
(0,-1,0):
                      (death of an euploid cell),
              \mu_a a_t
                       (aneuploid cell divides and becomes mutant),
(0,0,+1):
             v\lambda_a a_t
              \lambda_m m_t
(0,0,+1):
                       (birth of mutant cell),
(0,0,-1):
                       (death of mutant cell).
              \mu_m m_t
```

For the remaining of this paper we assume that the division rates for wildtype and aneuploid cells can be written as $\lambda_w w_t$ $(1-u-v) \approx \lambda_w w_t$ and $\lambda_a a_t$ $(1-v) \approx \lambda_a a_t$ because $u,v \ll 1$ (see Table 1). Each iteration of the simulation loop starts with computing the rates v_k of each event k. We then draw the time until the next event, Δt , from an exponential distribution whose rate parameter is the sum of the rates of all events, such that $\Delta t \sim Exp(\sum_j v_j)$. Then, we randomly determine which event occurred, where the probability for event k is $p_k = v_k/\sum_j v_j$. Finally, we update the number of cells of each type according to the event that occurred and update the time from t to $t + \Delta t$. We repeat these iterations until either the population becomes extinct (the number of cells of all types is zero) or the number of mutant cells is high enough so that its extinction probability is < 0.1%, that is until

$$m_t > \left| \frac{3 \log 10}{\log \left(\lambda_m / \mu_m \right)} \right| + 1,$$

which we obtain by solving $1 - (1 - p_m)^{m_t} = 0.999$ for m_t with $p_m = \Delta_m / \lambda_m$ as the probability that a single mutant escapes stochastic extinction (see Appendix A).

When simulations are slow (e.g., due to large population size), we use τ -leaping (Gillespie, 2001), where we assume that the change in the number of cells of genotype k in a fixed time interval Δt is Poisson distributed with mean $v_k \Delta t$. If the change in the number of cells is negative and larger than the subpopulation size then the subpopulation size is updated to be zero.

Parameterization. To parametrize the simulations, we assume that the cells under consideration are melanoma cells and rely on Rew and Wilson (2000) and Bozic et al. (2013) for the division and death rates, respectively. Rew and Wilson (2000) report *in vivo* measurements of the potential doubling times (the waiting time for the number of cells in the tumor to double disregarding cell death) for a large set of cancer types. The division rate is obtained as $\lambda = \log 2/T \approx 0.1$ per day. We select this to be the division rate for wildtype and mutant cells.

Bozic et al. (2013) report the growth rate Δ_w for wildtype melanoma cancer cells from which they deduce the death rate $0.11 \le \mu_w \le 0.17$. We use $\mu_w = 0.14$ per day. Additionally, they observe the growth rate of cancer cells prior to treatment to be 0.01, which we use as the growth rate of mutant cells, which are resistant to the drug. Thus, we use $\mu_m = 0.1 - 0.01 = 0.09$ per day as the death rate for mutant cells.

Aneuploid death rate μ_a is set to the same value as the mutant death rates, $\mu_m = 0.09$ per day, given that aneuploidy increases resistance to the drug, such as cisplatin, by antagonizing cell division (Replogle et al., 2020). Aneuploid division rate is selected such that the aneuploid growth rate $\Delta_w \ll \Delta_a \ll \Delta_m$ which means that $0.06 \le \lambda_a \le 0.1$. For most of our simulations we use $\lambda_a = 0.0899$ per day, such that aneuploidy can only act as a *stepping stone* for the generation of the mutant that rescues the cancer cell population.

We assume the mutation rate is 10^{-7} per gene per cell division (Loeb, 2001) and since we assume that a single target gene confers resistance to the drug, we use $v = 10^{-7}$ per cell division. Bakker et al. (2023) determined that the missegregation rate must be between $10^{-3} - 10^{-2}$ per chromosome per cell division. Given that a duplication is beneficial for cancer resistance only in one or two chromosomes (see details next paragraph) we use $u = 10^{-2}$ per cell division. For the missegregation rate in the drug-free environment, \tilde{u} , we use the lower-end value of $\tilde{u} = 10^{-3}$ per cell division, as some drugs increase the rate of aneuploidy (Mason et al., 2017, Wang et al., 2019).

The fitness cost s of an euploidy before the onset of therapy is difficult to estimate as we are interested in a specific type of an euploidy which improves the fitness of cancer cells in an environment altered by drugs. We derive s by using the formula $s = \tilde{u}\lambda_w/f$, where f is the fraction of an euploid cancer cells. To estimate f, we note that Lukow et al. (2021) mixed together wildtype and an euploid A375 melanoma cells at 50: 50 ratio, cultured them in drug-free environment and observed the ratio evolve as a function of time with the an euploid cells declining to 15% after 24 days. Ippolito et al.

(2021) observed that trisomy in Chr II and VI are most likely to confer increased resistance against the chemotherapeutic agent vemurafenib for A375 cells. We assume that if a tumor is aneuploid then it most likely has trisomy (Gisselsson et al., 2010) in the pre-treatment environment and that cells with more than one trisomy are very unlikely to survive. Additionally, we assume that all trisomies are equally likely and, as a result, we obtain the fraction of cancer cells with favorable aneuploidy as $f = 2/22 \times 0.15 \approx 0.0136$ (i.e. 1.36% of pre-treatment cancer cells have the beneficial aneuploidy). This gives us a fitness cost of $s = 7.3 \times 10^{-3}$ per day. We note that when we refer to wildtype cancer cells we include those cells that have any aneuploidy except trisomy in Chr II and VI as those are the aneuploid cells which are hypothesized to have higher fitness in the environment altered by drugs such as vemurafenib.

Density-dependent growth. In our analysis we assume that lineages produced by cells from the initial population divide and die independently of each other, which may be unrealistic, as cells usually compete for resources. This is a good approximation as we assume that the drugs cause the wildtype population to decline rapidly and, as a result, density-dependent effects can be neglected. A more realistic model includes competition for limited resources and spatial structure, which may play an important role in the development of cancer (e.g., Martens et al., 2011). To simulate division and death rates that depend on the number of cells in the population, we assume the rates of division and death to be

$$\begin{split} \lambda_w' &= \lambda_w, \\ \mu_w' &= \mu_w, \\ \lambda_a' &= \lambda_a, \\ \mu_a' &= \mu_a + \lambda_a \frac{w + a + m}{K}, \\ \lambda_m' &= \lambda_m, \\ \mu_m' &= \mu_m + \lambda_m \frac{w + a + m}{K}, \end{split}$$

where K is the tumor carrying capacity. The effective carrying capacity of this model is $K_e = K\Delta_a/\lambda_a \approx 10^6$ for $K = 10^8$, $\lambda_a = 0.0901$, $\mu_a = 0.09$, where we define the effective carrying capacity to be the population size at which the aneuploid division rate is equal to the aneuploid death rate.

Code and data availability. All source code is available online at https://github.com/yoavram-lab/EvolutionaryRescue.

Results

Evolutionary rescue probability

In our model, evolutionary rescue occurs when resistant cells appear and establish (avoid random extinction) in the population $(m_t \gg 1)$ before the population becomes extinct $(w_t = a_t = m_t = 0)$. Aneuploidy may contribute to evolutionary rescue by either preventing (when $\Delta_a > 0$) or delaying (when $0 > \Delta_a > \Delta_w$) the extinction of the population before mutant cells appear and establish. We assume independence between clonal lineages starting from an initial population of N wildtype cells (we check the effect of density-dependent growth on our results below). We therefore define p_w as the probability that a lineage starting from a single wildtype cell avoids extinction by acquiring drug resistance. Thus, $N^* = 1/p_w$ is the threshold tumor size above which evolutionary rescue is very likely, and the rescue probability is given by

$$p_{\text{rescue}} = 1 - (1 - p_w)^N \approx 1 - e^{-Np_w} = 1 - e^{-N/N^*},$$
 (2)

where the approximation $(1-p_w) \approx e^{-p_w}$ assumes that p_w (but not necessarily Np_w) is small. Indeed, when $N < 1/p_w$, then the probability for evolutionary rescue is $p_{\text{rescue}} \approx Np_w$ and when $N > 1/p_w$, it is $p_{\text{rescue}} \approx 1$, justifying the definition of N^* as the threshold tumor size for evolutionary rescue.

We use the theory of multi-type branching processes to find approximate expressions eqs. (A4), (A7) and (A11) for p_w in different regimes (see appendix A). Substituting these into $N^* = 1/p_w$, we find approximations for the threshold tumor size, N^* . For these approximations, an important quantity is $T^* = (4v\lambda_a^2\Delta_m/\lambda_m)^{-1/2}$, which is the critical time an aneuploid lineage needs to survive to produce a resistant mutant that avoids random extinction. First, if aneuploidy is very rare $(u\lambda_a T^* < 1)$, or if aneuploidy is rare $(u\lambda_a < -\Delta_a)$ and very sensitive to the drug $(\Delta_a T^* < -1)$, then evolutionary rescue will likely occur by a direct resistance mutation in a sensitive wildtype cell without the involvement of aneuploidy, such that

$$N_m^* \approx \frac{|\Delta_w|}{v\lambda_w} \cdot \frac{\lambda_m}{\Delta_m}.$$
 (3)

Here, $|\Delta_w|/(v\lambda_w)$ is the ratio of the rate at which wildtype cells are decreasing in number and the rate at which they are mutating. Notably, the aneuploidy parameters (u, λ_a, μ_a) do not affect N_m^* .

Otherwise, aneuploidy is frequent enough $(u\lambda_a > \max(-\Delta_a, 1/T^*))$ to affect the evolution of drug resistance. The threshold tumor size can then be approximated by one of the following cases, depending on $\Delta_a T^*$, the change in the aneuploid log-population size during the critical time,

$$N_a^* \approx \frac{|\Delta_w|}{u\lambda_w} \cdot \begin{cases} \frac{|\Delta_a|}{v\lambda_a} \cdot \frac{\lambda_m}{\Delta_m}, & \Delta_a T^* \ll -1 \text{ (tolerant aneuploids),} \\ 2\lambda_a T^*, & -1 \ll \Delta_a T^* \ll 1 \text{ (stationary aneuploids),} \\ \frac{\lambda_a}{\Delta_a}, & \Delta_a T^* \gg 1 \text{ (resistant aneuploids).} \end{cases}$$
(4)

These approximations perform very well when compared to results of stochastic evolutionary simulations (Figures 3 and 4). The first line describes the case in which aneuploid cells are still effectively killed by the treatment, but not as quickly as the wild type. In the second case, aneuploid cells are sufficiently resistant that the expected size of each aneuploid lineage is roughly 1. In both of these cases, aneuploidy increases the probability of rescue by slowing or halting the decrease of the cancer population, allowing more opportunities for producing resistant mutants. In the third case, aneuploid cells are sufficiently resistance for the cancer population to re-grow the tumor even without additional resistance mutations. Notably, in this case the mutant parameters $(v, \lambda_m, \text{ and } \Delta_m)$ do not affect N_a^* beyond their effect on T^* . In all cases, N_a^* is proportional to 1/u such that increasing the missegregation rate u will decrease the threshold tumor size (Figure 4B). Furthermore, increasing the aneuploid growth rate Δ_a (which appears both in the terms and in the conditions), also reduces the threshold tumor size, with a sharp decrease around $\Delta_a = 0$, but the effect is minor when $|\Delta_a|$ is small compared to T^* as this would result in the second case where $dN_a^*/d\Delta_a = 0$ (Figure 4A).

Using eqs. (3) and (4), we can find the ratio of threshold tumor size for rescue via an euploidy (u is high) or via direct mutation (u is low),

$$\frac{N_a^*}{N_m^*} \approx \begin{cases} \frac{|\Delta_a|}{u\lambda_a}, & \Delta_a T^* \ll -1, \\ \frac{1}{u} \left(v \cdot \frac{\Delta_m}{\lambda_m} \right)^{1/2}, & -1 \ll \Delta_a T^* \ll 1, \\ v \frac{\Delta_m}{\lambda_m} \cdot \left(u \frac{\Delta_a}{\lambda_a} \right)^{-1}, & \Delta_a T^* \gg 1. \end{cases}$$
(5)

As expected, this ratio increases with the mutation rate v and decreases with the aneuploidy rate u. In the first case, $|\Delta_a|/(u\lambda_a)$ is the ratio of the expected time for an aneuploid lineage to appear, $1/(u\lambda_a)$, and the expected time until that lineage disappears, $1/|\Delta_a|$. In the third case, $\left(v\frac{\Delta_m}{\lambda_m}\right)/\left(u\frac{\Delta_a}{\lambda_a}\right)$ is the ratio of the rates of formation of resistant mutants that avoid extinction and partially resistant

aneuploids that avoid extinction. In the second case, $\frac{1}{u} \left(v \cdot \frac{\Delta_m}{\lambda_m} \right)^{1/2} = \sqrt{\frac{\Delta_a}{u\lambda_a} \cdot v \frac{\Delta_m}{\lambda_m} \cdot \left(u \frac{\Delta_a}{\lambda_a} \right)^{-1}}$, which is the geometric mean of the first and third cases.

