

CRITICAL BRANCHING RANDOM WALK IN AN IID ENVIRONMENT

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ABSTRACT. Using a high performance computer cluster, we run simulations regarding an open problem about d -dimensional critical branching random walks in a random IID environment. The environment is given by the rule that at every site independently, with probability $p \in [0, 1]$, there is a *cookie*, completely suppressing the branching of any particle located there.

The simulations suggest *self averaging*: the asymptotic survival probability in n steps is the same in the annealed and the quenched case; it is $\frac{2}{qn}$, where $q := 1 - p$. This particular asymptotics indicates a non-trivial phenomenon: the tail of the survival probability (both in the annealed and the quenched case) is the same as in the case of non-spatial unit time critical branching, where the branching rule is modified: branching only takes place with probability q for every particle at every iteration.

CONTENTS

1. Introduction	1
1.1. Model	1
1.2. The problems we investigate	2
1.3. Motivation	2
2. Preliminaries	5
3. Implementation	6
3.1. Annealed simulation	7
3.2. Quenched simulation	8
4. Simulation results	9
4.1. Annealed simulation on \mathbb{Z}^2	9
4.2. Annealed simulation on \mathbb{Z}^1	10
4.3. Quenched simulation	12
5. Interpretation of the simulation results	13
5.1. Main finding	13
5.2. Interpretation of the fluctuations in the diagrams	13
6. Beyond the first order asymptotics	15
6.1. Two dimensions	16
6.2. One dimension	16
6.3. Comparison between one and two dimensions	16
7. Conclusion	17
References	17

1. INTRODUCTION

1.1. **Model.** We consider a model of a critical branching random walk in an IID random environment on the d -dimensional integer lattice as follows. The environment is determined by placing *cookies* on each site, with probability $0 \leq p \leq 1$, independently of each other.

Date: May 23, 2011.

2000 *Mathematics Subject Classification.* Primary: 60J80; Secondary: 60K37 .

Key words and phrases. Branching random walk, catalytic branching, mild obstacles, critical branching, random environment, simulation.

The authors thank S. Kuznetsov and R. St. Laurent for helpful conversations.

Given an environment, the initial particle, located at the origin, first moves according to a nearest neighbor simple random walk, and immediately afterwards, the following happens to her:

- (1) If there is no cookie at the new location, the particle either vanishes or splits into two offspring particles, with equal probabilities.
- (2) If there is a cookie at the new location, nothing happens to the particle.

The new generation then follows the same rule in the next unit time interval and produces the third generation, etc.

Let $p \in [0, 1]$. In the sequel \mathbb{P}_p will denote the law of the cookies and P^ω will denote the law of the BRW given the environment ω . So, if \mathbf{P}_p denotes the ‘mixed’ law in the environment with cookie probability p , we have

$$\mathbf{P}_p(\cdot) = \mathbb{E}_p P^\omega(\cdot).$$

Following standard terminology, P^ω and \mathbf{P}_p will be called *quenched* and *annealed* probabilities, respectively.

1.2. The problems we investigate. We are interested in the asymptotic behavior, as time tends to infinity, of the probability that there are surviving particles, and on its dependence on the parameters d and p . (Note that in the two extreme cases, when $p = 0$ and $p = 1$, the asymptotics is well known.) In particular, we are interested to know whether we can find some qualitative difference between the annealed and the quenched asymptotics, or they are the same (in which case one can talk about *self-averaging*). In the quenched case, one may, of course, only talk about the *almost sure* asymptotics, as the probability P^ω itself depends on the realization of the environment.

Furthermore, we are interested in the simulation aspects as well. We will see, that the efficiency will depend on the size of d and p , as well as on whether we are in the annealed or the quenched setting.

1.3. Motivation.

1.3.1. Motivation within mathematics. In [5] a spatial branching model has been studied, where the underlying motion is a d -dimensional ($d \geq 1$) Brownian motion, the particles perform dyadic branching, and the branching rate is affected by a random collection of reproduction suppressing sets dubbed *mild obstacles*. In fact the obstacle configuration was given by the union of balls with fixed radius, where the centers of the balls form a Poisson point process. The radius r plays no role in the results, but the Poisson intensity $\nu > 0$ does. The main result of [5] is the quenched Law of Large Numbers for the population for all $d \geq 1$. The environment ω (with law P^ω) has the property that the branching rate is β_1 inside the obstacles and β_2 outside of them, with $0 < \beta_1 < \beta_2$. The branching process given the environment ω was denoted by $Z(\omega)$ and the total population size at $t \geq 0$ was denoted by $|Z_t(\omega)|$. Define the average growth rate by $r_t = r_t(\omega) := \frac{\log |Z_t(\omega)|}{t}$. In [5] it was shown that for almost every environment,

$$\lim_{t \rightarrow \infty} (\log t)^{2/d} (r_t - \beta_2) = -c(d, \nu) \quad P^\omega\text{-probability,}$$

where $c(d, \nu) > 0$ is an explicit constant. (This is a kind of LLN, because the expectation of the total population size obeys the same asymptotics.)

It has also been shown that the branching Brownian motion with mild obstacles spreads less quickly than ordinary branching Brownian motion, and an upper estimate for its radial speed has been provided.

More general offspring distributions (beyond the dyadic one considered in the main theorems) were also discussed in [5]. In particular, the following question was posed. Consider the model where the offspring distribution is *critical*. One can easily prove (see Theorem 2.2 below) that, despite of the obstacles, the system still dies out with probability one.

Problem 1.1. What is the rate of decay for the survival probability? Is it still of order C/n as in the obstacle-free (non-spatial) case?

In the present paper we are going to investigate this problem in our discretized setting, where Brownian motion is replaced by branching random walk, and Poissonian obstacles are replaced by IID cookies on lattice points.

Remark 1.2. An alternative view on our setting is as follows. Let K denote the (random) set of those lattice points where there is a cookie. Then, our model can be viewed as a *catalytic* BRW as well — the catalytic set is then K^c (in the sense that branching is ‘made possible’ there). Catalytic spatial branching (mostly for superprocesses though) has been the subject of vigorous research in the last twenty years initiated by Dawson, Fleischmann and others — see the survey papers [10] and [4] and references therein. In those models the individual branching rates of particles moving in space depend on the amount of contact between the particle (‘reactant’) and a certain random medium called the catalyst. The random medium is usually assumed to be a ‘thin’ random set (that could even be just one