Interestingly, increasing both the aneuploid division rate, λ_a , and the aneuploid death rate, μ_a , such that the growth rate Δ_a remains constant, leads to decreases in T^* , and therefore to the second case. In this case, increasing the division rate λ_a should also increase the mutation rate $v\lambda_a$ in aneuploid cells, as mutations mostly occur during division, so overall the threshold tumor size N_a^* is unaffected by the division rate λ_a (i.e., $d\lambda_a T^*/d\lambda_a = 0$). Thus, if aneuploid cells rapidly die due to the drug but compensate by rapidly dividing, further increasing the division rate will *not* facilitate adaptation. This is consistent with experimental findings where aneuploidy confers resistance by decreasing the division rate (Replogle et al., 2020).

We can categorize tumors by their size: small tumors with size $N < N_a^*$, intermediate tumors with size $N_a^* < N < N_m^*$, and large tumors with size $N > N_m^*$. For the parameter values in Table 1 with $\lambda_a = 0.0899$, $\mu_w = 0.14$, $u = 10^{-2}$, $v = 10^{-7}$, we are in the tolerant aneuploid case, and substituting in eqs. (3) and (4), we have $N_a^* \approx 4.4 \cdot 10^6$ and $N_m^* \approx 4 \cdot 10^7$. Hence, we obtain the ratio $N_a^*/N_m^* \approx 0.11$ (eq. (5)), that is, aneuploidy reduces the threshold tumor size by approximately 90% Interestingly, the threshold between small and intermediate tumors, N_a^* , is similar to the tumor detection threshold of $4.19 \cdot 10^6$ cells for a wide variety of tumors as reported by Avanzini and Antal (2019).

Density-dependent growth. In our analysis we used branching processes, which assume that growth (division and death) is density-independent. However, growth may be limited by resources (oxygen, nutrients, etc.) and therefore depend on cell density. We therefore performed stochastic simulations of a logistic growth model with a carrying capacity (see Methods). We find that our density-independent approximations agree with results of simulations with density-dependent growth for biologically relevant parameter values (Figure S1).

Standing vs. de-novo genetic variation. In the above we assumed that at the onset of drug treatment, the initial tumor consisted entirely of wildtype cells that are drug sensitive. However, aneuploidy is likely produced even before onset of treatment at some rate $\tilde{u} \leq u$ and confer a deleterious fitness effect s in the absence of the drug (Giam and Rancati, 2015, Replogle et al., 2020), and the aneuploidy rate in the presence of drugs, u, is likely higher than in their absence, \tilde{u} (Mason et al., 2017, Wang et al., 2019). But if the number of cells in the tumor N is large (as expected if the tumor is to be treated with a drug), there may already be a fraction $f \approx \tilde{u} \lambda_w / s$ (we assume that the division rate of the wildtype in a drug-free environment is equal to the division rate of the wildtype in the environment affected by the anti-cancer drug) of aneuploid cells in the population.

Therefore, the ratio of threshold tumor size with standing generation variation, \tilde{N}_a^* , is similar to the ratio with de-novo variation, N_a^* , except that the sensitive growth rate $|\Delta_w|$ is replaced with the aneuploidy cost, s, such that

$$\frac{\tilde{N}_a^*}{N_a^*} = \frac{u}{\tilde{u}} \frac{s}{|\Delta_w|}.$$
 (6)

Comparing the above approximation to results of stochastic evolutionary simulations, we find that the approximations perform very well (Figure 5). Standing genetic variation will drive evolutionary rescue to the drug if wildtype growth rate Δ_w is very negative due to a stronger effect of the drug on sensitive cells, or if the aneuploidy cost s is very small due to a low cost of aneuploidy in the pre-drug conditions. In contrast, de-novo aneuploid cells will have a greater contribution to rescue if the aneuploidy cost, s, is large, the effect of the drug is weak (Δ_w is small), or if the drug induces the appearance of aneuploid cells ($u > \tilde{u}$). For example, with $\lambda_w = 0.1$, $\mu_w = 0.14$, $u = 10^{-2}$, $\tilde{u} = 10^{-3}$, $s = 7.3 \cdot 10^{-3}$, the ratio of the threshold tumor sizes for standing vs de-novo variation is $\tilde{N}_a^*/N_a^* \approx 1.825$, which means that de-novo genetic variation is the main driver of evolutionary rescue.

Using eqs. (3), (4) and (6), we can find the ratio of threshold tumor size for rescue via standing genetic variation to the threshold for direct mutation,

$$\frac{\tilde{N}_{a}^{*}}{N_{m}^{*}} = \frac{\tilde{N}_{a}^{*}}{N_{a}^{*}} \cdot \frac{N_{a}^{*}}{N_{m}^{*}} \approx \frac{s}{|\Delta_{w}|} \begin{cases} \frac{|\Delta_{a}|}{\tilde{u}\lambda_{a}}, & \Delta_{a}T^{*} \ll -1, \\ \frac{1}{\tilde{u}} \left(v \cdot \frac{\Delta_{m}}{\lambda_{m}}\right)^{1/2}, & -1 \ll \Delta_{a}T^{*} \ll 1, \\ v \frac{\Delta_{m}}{\lambda_{m}} \cdot \left(\tilde{u}\frac{\Delta_{a}}{\lambda_{a}}\right)^{-1}, & \Delta_{a}T^{*} \gg 1. \end{cases}$$
(7)

We observe that evolutionary rescue through direct mutation is more likely if the cost of aneuploidy s is very large or the effect of the drug Δ_w is small. In contrast, standing genetic variation will drive adaptation if the pre-treatment chromosome missegregation rate \tilde{u} is very large. The ratio does not depend on the rate of chromosome missegregation induced by the drug u. However, if the aneuploid growth rate Δ_a increases, then evolutionary rescue is driven by standing genetic variation. For the parameter values of $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\tilde{u} = 10^{-3}$, $v = 10^{-7}$, we are in the tolerant aneuploid case and obtain the ratio $\tilde{N}_a^*/N_m^* \approx 0.356$, which means that standing genetic variation reduces the threshold tumor size by approximately 65%. We observe that while standing genetic variation does not drive evolution of drug resistance when compared to de-novo aneuploidy, it does offer an advantage when compared with direct mutation.

Recurrence time due to evolutionary rescue

When evolutionary rescue occurs, the time until recurrence of the tumor may still be very long. We therefore explored the time until recurrence of the tumor, that is, the time until the tumor reaches its original size, N. When the expected number of resistant lineages that avoid extinction is small, the expected recurrence time can be estimated by adding two terms: the *mean evolutionary rescue time*, which is the waiting time for appearance of a resistant lineage that avoids extinction (conditioned on such an even occurring in the first place), and the *mean proliferation time*, which is the expected time for that lineage to grow to N cells. However, when the expected number of resistant lineages is large, the behavior of the size of the population of mutant cells is deterministic (i.e. it is modeled by a system of ODEs, see Equation (D2)) and the mean recurrence time cannot be separated into the mean evolutionary rescue time and mean proliferation time because multiple mutant lineages contribute towards the mutant population size reaching the initial tumor size. Of particular interest is the distribution of the evolutionary rescue time and recurrence time with tolerant aneuploid cells $(\Delta_a T^* \ll 1)$, for which we focus on the parameter values $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, μ_m

Evolutionary rescue time. We have derived approximations for τ_m , the mean evolutionary rescue time without aneuploidy (u = 0), and τ_a , the mean rescue time with aneuploidy (u > 0), both conditioned on evolutionary rescue occurring (see appendix C). These approximations are in good agreement with simulation results for small, intermediate, and large tumor sizes (Figures S2 and 6). The mean rescue time with aneuploidy for small and large tumors follows these expressions:

$$\tau_a \approx \begin{cases} \frac{1}{|\Delta_w|} + \frac{1}{|\Delta_a|}, & N \ll N_a^*, \\ \frac{1}{v\lambda_w N} \cdot \frac{\lambda_m}{\Delta_m}, & N \gg N_m^*. \end{cases}$$
(8)

For small tumors $(N \ll N_a^*)$, the mean rescue time is a function of the wildtype and aneuploid growth rates and independent of the other model parameters, including tumor size (blue line in Figure 6). Increasing the wildtype or aneuploid growth rates leads to an increase in the mean rescue time, because the corresponding cells will survive for longer and will produce additional rescue

mutations at latter times. In our focus parameter regime, we have $\Delta_w = -0.04$ and $\Delta_a = -10^{-4}$, such that the mean rescue time is mainly determined by the aneuploid growth rate, $\tau_a \approx 10^4$.

For large tumors $(N \gg N_m^*)$, the mean evolutionary rescue time (eq. (8)) is independent of parameters characterizing aneuploid cells or their production $(u, \lambda_a, \text{ and } \Delta_a)$. Increasing the per division mutation rate, v, leads to faster appearance of a rescue mutation and hence reduced mean rescue time. Finally, increasing the tumor size leads to shorter mean rescue time, as there are more wildtype cells that can mutate to become resistant.

When a fraction $f \approx 1.36\%$ of the initial cancer cell population is an euploid we calculate the mean evolutionary rescue time in eq. (C10) which we plot in Figure 9. We note that standing genetic variation reduces the mean evolutionary rescue time only for intermediary sized tumors but not for large or small tumors.

Recurrence time. In appendix D we approximate the mean time for the population of mutant cancer cells to reach the initial population size N which we denote as the recurrence time. Figure 7 shows the agreement between our approximations and simulations. For values of N smaller then 10^8 the mean recurrence time can be approximated as the sum of the mean time for the first rescue mutation to appear and the mean time for the lineage generated by that mutation to reach size N. For $N > 10^8$ the behavior of the mutant cancer cell population is deterministic and the mean recurrence time becomes constant. We note that τ_a^r has the following asymptotic behavior:

$$\tau_a^r \approx \begin{cases} \frac{1}{|\Delta_w|} + \frac{1}{|\Delta_a|} + \frac{\log p_m N}{\Delta_m}, & N \ll N_a^*, \\ \frac{1}{\Delta_m} \log \frac{\Delta_m - \Delta_w}{\nu \lambda_w}, & N \gg N_m^*. \end{cases}$$
(9)

In the first case, when tumor is small, the mean recurrence time grows logarithmically with tumor size N and is the same order of magnitude as the mean evolutionary rescue time (Figure 7). Increasing the mutant growth rate Δ_m leads to shorter recurrence times while increasing the wildtype and aneuploid growth rates Δ_m and Δ_a , retrospectively, increases the recurrence time. In the second case, when tumor is large, we observe that increasing either the mutant growth rare Δ_m or the mutation rate ν leads to a decrease in the time needed for the tumor to rebound to its initial size. Additionally, drugs which significantly increase the wildtype death rate μ_w and do not affect the division rate λ_w delays cancer relapse. Consequently, patients which take such drugs require a longer period of monitoring to guarantee the effectiveness of the treatment.

Distribution of evolutionary rescue time with and without aneuploidy. In Appendix E we derive the probability that by time *t* a successful mutant has been generated. Figure 6A show the agreement between our formula and simulation results for the case when aneuploidy is present and when it is absent. We observe that aneuploidy start to have an effect on adaptation for timescales greater then 100 days.

Distribution of recurrence time. In Appendix F we derive the probability that mutant cancer cell population reaches N at time t. Figure 6B shows agreement between our approximations and stochastic simulations for various values of N. Additionally, we derived the distribution of the recurrence time and plotted our result in Figure S4 alongside simulations for the case $N=10^6$ (small tumor) and we note that the distribution is wide and rights-skewed. It is highly unlikely to observe the recurrence of tumors at times smaller then $\frac{1}{\Delta_m} \log \frac{\Delta_m - \Delta_w}{\nu \lambda_w} \approx 1542$ days for the parameter values $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.0$

The recurrence can also be defined as the time necessary for the tumor size to reach a detection threshold M. We derive the mean recurrence time to detection size $M = 10^7$ in appendix D and plot in Figure 8 with simulations. We observe that for small and intermediate sized tumors the effect of

the detection size is negligible when compared to τ_a^r . However, for large tumors the mean recurrence time to detection size M decreases logarithmically with tumor size N.

Most clinical trials report data on the distribution of recurrence time measured from a varying time of surgery (Avanzini and Antal, 2019) with chemotherapy usually following after. Therefore, since only undetected secondary tumors are present at the time of the administration of the anti-cancer drug we lack knowledge of the size of the tumors and we cannot compare the empirical distributions to our predictions. However, we expect that the variability of secondary tumors to average out across large cohorts of patients and, as a result, we can use the mean recurrence time.

Discussion

We have modeled a tumor—a population of cancer cells—exposed to drug treatment that causes the population to decline in size towards potential extinction. In this scenario, the tumor can be "evolutionary rescued", or escape extinction, via two paths. In the direct path, a sensitive cell acquires a mutation that confers resistance that allows it to rapidly grow. In the indirect path, a sensitive cell first becomes aneuploid, which diminishes the effect of the drug, and then an aneuploid cell acquires a mutation that confers resistance (Figure 1).

Using multitype branching processes, we derived the probability of evolutionary rescue of the population of cancer cells under different scenarios for the effect of aneuploidy, ranging from tolerance to partial resistance. We obtained exact and approximate expressions for the probability of evolutionary rescue (eq. (2)). Our results show that the probability of evolutionary rescue increases with the initial tumor size N, the sensitive growth rate Δ_w , the mutation rate v, and the aneuploidy rate u.

Evolutionary rescue When an euploid cells are partially resistant to the drug ($\Delta_w \ll 0 \ll \Delta_a \ll \Delta_m$), evolutionary rescue can be approximated by a one-step process in which an euploidy itself rescues the population (Figure 4A). When an euploidy only provides tolerance to the drug ($\Delta_w \ll \Delta_a \ll 0 \ll \Delta_m$), it cannot rescue the population. Instead, it acts as a *stepping stone* through which the resistant mutant can appear more rapidly, given that the aneuploid cell population declines slower then the sensitive cell population (Figure 2). In this case, an euploidy provides two benefits. First, it delays the extinction of the population–providing more time for appearance of the resistance mutation. Second, it increases the population size relative to a sensitive population–providing more cells in which mutations can occur, i.e., it increases the mutation supply, $Nuv\lambda_w\lambda_a/\Delta_a/\Delta_w\Delta_a$.

We find that aneuploidy can have a significant effect on evolutionary rescue as it reduces the threshold tumor size by at least one order of magnitude even when aneuploidy is tolerant (Figure 3). Interestingly, aneuploidy is unlikely to contribute to evolutionary rescue in primary tumors in which the number of cells is large enough $(N \gg N_m^* \approx 4 \times 10^7)$ for the appearance of resistant mutation directly in sensitive cells before these cells become extinct (Figure 3). However, aneuploidy can have a crucial role in evolutionary rescue of secondary tumors, in which the number of sensitive cells may be below the detection threshold of $\sim 10^7$ (Bozic et al., 2013). This can have an impact on the recurrence of cancer after the resection of the primary tumor through secondary tumors which are too small to be detected and for which chemotherapy is employed to prevent cancer relapse and are estimated to cause the majority of cancer-related deaths (Chaffer and Weinberg, 2011). This is reinforced by the fact that metastases have been shown to have higher chromosome missagregation rate two to three orders of magnitude higher then primary tumors (Kimmel et al., 2023).

Given the fact that the mean time for such secondary tumors to overcome chemotherapy can be of the order of 1000 days (Figure S2A), this can explain the reappearance of cancer even after initial remission. The theoretical predictions for the mean rescue time for tumors smaller then 10^8 is greater then 4 years which is consistent with previous estimates of the recurrence time of tumors after resection (Avanzini and Antal, 2019).

We hypothesized that presence of *standing variation*—the existence of a subpopulation of aneuploid cancer cells before therapy begins—can facilitate evolutionary rescue by reducing the waiting time for the appearance of aneuploid cells. However, we observe that for reasonable parameter values evolutionary rescue is more likely to occur through de-novo aneuploidy (Figure 5). If the the fraction of the tumor cells which have the "beneficial" aneuploidy is $f \gg \frac{u\lambda_w}{|\Delta_w|} \approx 2.5\%$ then evolutionary rescue is more likely to occur through this existing standing variation, rather then through *de novo* aneuploid cells. In that case, evolutionary rescue occurs through a single step by the aneuploid cells acquiring a beneficial mutation and the probability of evolutionary rescue is given by $1 - \exp\left(-N/\tilde{N}_a^*\right)$ (Figure S5).

Experimental future direction Our model's predictions may be tested by experiments (Martin et al., 2013). For example, to study the effects of initial tumor size on the probability of evolutionary rescue, a large culture mass can be propagated from a single cancer cell in permissive conditions and then diluted to a range of starting tumor sizes. Afterwards, these tumors may be exposed to anti-cancer drugs that induces aneuploidy or to saline solution for control (Ippolito et al., 2021). Cell density can then be measured by optical density and a population exposed to the drug is considered extinct if the optical density is lower when compared to the control case with no cells present. We compare to the predictions of our model to see if the tumors with initial size bellow the threshold (4) are more likely to go extinct.

Additionally, we can test our model's predictions with data from patients. Given the type of cancer they have and the degree of "beneficial" aneuploidy, we can obtain the parameters for our model and predict with what probability will the cancer relapse after treatment.

Directions of future research Even though our model has been designed for cancer it can be extended to understand evolutionary rescue in different biological contexts, for example, how yeast subject to stress can overcome extinction with the help of aneuploidy (Pompei and Cosentino Lagomarsino, 2023). Additionally, the heterogeneity of aneuploidy is a fact the we did not account for, as not all the aneuploidy lineages generated have the same growth rate Δ_a . This can be corrected by fixing the aneuploidy death rates and sampling the aneuploidy growth rates from a distribution $f(\Delta_a|\mu_a)$ Martin et al. (2013). Furthermore, tumor heterogeneity is also a factor for pre-existing genetic variation where the fitness of the aneuploid cells can be drawn from a distribution upon which selection acts when the tumor is exposed to a strong stress.

We have assumed that cancer cell lineages are independent of each other. However, this may not be the case, as cancer cells compete for resources (e.g., blood supply). Nevertheless, we find that when the effective carrying capacity is not too large ($K \sim 10^8, K_e \sim 10^6$) our approximation for the probability of evolutionary rescue agrees with results of stochastic simulations with density-dependent growth (Figure S1). Future work may focus on scenarios with small carrying capacity by analyzing density-dependent branching processes (Harris, 1963, Klebaner, 1997).

Solid tumors are spatially heterogeneous with different genotypes inhabiting cellular niches and immune infiltration impacting growth in affected regions (Galon et al., 2010, Varrone et al., 2023). This has the potential to impact the probability of overcoming chemotherapy (Martens et al., 2011). Future work should take into consideration the spatial structure of the tumor and its effect on the probability of evolutionary rescue.

An additional limitation of our model is the choice of parameters which are based on a specific set of assumptions which might not be true for many tumor. As a result, incorporating a wider distribution of parameters would be a beneficial extension for our model.

Conclusions Our paper analyzed the role that aneuploidy plays in tumor adaptation to the effects of anti-cancer drugs. Our study quantitatively confirms that aneuploidy plays an important role in

tumors overcoming exposure to chemotherapeutic drugs when the tumor size is small or intermediate. Very large tumors can escape anti-cancer drugs through direct mutation while smaller ones are able to obtain the beneficial mutation through an aneuploid intermediary (Figure 3). As a result, therapies which increase the rate of aneuploidy in tumors in order to combat cancer might have an adverse effect on patient outcomes.

Acknowledgements

We thank Hildegard Uecker for discussions and comments. This work was supported in part by the Israel Science Foundation (ISF 552/19, YR), the US-Israel Binational Science Foundation (BSF 2021276, YR), Minerva Stiftung Center for Lab Evolution (YR), Ela Kodesz Institute for Research on Cancer Development and Prevention (RS), the Simons Foundation (Investigator in Mathematical Modeling of Living Systems #508600, DBW), the Sloan Foundation Research Fellowship (FG-2021-16667, DBW), the NSF grant (#2146260, DBW),

	Name	Value	Units	References
N	Initial tumor size	$10^7 - 10^9$	cells	Del Monte (2009)
λ_w	Wildtype division rate	0.1	1/days	Bozic et al. (2013), Rew and Wilson (2000)
μ_w	Wildtype death rate	0.11 – 0.17	1/days	Bozic et al. (2013)
λ_a	Aneuploid division rate*	0.06 - 0.1	1/days	-
μ_a	Aneuploid death rate*	0.09	1/days	-
λ_m	Mutant division rate	0.1	1/days	Bozic et al. (2013), Rew and Wilson (2000)
μ_m	Mutant death rate	0.09	1/days	Bozic et al. (2013), Carlson (2003)
и	Missegregation rate	$10^{-3} - 10^{-2}$	1/cell division	Bakker et al. (2023)
v	Mutation rate	$10^{-9} - 10^{-7}$	1/cell division	Bozic et al. (2013), Loeb (2001)
ũ	Missegregation rate in the drug free environment*	10^{-3}	1/cell division	-
S	Selection coefficient of aneu- ploidy in the drug free envi- ronment	$7.3 \cdot 10^{-3}$	1/days	Lukow et al. (2021)

Table 1: Model parameters. We have modified the parameters from Bozic et al. (2013) such that wildtype/mutant division rate is $\lambda_{w,m} = \log 2/T \approx 0.1$ instead of their value of 0.14 where T is the doubling time in the absence of cellular death obtained from Rew and Wilson (2000).

References

- Allen, L. J. (2010), An introduction to stochastic processes with applications to biology, CRC press.
- Avanzini, S. and Antal, T. (2019), 'Cancer recurrence times from a branching process model', *PLoS computational biology* **15**(11), e1007423.
- Bakker, B., Schubert, M., Bolhaqueiro, A. C., Kops, G. J., Spierings, D. C. and Foijer, F. (2023), 'Predicting CIN rates from single-cell whole genome sequencing data using an *in silico* model', *bioRxiv* pp. 2023–02.
- Barton, G. (1989), *Elements of Green's functions and propagation: potentials, diffusion, and waves*, Oxford University Press.
- Ben-David, U. and Amon, A. (2020), 'Context is everything: aneuploidy in cancer', *Nature Reviews Genetics* **21**(1), 44–62.
- Bozic, I., Reiter, J. G., Allen, B., Antal, T., Chatterjee, K., Shah, P., Moon, Y. S., Yaqubie, A., Kelly, N., Le, D. T. et al. (2013), 'Evolutionary dynamics of cancer in response to targeted combination therapy', *eLife* 2, e00747.

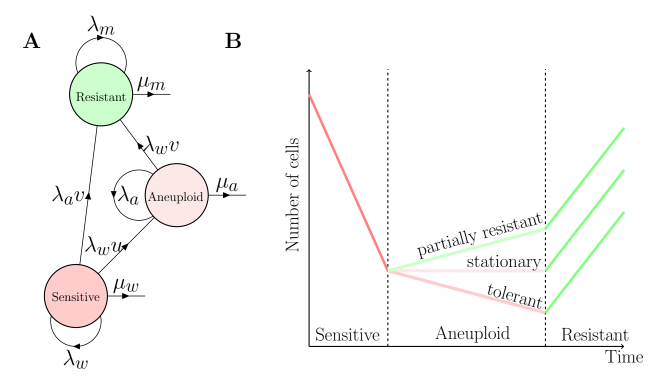


Figure 1: Model illustration. (**A**) A population of cancer cells is composed of wildtype, aneuploid, and mutant cells, which divide with rates λ_w , λ_a , and λ_m and die at rates μ_w , μ_a , and μ_m , respectively. Wildtype cells can divide and become aneuploid at rate $u\lambda_w$. Both aneuploid and wildtype cells can divide and acquire a beneficial mutation with rate $v\lambda_a$ and $v\lambda_w$, respectively. Color denotes the relative growth rates of the three genotypes such that $\lambda_w - \mu_w < \lambda_a - \mu_a < \lambda_m - \mu_m$. (**B**) The wildtype and the mutant are susceptible and resistant, respectively, to the drug. The aneuploid may be tolerant, stationary and partially resistant.

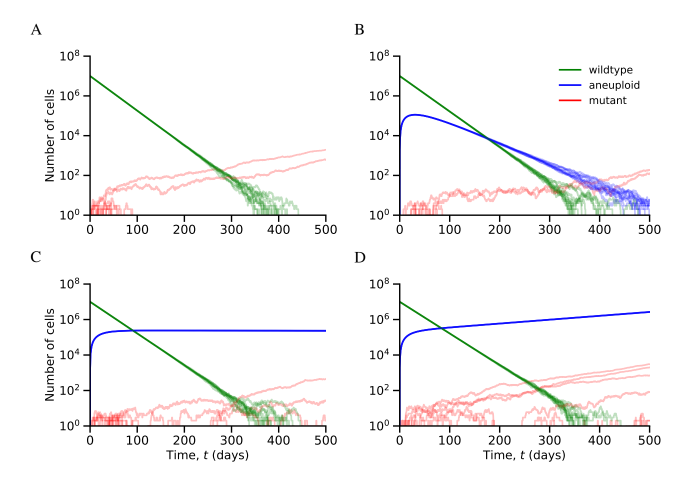


Figure 2: Sample trajectories of the different genotype frequencies. (A) When the missegregation rate u=0 evolutionary rescue is only possible through direct mutation and in most cases the tumor will be killed by the drug. (B) When the aneuploidy growth rate $\Delta_a \ll 0$ we observe similar dynamic to case (A) as direct mutation is the only viable route toward evolutionary rescue for the tumor. (C) Intermediate aneuploid growth rates $\Delta_a \approx 0$ leads to the appearance of aneuploid lineages even after the wildtype population has gone extinct thus increasing the chance of evolutionary rescue. (D) As the growth rate of the aneuploid becomes positive we observe that the tumour is rescued by the aneuploid cancer cell population. Each plot features 10 trajectories of (w_t, a_t, m_t) as a function of time t for the following parameter values: $\lambda_w = 0.1$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\nu = 10^{-7}$, $\nu = 10^{-7}$. For (A) we set $\nu = 0$, for (B) $\nu = 0.065$, $\nu = 10^{-2}$, for (C) $\nu = 0.0899$, $\nu = 10^{-2}$ and for (D) $\nu = 0.095$, $\nu = 10^{-2}$.

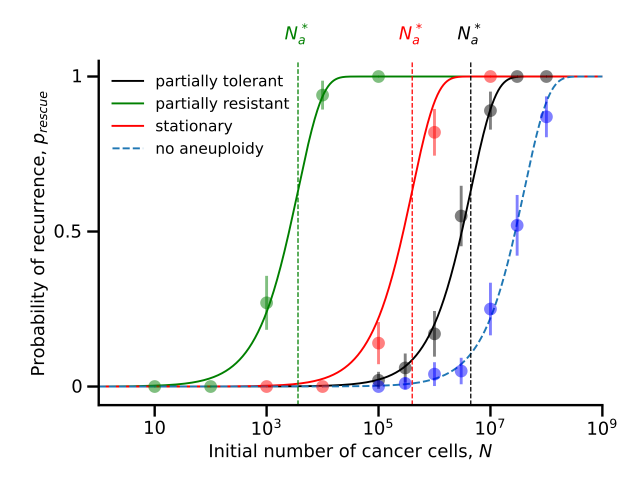


Figure 3: Aneuploidy facilitates evolutionary rescue of cancer under drug treatment. The probability of evolutionary rescue (i.e. the probability that the population does not go to extinction), p_{rescue} , as a function of the initial tumor size, N (see eq. (2)). Dashed vertical line shows the threshold tumor size, N_a^* , above which the probability is very high (see eq. (4)). Blue dashed line represents the probability of evolutionary rescue as a function of N without aneuploidy (u = 0). The black line represents the case with tolerant aneuploidy ($u = 10^{-2}$, $\lambda_a = 0.0899$), the red line represents the case with stationary aneuploidy ($u = 10^{-2}$, $\lambda_a = 0.08999$) and the green line represents the case with partially resistant aneuploidy ($u = 10^{-2}$, $\lambda_a = 0.095$). The dots represent simulations and the error bars represent 95% confidence interval of the form $p \pm 1.96\sqrt{p(1-p)/n}$ where p is the fraction of simulations in which the tumor has adapted to the stress and n = 100 is the number of simulations. Parameters: $\lambda_w = 0.1$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\nu = 10^{-7}$.

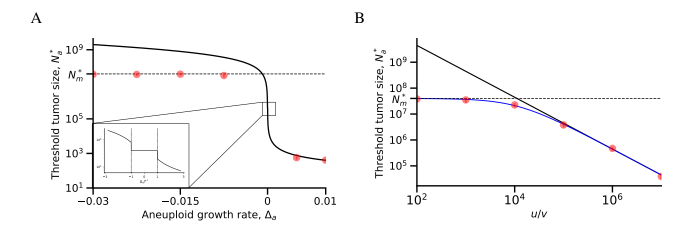


Figure 4: Aneuploidy facilitates the evolutionary rescue of cancer under drug treatment. (A) The threshold tumor size N_a^* as a function of the aneuploid growth rate Δ_a . The dashed horizontal line shows N_m^* , the threshold tumor size without aneuploidy (u=0). When aneuploid growth rate is close to or higher than zero, aneuploidy decreases the threshold tumor size, thereby facilitating evolutionary rescue. The inset highlights the case when aneuploidy cancer cells are non-growing. The red dots represents simulations and the error bars represent the 95% confidence intervals obtained with bootstrapping, see Appendix G. Parameters: $\lambda_w = 0.1, \lambda_m = 0.1, \mu_w = 0.14, \mu_a = 0.09, \mu_m = 0.09, u = 10^{-2}, v = 10^{-7}$. (B) The threshold tumor size N_a^* as a function of the ratio of aneuploidy and mutation rates, u/v. The dashed horizontal line shows N_m^* , the threshold tumor size without aneuploidy (u=0). When the aneuploidy rate is much higher than the mutation rate, aneuploidy decreases the threshold tumor size, thereby facilitating evolutionary rescue. The blue line represents the exact formula for threshold tumor size N_a^* while the solid black line represents the approximation eq. (4). The red dots represents simulations and the error bars represent the 95% confidence intervals obtained with bootstrapping, see Appendix G. Parameters: $\lambda_w = 0.1, \lambda_m = 0.0899, \lambda_m = 0.1, \mu_w = 0.14, \mu_a = 0.09, \mu_m = 0.09, v = 10^{-7}$.

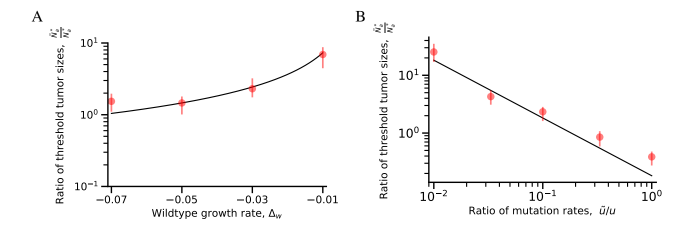


Figure 5: Standing genetic variation facilitates evolutionary rescue of cancer under drug treatment. (A) The ratio of threshold tumor size \tilde{N}_a^* when a fraction $\frac{\tilde{u}\lambda_w}{s}$ is an euploid at the start of treatment and N_a^* as a function of the wildtype growth rate Δ_w . Standing genetic variation will drive adaptation to the drug if Δ_w is very negative due to a stronger effect of the drug on sensitive cells. The red dots represents simulations and the error bars represent the 95% confidence intervals obtained with bootstrapping, see Appendix G. Parameters: $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\tilde{u} = 10^{-3}$, $u = 10^{-2}$, $v = 10^{-7}$. (B) The ratio of threshold tumor size \tilde{N}_a^* when a fraction $\frac{\tilde{u}\lambda_w}{s}$ is an euploid at the start of treatment and N_a^* as a function of the the ratio of an euploidy rates \tilde{u}/u . De-novo an euploids will have larger contribution to the appearance of drug resistance if the drug induces the appearance of an euploid cells ($u \gg \tilde{u}$). The red dots represents simulations and the error bars represent the 95% confidence intervals obtained with bootstrapping, see Appendix G. Parameters: $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\tilde{u} = 10^{-3}$, $v = 10^{-7}$.

Brauner, A., Fridman, O., Gefen, O. and Balaban, N. Q. (2016), 'Distinguishing between resistance, tolerance and persistence to antibiotic treatment', *Nature Reviews Microbiology* **14**(5), 320–330.

Carja, O. and Plotkin, J. B. (2017), 'The evolutionary advantage of heritable phenotypic heterogeneity', *Scientific reports* **7**(1), 1–12.

Carja, O. and Plotkin, J. B. (2019), 'Evolutionary rescue through partly heritable phenotypic variability', *Genetics* **211**(3), 977–988.

Carlson, J. A. (2003), 'Tumor doubling time of cutaneous melanoma and its metastasis', *The American journal of dermatopathology* **25**(4), 291–299.

Chaffer, C. L. and Weinberg, R. A. (2011), 'A perspective on cancer cell metastasis', *science* **331**(6024), 1559–1564.

Cobbold, C. A. and Stana, R. (2020), 'Should I stay or should I go: partially sedentary populations can outperform fully dispersing populations in response to climate-induced range shifts', *Bulletin of Mathematical Biology* **82**(2), 1–21.

Del Monte, U. (2009), 'Does the cell number 10⁹ still really fit one gram of tumor tissue?', *Cell cycle* **8**(3), 505–506.

Galon, J., Dieu-Nosjean, M., Tartour, E., Sautes-Fridman, C., Fridman, W. et al. (2010), 'Immune infiltration in human tumors: a prognostic factor that should not be ignored', *Oncogene* **29**(8), 1093–1102.

Giam, M. and Rancati, G. (2015), 'Aneuploidy and chromosomal instability in cancer: a jackpot to chaos', *Cell division* **10**(1), 1–12.

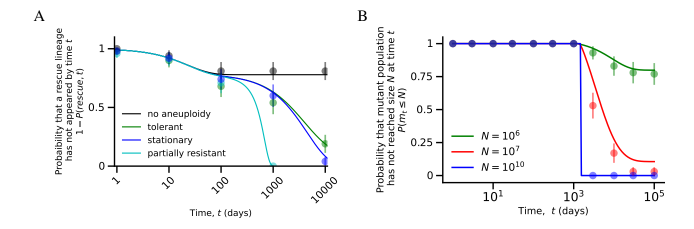


Figure 6: Aneuploidy has a impact on cancer relapse early on. (A) The probability that a successful mutant has not appeared by time t. The green line represents the case with tolerant aneuploidy ($u > 0, \lambda_a = 0.0899$), the blue line represents the case with non-growing aneuploidy $(u > 0, \lambda_a = 0.089999)$, the cyan line represents the case with partially resistant aneuploidy $(u > 0, \lambda_a = 0.089999)$ $0, \lambda_a = 0.095$) and the black line represents the case without an euploidy (u = 0). As time increases, aneuploidy plays an important role in helping the cancer cell population escape extinction. The markers represent simulations and the error bars represent 95% confidence interval of the form $p \pm 1.96 \sqrt{p(1-p)}/n$ where p is the fraction of simulations in which a successful mutant has not been generated and n=100 is the number of simulations. Parameters: $\lambda_w=0.1, \lambda_m=0.1, \mu_w=0.1$ $0.14, \mu_a = 0.09, \mu_m = 0.09, u = 10^{-2}, v = 10^{-7}, N = 10^{7}$. (B) The probability that a mutant cancer cell population has not reached size N at time t. The green line represents the case where $N = 10^6$ (small tumor), the red line represents $N = 10^7$ (intermediate sized tumor) and the blue line represents the case where $N = 10^{10}$ (large tumor). Increasing the initial tumor size guarantees that the tumor will regrow. The markers represent simulations and the error bars represents 95% confidence interval of the form $p \pm 1.96\sqrt{p(1-p)}/n$ where p is the fraction of the simulations in which the mutant population size has not reached N and n = 100 is the number of simulations. Parameters: $\lambda_w = 0.1, \lambda_a = 0.0899, \lambda_m = 0.1, \mu_w = 0.14, \mu_a = 0.09, \mu_m = 0.09, u = 10^{-2}, v = 10^{-7}.$

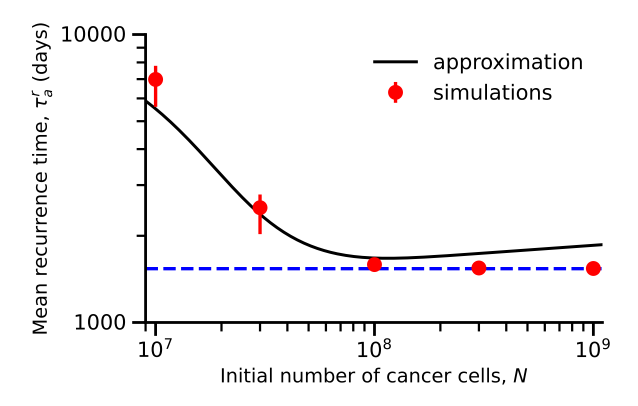


Figure 7: Tumor size decreases the mean recurrence time. The mean time for the mutant cell population to reach size N, the initial number of cancer cells. Our inhomogeneous Poisson-process approximation (solid black line, eq. (D1)) is in agreement with simulation results (red markers with 95% confidence interval obtained with bootstrapping, see Appendix G) for intermediary N. The simulations converge to eq. (D4) (blue dashed line) for large values of N. Parameters: $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\mu_m = 10^{-2}$, $\nu = 10^{-7}$.

- Gillespie, D. T. (1976), 'A general method for numerically simulating the stochastic time evolution of coupled chemical reactions', *Journal of computational physics* **22**(4), 403–434.
- Gillespie, D. T. (1977), 'Exact stochastic simulation of coupled chemical reactions', *The journal of physical chemistry* **81**(25), 2340–2361.
- Gillespie, D. T. (2001), 'Approximate accelerated stochastic simulation of chemically reacting systems', *The Journal of chemical physics* **115**(4), 1716–1733.
- Gisselsson, D., Jin, Y., Lindgren, D., Persson, J., Gisselsson, L., Hanks, S., Sehic, D., Mengelbier, L. H., Øra, I., Rahman, N. et al. (2010), 'Generation of trisomies in cancer cells by multipolar mitosis and incomplete cytokinesis', *Proceedings of the National Academy of Sciences* **107**(47), 20489–20493.
- Gomulkiewicz, R. and Holt, R. D. (1995), 'When does evolution by natural selection prevent extinction?', *Evolution* pp. 201–207.
- Gunnarsson, E. B., De, S., Leder, K. and Foo, J. (2020), 'Understanding the role of phenotypic switching in cancer drug resistance', *Journal of theoretical biology* **490**, 110162.
- Harris, T. E. (1963), The theory of branching processes, Vol. 6, Springer Berlin.
- Harsch, M. A., Zhou, Y., HilleRisLambers, J. and Kot, M. (2014), 'Keeping pace with climate change: stage-structured moving-habitat models', *The American Naturalist* **184**(1), 25–37.
- Ippolito, M. R., Martis, V., Martin, S., Tijhuis, A. E., Hong, C., Wardenaar, R., Dumont, M., Zerbib, J., Spierings, D. C., Fachinetti, D. et al. (2021), 'Gene copy-number changes and chromosomal instability induced by aneuploidy confer resistance to chemotherapy', *Developmental cell* **56**(17), 2440–2454.
- Kimmel, G. J., Beck, R. J., Yu, X., Veith, T., Bakhoum, S., Altrock, P. M. and Andor, N. (2023), 'Intratumor heterogeneity, turnover rate and karyotype space shape susceptibility to missegregation-induced extinction', *PLOS Computational Biology* **19**(1), e1010815.
- Klebaner, F. (1997), Population and density dependent branching processes, *in* 'Classical and modern branching processes', Springer, pp. 165–169.
- Kocarnik, J. M., Compton, K., Dean, F. E., Fu, W., Gaw, B. L., Harvey, J. D., Henrikson, H. J., Lu, D., Pennini, A., Xu, R. et al. (2022), 'Cancer incidence, mortality, years of life lost, years lived with disability, and disability-adjusted life years for 29 cancer groups from 2010 to 2019: a systematic analysis for the global burden of disease study 2019', *JAMA oncology* **8**(3), 420–444.
- Lee, H.-S., Lee, N. C., Kouprina, N., Kim, J.-H., Kagansky, A., Bates, S., Trepel, J. B., Pommier, Y., Sackett, D. and Larionov, V. (2016), 'Effects of anticancer drugs on chromosome instability and new clinical implications for tumor-suppressing therapies', *Cancer research* **76**(4), 902–911.
- Levien, E., Min, J., Kondev, J. and Amir, A. (2021), 'Non-genetic variability in microbial populations: survival strategy or nuisance?', *Reports on Progress in Physics* **84**(11), 116601.
- Loeb, L. A. (2001), 'A mutator phenotype in cancer', Cancer research 61(8), 3230–3239.
- Lukow, D. A., Sausville, E. L., Suri, P., Chunduri, N. K., Wieland, A., Leu, J., Smith, J. C., Girish, V., Kumar, A. A., Kendall, J. et al. (2021), 'Chromosomal instability accelerates the evolution of resistance to anti-cancer therapies', *Developmental cell* **56**(17), 2427–2439.

- Martens, E. A., Kostadinov, R., Maley, C. C. and Hallatschek, O. (2011), 'Spatial structure increases the waiting time for cancer', *New journal of physics* **13**(11), 115014.
- Martin, G., Aguilée, R., Ramsayer, J., Kaltz, O. and Ronce, O. (2013), 'The probability of evolutionary rescue: towards a quantitative comparison between theory and evolution experiments', *Philosophical Transactions of the Royal Society B: Biological Sciences* **368**(1610), 20120088.
- Mason, J. M., Wei, X., Fletcher, G. C., Kiarash, R., Brokx, R., Hodgson, R., Beletskaya, I., Bray, M. R. and Mak, T. W. (2017), 'Functional characterization of cfi-402257, a potent and selective mps1/ttk kinase inhibitor, for the treatment of cancer', *Proceedings of the National Academy of Sciences* **114**(12), 3127–3132.
- Orr, H. A. and Unckless, R. L. (2008), 'Population extinction and the genetics of adaptation', *The American Naturalist* **172**(2), 160–169.
- Orr, H. A. and Unckless, R. L. (2014), 'The population genetics of evolutionary rescue', *PLoS genetics* **10**(8), e1004551.
- Pompei, S. and Cosentino Lagomarsino, M. (2023), 'A fitness trade-off explains the early fate of yeast aneuploids with chromosome gains', *Proceedings of the National Academy of Sciences* **120**(15), e2211687120.
- Replogle, J. M., Zhou, W., Amaro, A. E., McFarland, J. M., Villalobos-Ortiz, M., Ryan, J., Letai, A., Yilmaz, O., Sheltzer, J., Lippard, S. J. et al. (2020), 'Aneuploidy increases resistance to chemotherapeutics by antagonizing cell division', *Proceedings of the National Academy of Sciences* **117**(48), 30566–30576.
- Rew, D. and Wilson, G. (2000), 'Cell production rates in human tissues and tumours and their significance. part ii: clinical data', *European Journal of Surgical Oncology (EJSO)* **26**(4), 405–417.
- Rutledge, S. D., Douglas, T. A., Nicholson, J. M., Vila-Casadesús, M., Kantzler, C. L., Wangsa, D., Barroso-Vilares, M., Kale, S. D., Logarinho, E. and Cimini, D. (2016), 'Selective advantage of trisomic human cells cultured in non-standard conditions', *Scientific reports* **6**(1), 22828.
- Schukken, K. M. and Foijer, F. (2018), 'CIN and aneuploidy: different concepts, different consequences', *Bioessays* **40**(1), 1700147.
- Smith, J. C. and Sheltzer, J. M. (2018), 'Systematic identification of mutations and copy number alterations associated with cancer patient prognosis', *elife* 7, e39217.
- Tanaka, M. M. and Wahl, L. M. (2022), 'Surviving environmental change: when increasing population size can increase extinction risk', *Proceedings of the Royal Society B* **289**(1976), 20220439.
- Uecker, H. and Hermisson, J. (2011), 'On the fixation process of a beneficial mutation in a variable environment', *Genetics* **188**(4), 915–930.
- Uecker, H. and Hermisson, J. (2016), 'The role of recombination in evolutionary rescue', *Genetics* **202**(2), 721–732.
- Uecker, H., Otto, S. P. and Hermisson, J. (2014), 'Evolutionary rescue in structured populations', *The American Naturalist* **183**(1), E17–E35.
- Uecker, H., Setter, D. and Hermisson, J. (2015), 'Adaptive gene introgression after secondary contact', *Journal of mathematical biology* **70**, 1523–1580.
- Van Rossum, G. and Others (2007), Python programming language, in 'USENIX Annu. Tech. Conf.'.

- Varrone, M., Tavernari, D., Santamaria-Martínez, A., Walsh, L. A. and Ciriello, G. (2023), 'Cellcharter reveals spatial cell niches associated with tissue remodeling and cell plasticity', *Nature Genetics* pp. 1–11.
- Wang, S., Zhang, M., Liang, D., Sun, W., Zhang, C., Jiang, M., Liu, J., Li, J., Li, C., Yang, X. et al. (2019), 'Molecular design and anticancer activities of small-molecule monopolar spindle 1 inhibitors: A medicinal chemistry perspective', *European Journal of Medicinal Chemistry* **175**, 247–268.
- Weissman, D. B., Desai, M. M., Fisher, D. S. and Feldman, M. W. (2009), 'The rate at which asexual populations cross fitness valleys', *Theoretical population biology* **75**(4), 286–300.
- Weissman, D. B., Feldman, M. W. and Fisher, D. S. (2010), 'The rate of fitness-valley crossing in sexual populations', *Genetics* **186**(4), 1389–1410.
- Wilson, B. A., Pennings, P. S. and Petrov, D. A. (2017), 'Soft selective sweeps in evolutionary rescue', *Genetics* **205**(4), 1573–1586.
- Zhou, Y. (2022), 'Range shifts under constant-speed and accelerated climate warming', *Bulletin of Mathematical Biology* **84**(1), 1.

Appendices

Appendix A Survival probability of a single lineage

To analyze evolutionary rescue in this model, we use the framework of *multitype branching processes* (Harris, 1963, Weissman et al., 2009). This allows us to find explicit expressions for the *survival probability*: the probability that a lineage descended from a single cell does not become extinct.

Let p_w , p_a , and p_m be the survival probabilities of a population consisting initially of single wildtype cell, an euploid cell, or mutant cell, respectively. The complements $1 - p_w$, $1 - p_a$, and $1 - p_m$ are the extinction probabilities, which satisfy each its respective equation (Harris, 1963),

$$1 - p_{w} = \frac{\mu_{w}}{\lambda_{w} + \mu_{w} + u\lambda_{w} + v\lambda_{w}} + \frac{u\lambda_{w}}{\lambda_{w} + \mu_{w} + u\lambda_{w} + v\lambda_{w}} (1 - p_{a}) (1 - p_{w}) + \frac{\lambda_{w}}{\lambda_{w} + \mu_{w} + u\lambda_{w} + v\lambda_{w}} (1 - p_{w})^{2} + \frac{v\lambda_{w}}{\lambda_{w} + \mu_{w} + u\lambda_{w} + v\lambda_{w}} (1 - p_{m}) (1 - p_{w}),$$

$$1 - p_{a} = \frac{\mu_{a}}{\lambda_{a} + \mu_{a} + v\lambda_{a}} + \frac{v\lambda_{a}}{\lambda_{a} + \mu_{a} + v\lambda_{a}} (1 - p_{m}) (1 - p_{a}) + \frac{\lambda_{a}}{\lambda_{a} + \mu_{a} + v\lambda_{a}} (1 - p_{a})^{2},$$

$$1 - p_{m} = \frac{\mu_{m}}{\lambda_{m} + \mu_{m}} + \frac{\lambda_{m}}{\lambda_{m} + \mu_{m}} (1 - p_{m})^{2}.$$
(A1)

The survival probabilities are given by the smallest solution for each quadratic equation (Uecker et al., 2015). Therefore we have

$$p_{w} = \frac{\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m} + \sqrt{\left(\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m}\right)^{2} + 4\lambda_{w}^{2}\left(up_{a} + vp_{m}\right)}}{2\lambda_{w}},$$

$$p_{a} = \frac{\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m} + \sqrt{\left(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m}\right)^{2} + 4\lambda_{a}^{2}vp_{m}}}{2\lambda_{a}},$$

$$p_{m} = \frac{\lambda_{m} - \mu_{m}}{\lambda_{m}}.$$
(A2)

Note that the equation for p_w depends on both p_a and p_m , and the equation for p_a depends on p_m . To proceed, we can plug the solution for p_m and p_a into the solution for p_w . We perform this for three different scenarios.

Scenario 1: Aneuploid cells are partially resistant

We first assume that an euploidy provides partial resistance to drug therapy, $\lambda_a > \mu_a$, and that this resistance is significant, $(\lambda_a - \mu_a - v\lambda_a p_m)^2 > 4\lambda_a^2 v p_m$. We thus rewrite eq. (A2) as

$$p_{w} = \frac{\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m}}{2\lambda_{w}} \left(1 - \sqrt{1 + \frac{4\lambda_{w}^{2}\left(vp_{m} + up_{a}\right)}{\left(\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m}\right)^{2}}}\right), \text{ and}$$

$$p_{a} = \frac{\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m}}{2\lambda_{a}} \left(1 + \sqrt{1 + \frac{4\lambda_{a}^{2}vp_{m}}{\left(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m}\right)^{2}}}\right).$$

Using the quadratic Taylor expansion $\sqrt{1+x} = 1 + x/2 + O(x^2)$ and assuming $u, v \ll 1$, we obtain the following approximation for the survival probability of a population initially consisting of a single wildtype cell,

$$p_{w} \approx -\frac{v\lambda_{w}p_{m} + u\lambda_{w}p_{a}}{\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m}}$$
(A3)

$$\approx -\frac{1}{\lambda_w - \mu_w} \left[\frac{u\lambda_w \left(\lambda_a - \mu_a \right)}{\lambda_a} + \frac{uv\lambda_w \lambda_a \left(\lambda_m - \mu_m \right)}{\lambda_m \left(\lambda_a - \mu_a \right)} + \frac{v\lambda_w \left(\lambda_m - \mu_m \right)}{\lambda_m} \right].$$

Now *uv* is very small, and if we use the fact that $v \ll u$, we have:

$$p_w \approx \frac{u\lambda_w}{|\Delta_w|} \cdot \frac{\Delta_a}{\lambda_a}.\tag{A4}$$

However, if aneuploidy is very rare such that

$$\frac{u\lambda_w\Delta_a}{\lambda_a}<\frac{v\lambda_w\Delta_m}{\lambda_m}\Rightarrow u\lambda_a<\frac{v\lambda_a^2\Delta_m}{\lambda_m}\cdot\frac{1}{\Delta_a}<\frac{v\lambda_a^2\Delta_m}{\lambda_m}\cdot\frac{1}{\sqrt{4\lambda_a^2vp_m}}\Rightarrow u\lambda_a< T^*,$$

where $T^* = (4v\lambda_a^2\Delta_m/\lambda_m)^{-1/2}$ and in the second inequality we used the fact that $\Delta_a^2 > 4\lambda_a^2vp_m$. In this case adaptation is through direct mutation and:

$$p_w pprox rac{v\lambda_w}{|\Delta_w|} \cdot rac{\Delta_m}{\lambda_m}.$$

Scenario 2: Aneuploid cells are tolerant.

We now assume that an euploidy provides tolerance to drug therapy, that is, the number of an euploid cells significantly declines over time, but at a lower rate than the number of wildtype cells, $\lambda_w - \mu_w < \lambda_a - \mu_a < 0$. We also assume that the decline are significant, $(\lambda_a - \mu_a - v\lambda_a p_m)^2 > 4\lambda_a^2 v p_m$. We rewrite eq. (A2) as

$$p_{w} = \frac{\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m}}{2\lambda_{w}} \left(1 - \sqrt{1 + \frac{4\lambda_{w}^{2}\left(vp_{m} + up_{a}\right)}{\left(\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m}\right)^{2}}}\right),$$

$$p_{a} = \frac{\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m}}{2\lambda_{a}} \left(1 - \sqrt{1 + \frac{4\lambda_{a}^{2}vp_{m}}{\left(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m}\right)^{2}}}\right).$$
(A5)

Since $u, v \ll 1$, the term in the root can be approximated using a 1st-order Taylor expansion. So, substituting the expressions for p_a and p_m , we have

$$p_{w} \approx -\frac{v\lambda_{w}p_{m} + u\lambda_{w}p_{a}}{\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m}}$$

$$\approx \frac{1}{\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m}} \left[\frac{uv\lambda_{w}\lambda_{a} (\lambda_{m} - \mu_{m})}{\lambda_{m} (\lambda_{a} - \mu_{a} - v\lambda_{a})} - \frac{v\lambda_{w} (\lambda_{m} - \mu_{m})}{\lambda_{m}} \right]$$

$$\approx \frac{v\lambda_{w} (\lambda_{m} - \mu_{m})}{\lambda_{m} (\lambda_{w} - \mu_{w})} \left[\frac{u\lambda_{a}}{(\lambda_{a} - \mu_{a})} - 1 \right]$$

$$= \frac{v\lambda_{w}\Delta_{m}}{\lambda_{m}|\Delta_{w}|} \left(\frac{u\lambda_{a}}{|\Delta_{a}|} + 1 \right).$$
(A6)

If we assume that $u\lambda_a > |\Delta_a|$ then we have:

$$p_w \approx \frac{u\lambda_w}{|\Delta_w|} \cdot \frac{v\lambda_a}{|\Delta_a|} \cdot \frac{\Delta_m}{\lambda_m}.$$
 (A7)

Scenario 3: Aneuploid cells are stationary

We now assume that the growth rate of an euploid cells is close to zero (either positive or negative), such that $(\Delta_a - v\lambda_a p_m)^2 \ll 4\lambda_a^2 v p_m$. We rewrite eq. (A2) as

$$p_{a} = \frac{\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m} + 2\sqrt{\lambda_{a}^{2}vp_{m}} \left(1 + \frac{\left(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m}\right)^{2}}{4\lambda_{a}^{2}vp_{m}}\right)^{\frac{1}{2}}}{2\lambda_{a}}.$$
(A8)

Using a following Taylor series expansion for small $(\lambda_a - \mu_a - v\lambda_a p_m)^2 / 4\lambda_a^2 v p_m$,

$$\left(1+\frac{\left(\lambda_a-\mu_a-v\lambda_a p_m\right)^2}{4\lambda_a^2 v p_m}\right)^{\frac{1}{2}}=1+\frac{\left(\lambda_a-\mu_a-v\lambda_a p_m\right)^2}{8\lambda_a^2 v p_m}+\cdots,$$

we obtain the approximation

$$p_{a} \approx \frac{\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m} + 2\sqrt{\lambda_{a}^{2}vp_{m}} \left[1 + \frac{(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m})^{2}}{8\lambda_{a}^{2}vp_{m}}\right]}{2\lambda_{a}}$$

$$= \frac{\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m} + 2\sqrt{\lambda_{a}^{2}vp_{m}} + \frac{(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m})^{2}}{4\sqrt{\lambda_{a}^{2}vp_{m}}}}{2\lambda_{a}}$$

$$= \frac{\left(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m} + 2\sqrt{\lambda_{a}^{2}vp_{m}}\right)^{2} + 4\lambda_{a}^{2}vp_{m}}{8\lambda_{a}\sqrt{\lambda_{a}^{2}vp_{m}}}$$

$$= \frac{4\lambda_{a}^{2}vp_{m} + 4\lambda_{a}^{2}vp_{m}\left(1 + \frac{\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m}}{2\sqrt{\lambda_{a}^{2}vp_{m}}}\right)^{2}}{8\lambda_{a}\sqrt{\lambda_{a}^{2}vp_{m}}}$$

$$= \frac{1}{2\lambda_{a}}\left(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m} + 2\sqrt{\lambda_{a}^{2}vp_{m}}\right).$$
(A9)

Plugging this in eq. (A3), the survival probability of a population starting from one wildtype individual is

$$p_{w} \approx -\frac{1}{\lambda_{w} - \mu_{w} - u\lambda_{w}p_{a} - v\lambda_{w}p_{m}} \left[v\lambda_{w} \frac{\lambda_{m} - \mu_{m}}{\lambda_{m}} + \frac{u\lambda_{w}}{2\lambda_{a}} \left(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m} + 2\sqrt{\lambda_{a}^{2}vp_{m}} \right) \right]$$

$$= -\frac{1}{\lambda_{w} - \mu_{w} - u\lambda_{w} - v\lambda_{w}} \left[v\lambda_{w} \frac{\lambda_{m} - \mu_{m}}{\lambda_{m}} + \frac{u\lambda_{w}}{2\lambda_{a}} \left(\lambda_{a} - \mu_{a} - v\lambda_{a}p_{m} \right) + u\lambda_{w} \sqrt{\frac{v\left(\lambda_{m} - \mu_{m}\right)}{\lambda_{m}}} \right]$$

$$\approx -\frac{1}{\Delta_{w}} \left[v\lambda_{w} \frac{\Delta_{m}}{\lambda_{m}} + \frac{u\lambda_{w}\left(\Delta_{a} - v\lambda_{a}\right)}{2\lambda_{a}} + u\lambda_{w} \sqrt{\frac{v\Delta_{m}}{\lambda_{m}}} \right]. \tag{A10}$$

Using the fact that

$$\left(\Delta_a - v\lambda_a p_m\right)^2 \ll 4\lambda_a^2 v p_m \Rightarrow \frac{\Delta_a - v\lambda_a p_m}{2\lambda_a} \ll \sqrt{\frac{v\lambda_a \Delta_m}{\lambda_m}},$$

and $v \ll u$ we obtain:

$$p_w \approx \frac{u\lambda_w}{|\Delta_w|} \cdot \sqrt{\frac{v\lambda_a \Delta_m}{\lambda_m}}.$$
 (A11)

Appendix B Evolutionary rescue probability

Using the fact that $\Delta_a - v\lambda_a p_m \approx \Delta_a$ we write the condition $(\Delta_a - v\lambda_a p_m)^2 \ll 4\lambda_a^2 v p_m$ as:

$$\Delta_a^2 \ll 4\lambda_a^2 v p_m \Rightarrow -1 \ll \Delta_a T^* \ll 1,$$

where $T^* = (4v\lambda_a^2\Delta_m/\lambda_m)^{-1/2}$. Substituting eqs. (A4), (A7) and (A11) into eq. (2), the evolutionary rescue probability can be approximated by

$$p_{\text{rescue}} \approx \begin{cases} 1 - \exp\left[-\frac{u\lambda_{a}}{|\Delta_{w}|} \cdot \frac{v\lambda_{w}}{|\Delta_{a}|} \cdot \frac{\Delta_{m}}{\lambda_{m}} \cdot N\right], & \Delta_{a}T^{*} \ll -1, \\ 1 - \exp\left[-\frac{u\lambda_{w}}{|\Delta_{w}|} \cdot \sqrt{\frac{v\lambda_{a}\Delta_{m}}{\lambda_{m}}} \cdot N\right], & -1 \ll \Delta_{a}T^{*} \ll 1, \\ 1 - \exp\left[-\frac{u\lambda_{w}}{|\Delta_{w}|} \cdot \frac{\Delta_{a}}{\lambda_{a}} \cdot N\right], & 1 \ll \Delta_{a}T^{*}. \end{cases}$$
(B1)

Appendix C Evolutionary rescue time

We first calculate the expected time for the appearance of the first mutant that rescues the cell population. This can occur either through the evolutionary trajectory $wildtype \rightarrow mutant$ or through the trajectory $wildtype \rightarrow aneuploid \rightarrow mutant$. We start with the former.

Assuming no aneuploidy (u=0), we define T_m to be the time at which the first mutant cell appears that will avoid extinction and will therefore rescue the population. Note that if extinction occurs, that is the frequency of mutants after a very long time is zero, $m_{\infty}=0$, then it is implied that $T_m=\infty$, and vice versa if $T_m<\infty$ then $m_{\infty}>0$.

The number of successful mutants generated until time t can be approximated by an inhomogeneous Poisson process with rate $R_m(t) = v\lambda_w p_m w_t$, where $w_t = Ne^{\Delta_w t}$ is the number of wildtype cells at time t. Note that

$$\int_0^t R_m(z) dz = v \lambda_w p_m N \frac{\exp[\Delta_w t] - 1}{\Delta_w} \approx v \lambda_w p_m N t, \tag{C1}$$

by integrating the exponential and because $\frac{\exp[\Delta_w t]-1}{\Delta_w} = \frac{1+\Delta_w t+O(t^2)-1}{\Delta_w} = t+O(t^2)$. The probability density function of T_m is thus $R_m(t) \exp\left(-\int_0^t R_m(z) dz\right)$ (Allen, 2010). Therefore, the probability density function of the conditional random variable $(T_m \mid T_m < \infty)$ is $f_m(t) = \frac{R_m(t) \exp\left(-\int_0^t R_m(z) dz\right)}{D_{\text{Procuse}}}$.

We are interested in the mean conditional time, $\tau_m = \mathbb{E}\left[T_m \mid T_m < \infty\right]$, which is given by

$$\tau_m = \int_0^\infty t f_m(t) dt = \frac{\int_0^\infty t R_m(t) \exp\left(-\int_0^t R_m(z) dz\right) dt}{p_{rescue}},$$
 (C2)

Therefore, plugging eqs. (2) and (C1) in eq. (C2),

$$\tau_m = \int_0^\infty t v \lambda_w N e^{\Delta_w t} \frac{e^{-v \lambda_w N p_m e^{\Delta_w t} - 1}}{1 - (1 - p_w)^N} dt \approx \int_0^\infty t v \lambda_w N e^{\Delta_w t} \frac{e^{-v \lambda_w N p_m t}}{1 - e^{-N p_w}} dt.$$
 (C3)

Figure S2B show the agreement between this approximating and simulation results. Assuming an euploidy is possible (u > 0), we define T_a to be the time at which the first mutant

cell appears that will rescue the population. We are interested in the mean conditional time, $\tau_a = \mathbb{E}[T_a \mid T_a < \infty]$.

When $Nu\lambda_w/|\Delta_w| \gg 1$ the aneuploid frequency dynamics is roughly deterministic and therefore can be approximated by

$$a_t \approx \frac{Nu\lambda_w e^{\Delta_w t}}{\Delta_w - \Delta_a} \left[1 - e^{-(\Delta_w - \Delta_a)t} \right].$$
 (C4)

As a result, the number of successful mutants created by direct mutation and via an euploidy can be approximated by inhomogeneous Poisson processes with the rates

$$r_1(t) = v\lambda_a p_m \int_0^t a_z dz = \frac{uv\lambda_w \lambda_a N p_m}{\Delta_w - \Delta_a} \left(\frac{e^{\Delta_w t} - 1}{\Delta_w} - \frac{e^{\Delta_a t} - 1}{\Delta_a} \right), \tag{C5}$$

$$r_2(t) = v\lambda_w p_m \int_0^t w_z dz = v\lambda_w N p_m \frac{e^{\Delta_w t} - 1}{\Delta_w}.$$
 (C6)

For large initial population sizes we assume that the two processes are independent and as a result, they can be merged into a single Poisson process with rate $R_a(t) = (r_1 + r_2)(t)$. Consequently, the mean time to the appearance of the first rescue mutant is

$$\tau_{a} = \frac{\int_{0}^{\infty} t R_{a}(t) \exp\left(-\int_{0}^{t} R_{a}(z) dz\right) dt}{p_{rescue}}$$

$$= \int_{0}^{\infty} t \left(v \lambda_{a} p_{m} a_{t} + v \lambda_{w} p_{m} w_{t}\right) \frac{\exp\left[-\frac{u v \lambda_{w} \lambda_{a} N p_{m}}{\Delta_{w} - \Delta_{a}} \left(\frac{e^{\Delta_{w} t} - 1}{\Delta_{w}} - \frac{e^{\Delta_{a} t} - 1}{\Delta_{a}}\right) - v \lambda_{w} N p_{m} \frac{e^{\Delta_{w} t} - 1}{\Delta_{w}}\right]}{1 - e^{-N p_{w}}} dt, \quad (C7)$$

which we plot in Figure S2A as a function of the initial population size, N.

Paradoxically, we observe from Figure S2 that the mean time of a rescue mutation to appear is significantly shorter for the case when u = 0 when compared to the case u > 0, however this can be explained by the fact this mean time is conditioned on evolutionary rescue and, as a result, an euploidy increase the *window of opportunity* in which a rescue mutation could appear thus increasing the mean time as well (Figure 2).

If $N \gg N_m^*$ then the mean time τ_a can be written as:

$$\tau_{a} = \int_{0}^{\infty} e^{-R_{a}(\tau)} d\tau = \int_{0}^{\infty} \exp \left[-\frac{uv\lambda_{w}\lambda_{a}Np_{m}}{\Delta_{w} - \Delta_{a}} \left(\frac{e^{\Delta_{w}\tau} - 1}{\Delta_{w}} - \frac{e^{\Delta_{a}\tau} - 1}{\Delta_{a}} \right) - v\lambda_{w}Np_{m} \frac{e^{\Delta_{w}\tau} - 1}{\Delta_{w}} \right] d\tau,$$

and we use the following Taylor series expansions:

$$\frac{e^{\Delta_w \tau} - 1}{\Delta_w} = \frac{1 + \Delta_w \tau + O(\tau^2) - 1}{\Delta_w} = \tau + O(\tau^2).$$
$$\frac{e^{\Delta_a \tau} - 1}{\Delta_a} = \frac{1 + \Delta_a \tau + O(\tau^2) - 1}{\Delta_a} = \tau + O(\tau^2),$$

to obtain a simpler approximation for τ_a :

$$\tau_a \approx \int_0^\infty e^{-v\lambda_w N p_m \tau} d\tau = \frac{1}{v\lambda_w N p_m}.$$
 (C8)

If $N \ll N_a^*$ then we can write Equation (C7) as:

$$\tau_{a} \approx \frac{\int_{0}^{\infty} t v \lambda_{a} p_{m} a_{\tau} d\tau}{1 - e^{-Np_{w}}} \approx \frac{u v \lambda_{a} \lambda_{w} p_{m} |\Delta_{w} + \Delta_{a}|}{p_{w} \Delta_{a}^{2} \Delta_{w}^{2}}$$

$$=\frac{1}{|\Delta_w|} + \frac{1}{|\Delta_a|},\tag{C9}$$

where in the last line we used the fact that $1/p_w = N_a^*$ and Equation (4).

If a fraction f of the cancer cells are an uploid when the drug is administered then the rates at which the rescue mutations are generated can be written as:

$$r_1^f(t) = v\lambda_a p_m \int_0^t a_z dz = (1 - f) \frac{uv\lambda_w \lambda_a N p_m}{\Delta_w - \Delta_a} \left(\frac{e^{\Delta_w t} - 1}{\Delta_w} - \frac{e^{\Delta_a t} - 1}{\Delta_a} \right) + fv\lambda_a N p_m \frac{e^{\Delta_a t} - 1}{\Delta_a},$$

$$r_2^f(t) = v\lambda_w p_m \int_0^t w_z dz = (1 - f) v\lambda_w N p_m \frac{e^{\Delta_w t} - 1}{\Delta_w},$$

and the mean evolutionary rescue time is given by:

$$\tau_a = \frac{\int_0^\infty t R_a^f(t) \exp\left(-\int_0^t R_a^f(z) dz\right) dt}{p_{rescue}},$$
 (C10)

where $R_a^f(t) = r_1^f(t) + r_2^f(t)$ and $p_{rescue} = 1 - \exp\left[-\left(1 - f\right)p_wN - fp_aN\right]$. We plot our approximation in Figure 9 together with simulated data.

Appendix D Recurrence time

We define the proliferation time τ_a^r to be the time it takes the population of mutant cancer cells to reach the initial tumor size N. The number of rescue lineages generated by the wildtype population is given by eq. (C5) (see Figure S3):

$$r_1(\infty) = \frac{uv\lambda_w\lambda_a Np_m}{|\Delta_w||\Delta_a|} = \frac{N}{N_a^*},$$

where we ignore lineages created by direct mutation because we assumed $u\lambda_a > \max(-\Delta_a, 1/T^*)$, $N \ll N_m^*$ and used Equation (4).

This helps us distinguish between two cases for the proliferation time. Firstly, when we have at most one lineages which rescues the cancer cell population:

$$N \ll N_a^*$$

As a result, the recurrence time is given by (Avanzini and Antal, 2019):

$$\tau_a^r \approx \tau_a + \frac{\log p_m N}{\Delta_m}.$$
 (D1)

The factor of p_m in the second term of eq. (D1) is due to the fact that the lineage is conditioned to survive genetic drift and the time to reach N is shorter then the case without this property.

The second case is when the wildtype population produces a large number of rescue lineages in a short period of time. This is given by the condition:

$$N \gg N_a^*$$
.

As a result, the recurrence time is obtained by solving the following system of ODEs:

$$\frac{dw}{dt} = \Delta_w w,
\frac{da}{dt} = \Delta_a a + u \lambda_w w,
\frac{dm}{dt} = \Delta_m m + v \lambda_a a + v \lambda_w w.$$
(D2)

Solving the system of ODEs for initial condition (w(0), a(0), m(0)) = (N, 0, 0) we obtain:

$$m\left(t\right) = \frac{Nuv\lambda_{a}\lambda_{w}}{\Delta_{w} - \Delta_{a}}\left[\frac{\mathrm{e}^{\Delta_{w}t} - \mathrm{e}^{\Delta_{m}t}}{\Delta_{w} - \Delta_{m}} - \frac{\mathrm{e}^{\Delta_{a}t} - \mathrm{e}^{\Delta_{m}t}}{\Delta_{a} - \Delta_{m}}\right] + Nv\lambda_{w}\frac{\mathrm{e}^{\Delta_{w}t} - \mathrm{e}^{\Delta_{m}t}}{\Delta_{w} - \Delta_{m}}.$$

We obtain τ_a^r such that $m\left(\tau_a^r\right) = N$ by solving:

$$1 = \frac{uv\lambda_a\lambda_w}{\Delta_w - \Delta_a} \left[\frac{e^{\Delta_w\tau_a^r} - e^{\Delta_m\tau_a^r}}{\Delta_w - \Delta_m} - \frac{e^{\Delta_a\tau_a^r} - e^{\Delta_m\tau_a^r}}{\Delta_a - \Delta_m} \right] + v\lambda_w \frac{e^{\Delta_w\tau_a^r} - e^{\Delta_m\tau_a^r}}{\Delta_w - \Delta_m}.$$
 (D3)

This is a transcendental equation which cannot be solved exactly but we can obtain an approximation by noting that for large τ_a^r the above equation can be written as:

$$1 = v\lambda_w \frac{\mathrm{e}^{\Delta_m \tau_a^r}}{|\Delta_w - \Delta_m|},$$

which has solution:

$$\tau_a^r \approx \frac{1}{\Delta_m} \log \frac{\Delta_m - \Delta_w}{v \lambda_w}.$$
 (D4)

We observe that the terms given by the evolutionary trajectory $wildtype \rightarrow aneuploid \rightarrow mutant$ do not contribute to the above approximation and, as a result, we deduce that it accurate only for $N \gg N_m^* > N_a^*$.

Additionally, we note that if we are interested in the time until the tumor reaches a detectable size *M* then our above analysis is valid but in Equation (D1) we change:

$$\tau_a^{r,M} \approx \tau_a + \frac{\log p_m M}{\Delta_m},$$
 (D5)

and Equation (D4) becomes:

$$\tau_a^{r,M} \approx \frac{1}{\Delta_m} \log \frac{M (\Delta_m - \Delta_w)}{v \lambda_w N},$$
(D6)

which we plot in Figure 8 and observe that our approximations are in agreement with simulations.

Appendix E Distribution of evolutionary rescue time

The probability that a successful mutant has been generated by time t is given by:

$$\begin{split} P\left(rescue,t\right) &= P\left(T_{a} < t\right) \\ &= 1 - \exp\left\{-\left[r_{1}\left(t\right) + r_{2}\left(t\right)\right]\right\} \\ &= 1 - \exp\left\{-\left[\frac{uv\lambda_{w}\lambda_{a}Np_{m}}{\Delta_{w} - \Delta_{a}}\left(\frac{\mathrm{e}^{\Delta_{w}t} - 1}{\Delta_{w}} - \frac{\mathrm{e}^{\Delta_{a}t} - 1}{\Delta_{a}}\right) + v\lambda_{w}Np_{m}\frac{\mathrm{e}^{\Delta_{w}t} - 1}{\Delta_{w}}\right]\right\}, \end{split}$$

where T_a is the time at which the first mutant cell appears that will avoid extinction and which was defined in appendix C.

As a result, the probability that a successful mutant has not been generated by time t is:

$$1 - P\left(rescue, t\right) = \exp\left\{-\left[\frac{uv\lambda_w\lambda_a Np_m}{\Delta_w - \Delta_a}\left(\frac{e^{\Delta_w t} - 1}{\Delta_w} - \frac{e^{\Delta_a t} - 1}{\Delta_a}\right) + v\lambda_w Np_m \frac{e^{\Delta_w t} - 1}{\Delta_w}\right]\right\}.$$
(E1)

Appendix F Distribution of recurrence time

The probability distribution of the time that a lineage, consisting initially of a single cell, will reach size N as time t is given by the Gumbel distribution $\operatorname{Gumb}_{max}\left(\frac{\log Np_m}{\Delta_m}, \frac{1}{\Delta_m}\right)$ (Avanzini and Antal, 2019) with probability density function:

$$G(t) = e^{-p_m N e^{-\Delta_m t}}.$$

A mutant lineage initiated at time s, through an euploidy, at rate $v\lambda_a p_m a_s$ reaches size N before time t with probability G(t-s) where $s \le t$. As a result, the number of successful mutant lineages which reach size N by time t can be approximated by inhomogeneous Poisson random variable with rate:

$$r(t) = v\lambda_a p_m \int_0^t a_s G(t - s) \, ds$$

where a_s is an euploid population size at time s defined in eq. (C4). The proliferation time is defined as the first time the size of all lineages reaches N. When $N \ll |\Delta_w| |\Delta_a| / uv \lambda_w \lambda_a p_m$ there is at most a single mutant lineage that will survive and reach size N (Figure S3) and the probability that the size of that lineage has not reached N by time t is given by:

$$P(m_{t} \leq N) = \exp\left[-r(t)\right]$$

$$= \exp\left[-\frac{Nuv\lambda_{w}\lambda_{a}p_{m}}{\Delta_{w} - \Delta_{a}} \int_{0}^{t} \left[e^{\Delta_{w}s} - e^{\Delta_{a}s}\right] e^{-p_{m}Ne^{-\Delta_{m}(t-s)}} ds\right].$$
 (F1)

When $N \gg |\Delta_w| |\Delta_a| / uv \lambda_w \lambda_a p_m$ the dynamics of the cancer cell populations is deterministic and approximated by the system of ODEs shown in eq. (D2). As a result, the size of the mutant cell population will always be below N until time τ_a^r and will always be greater after:

$$P\left(m_{t} \leq N\right) = 1 - H\left(t - \tau_{a}^{r}\right),\tag{F2}$$

where H(x) is the Heaviside function:

$$H\left(x\right) = \begin{cases} 0, & x < 0, \\ 1, & x \ge 0. \end{cases}$$

We plot eq. (F1) and eq. (F2) in Figure and compare with stochastic simulations and observe that our approximation are in agreement.

We observe that for $N=10^7$ our formula overestimates the probability that the mutant population will be smaller then N at time t. This can be explained by the fact that $N=10^7$ is an intermediary case where the wildtype population produces a number of rescue lineages that is greater then one but still sufficiently small such that stochasticity plays an important role in the population dynamics. As a result, the number of mutant cancer cells will reach N faster then the case with a single mutant lineage. Additionally, we observe from Figure 6B that the probability of the mutant cell population reaching size N is approximately zero before time τ_a^r which is the recurrence time for the deterministic case. This can be explained as follows: in the deterministic case there is a sufficient number of lineages produced such that there exists a lineage where each descendant will only reproduce and not die; the time it takes for this lineage to reach N is the lower bound for the time of all other lineages to reach N and this time cannot be smaller then τ_a^r by definition. Given that for small values of N we expect that at most a single lineage will rescue the tumor, this lineage cannot reach N before τ_a^r for the deterministic case eq. (D4).

From eq. (F2) we obtain the distribution of the recurrence time conditional of evolutionary rescue:

$$f(t) = \frac{d}{dt} \left[\frac{P(m_t \ge N)}{p_{rescue}} \right] = r'(t) \frac{\exp\left[-r(t)\right]}{p_{rescue}},$$
 (F3)

which we plot in Figure and compare with simulations. We note that in the case $N \gg |\Delta_w| |\Delta_a| / uv \lambda_w \lambda_a p_m$ the distribution becomes the Dirac δ -function (Barton, 1989).

Appendix G: Bootstrapping

For the mean times the 95% confidence interval is obtained through bootstrapping in the following steps: (1) we simulate T 100 times; (2) we sample with replacement which we store in T'; (3) for each element of this sample we obtain $\tau = \mathbb{E}[T']$; (4) we repeat steps (2)-(3) 100 times to obtain τ and we select the upper and lower limits such that 95% of the values of τ lie in the interval given by the bounds.

For the threshold tumor sizes the 95% confidence interval is obtained through bootstrapping in the following steps: (1) we simulate p_{rescue} 100 times; (2) we sample with replacement which we store in S; (3) for each element of this sample we obtain $N_a^* = 1/p_w$ using $p_w = -1/N_s \log \left(1 - \bar{S}\right)$ where \bar{S} is the mean of S and N_s is an arbitrary value of the initial population size we selected in order to calculate p_{rescue} ; (4) we repeat steps (2)-(3) 100 times to obtain N_a^* and we select the upper and lower limits such that 95% of the values of N_a^* lie in the interval given by the bounds.

For the ratio of the threshold tumor sizes the 95% confidence interval is obtained through bootstrapping in the following steps: (1) we simulate p_{rescue} 100 times for both the case when $f = \tilde{u}\lambda_w/s$ and f = 0; (2) we sample with replacement which we store in S_0 and S_f ; (4) for each element of S_0 we obtain $N_a^* = 1/p_w$ using $p_w = -1/N_s \log\left(1 - \bar{S}\right)$ where \bar{S} is the mean of S_0 and S_0 and S_0 is an arbitrary value of the initial population size we selected in order to calculate p_{rescue} ; (5) for each element of S_f we obtain $\tilde{N}_a^* = 1/p_a$ using $p_a = -f/N_s \log\left(1 - \bar{S}\right)$ where \bar{S}_f is the mean of S_f and S_f is an arbitrary value of the initial population size we selected in order to calculate S_0 we repeat steps (2)-(5) 100 times to obtain \tilde{N}_a^*/N_a^* and we select the upper and lower limits such that 95% of the values of \tilde{N}_a^*/N_a^* lie in the interval given by the bounds.

Supplementary Figures

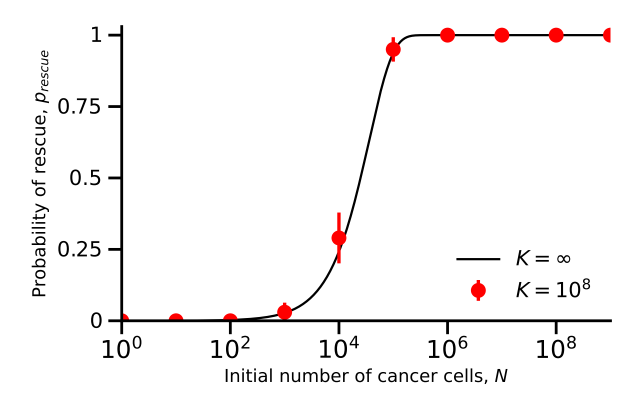


Figure S1: Density dependent growth does not affect the accuracy of our model. Comparison of results of simulations with density-dependent growth (red markers with with 95% CI) and the approximation formula (black line, eq. (4) in eq. (2)) with maximum carrying capacity $K = 10^8$ and effective carrying capacity $K_e = K\Delta_a/\lambda_a \approx 10^6$. The error bars represent 95% confidence interval of the form $p \pm 1.96\sqrt{p(1-p)/n}$ where p is the fraction of simulations in which the tumor has adapted to the stress and n = 100 is the number of simulations. Parameters: $\lambda_w = 0.1$, $\lambda_a = 0.0901$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\mu_m = 10^{-2}$, $\nu = 10^{-7}$, $\nu = 10^{-8}$.

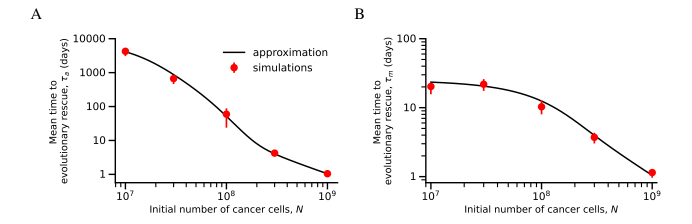


Figure S2: Evolutionary rescue time. Shown is the mean time for appearance of a resistance mutation the leads to evolutionary rescue (A) with aneuploidy (u > 0) and (B) without aneuploidy (u = 0). Our inhomogeneous Poisson-process approximations (solid black lines, right: eq. (C2), left: eq. (C7)) is in agreement with simulation results (red markers with 95% quantile intervals obtained with bootstrapping, see see Appendix G). Parameters: $\lambda_w = 0.1$, $\lambda_m = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\mu_m = 0.09$, $\mu_m = 0.09$.

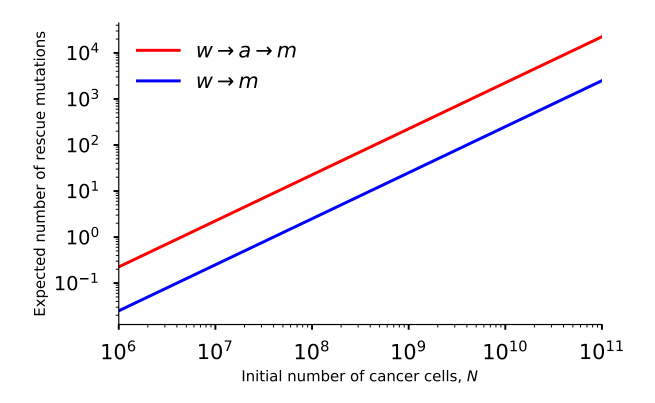


Figure S3: Aneuploidy increases the number of mutations which rescue the tumor. Shown is the expected number of mutation, which will rescue the cancer cell population, produced through the evolutionary trajectory wildtype \rightarrow mutant (blue line, eq. (C6)) or through the trajectory wildtype \rightarrow aneuploid \rightarrow mutant (red line, eq. (C5)). Parameters: $\lambda_w = 0.1$, $\lambda_m = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\mu = 10^{-2}$, $\nu = 10^{-7}$.



Figure S4: Distribution of the recurrence time. Shown is the distribution of the time for the mutant cell population to reach size N, where N is the initial number of cancer cells. The red line is analytic result eq. (F3) overlaid over the histogram of simulations. Parameters: $N = 10^6$, $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\mu_m = 0.09$, $\mu_m = 0.09$.

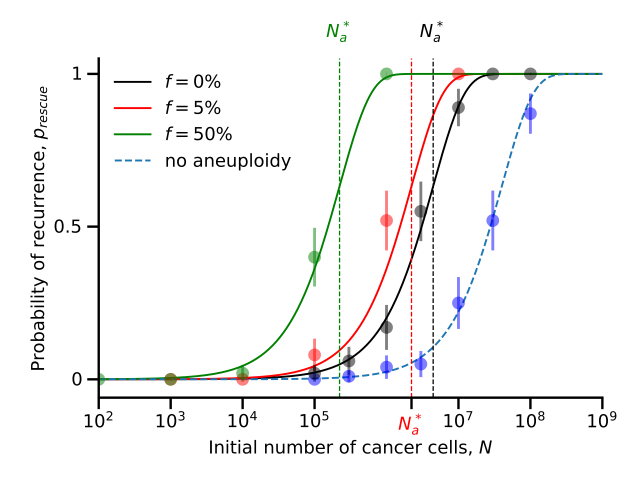


Figure S5: The probability of evolutionary rescue (i.e. the probability that the population does not go to extinction), p_{rescue} , as a function of the initial tumor size, N. Dashed vertical line shows the threshold tumor size, above which the probability is very high. Blue dashed line represents the probability of evolutionary rescue as a function of N without aneuploidy (u = 0). The black line represents the case where a fraction f = 0% of the initial tumor is aneuploid, the red line represents the case with f = 5% and the green line represents the case with f = 50%. The dots represent simulations and the error bars represent 95% confidence interval of the form $p \pm 1.96\sqrt{p(1-p)/n}$ where p is the fraction of simulations in which the tumor has adapted to the stress and n = 100 is the number of simulations. Parameters: $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\mu_a = 0.09$,

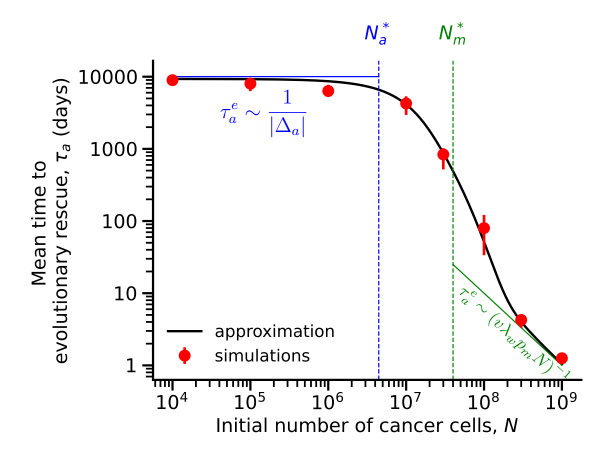


Figure 6: Shown is the mean time for appearance of a resistance mutation the leads to evolutionary rescue with aneuploidy (u > 0). Our inhomogeneous Poisson-process approximations (solid black lines, right: eq. (C7)) is in agreement with simulation results (red markers with 95% confidence intervals obtained with bootstrapping, see Appendix G). Dashed vertical blue line represents the threshold tumor size above which evolutionary rescue is very likely through aneuploidy eq. (4) and the dashed vertical green line represents the threshold tumor size above which evolutionary rescue is very likely through direct mutation eq. (3). Solid lines represents the approximations eq. (8) ($N < N_a^*$ blue line and $N > N_m^*$ green line). Parameters: $\lambda_w = 0.1$, $\lambda_m = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$.

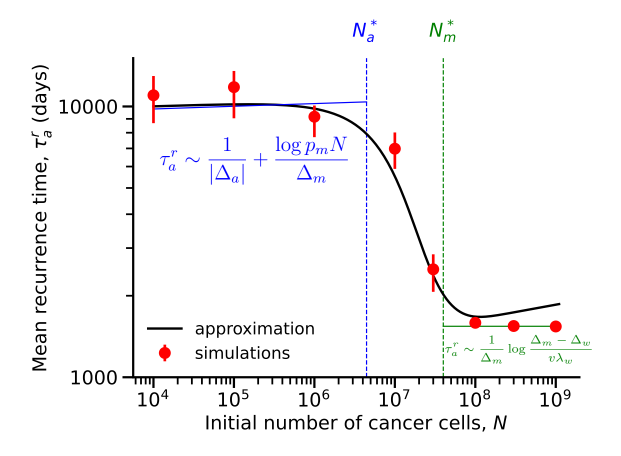


Figure 7: The mean time for the mutant cell population to reach size N, where N is the initial number of cancer cells. Our inhomogeneous Poisson-process approximation (solid black line, eq. (D1)) is in agreement with simulation results (red markers with 95% confidence intervals obtained with bootstrapping, see Appendix G) for small and intermediate values of N. Dashed vertical blue line represents the threshold tumor size above which evolutionary rescue is very likely through aneuploidy eq. (4) and the dashed vertical green line represents the threshold tumor size above which evolutionary rescue is very likely through direct mutation eq. (3). Solid lines represents the approximations eq. (9) $(N < N_a^*)$ blue line and $N > N_m^*$ green line). The simulations converge to eq. (D4) (green line) for large values of $N \gg N_m^*$. Parameters: $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$

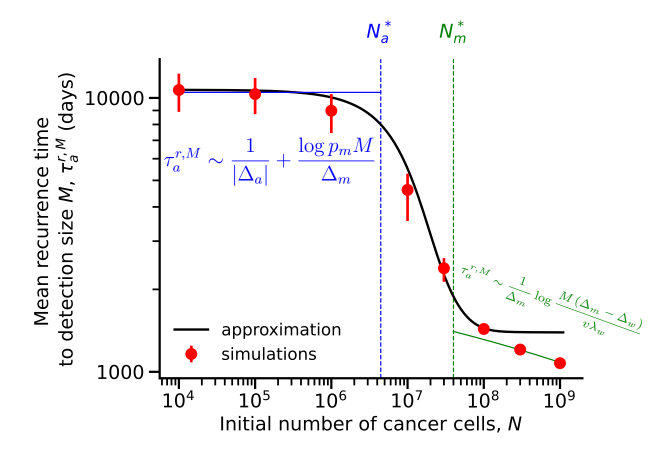


Figure 8: The mean time for the mutant cell population to reach size M, where M is the tumor detection size. Our inhomogeneous Poisson-process approximation (solid black line, eq. (D5)) is in agreement with simulation results (red markers with 95% confidence intervals obtained with bootstrapping, see Appendix G) for small and intermediate values of N. Dashed vertical blue line represents the threshold tumor size above which evolutionary rescue is very likely through aneuploidy eq. (4) and the dashed vertical green line represents the threshold tumor size above which evolutionary rescue is very likely through direct mutation eq. (3). Solid blue line represents the approximation eq. (D5) with τ_a from eq. (8) for $N < N_a^*$ and the solid green line represents the approximation eq. (D6) for $N > N_m^*$. The simulations converge to eq. (D6) (green line) for large values of $N \gg N_m^*$. Parameters: $\lambda_w = 0.1$, $\lambda_a = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\mu_m = 10^{-2}$, $\nu = 10^{-7}$.

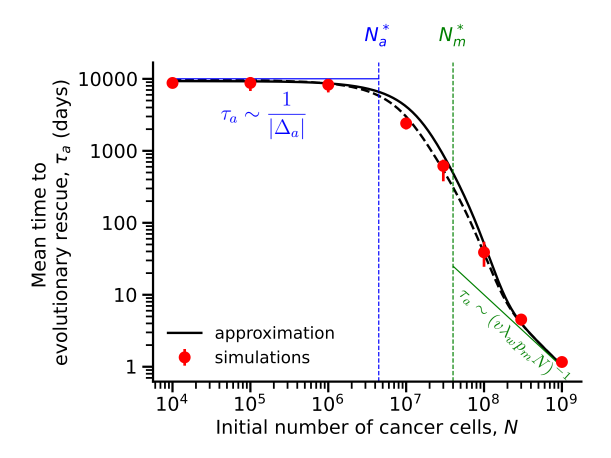


Figure 9: Shown is the mean time for appearance of a resistance mutation the leads to evolutionary rescue with aneuploidy (u > 0). Black lines represent our inhomogeneous Poisson-process approximations (solid black line, eq. (C7); dashed black line eq. (C10)). Dashed black line is the inhomogeneous Poisson-process approximation where a fraction f of tumor is aneuploid at the onset of drug therapy which is in agreement with simulation results (red markers with 95% confidence intervals obtained with bootstrapping, see Appendix G). Dashed vertical blue line represents the threshold tumor size above which evolutionary rescue is very likely through aneuploidy eq. (4) and the dashed vertical green line represents the threshold tumor size above which evolutionary rescue is very likely through direct mutation eq. (3). Solid lines represents the approximations eq. (8) $(N < N_a^*)$ blue line and $N > N_m^*$ green line). Parameters: $\lambda_w = 0.1$, $\lambda_m = 0.0899$, $\lambda_m = 0.1$, $\mu_w = 0.14$, $\mu_a = 0.09$, $\mu_m = 0.09$, $\mu_m = 0.09$, $\mu_m = 0.07$, $\mu_m = 0.07$, $\mu_m = 0.0899$, $\mu_m = 0.0899$, $\mu_m = 0.09$, $\mu_m =$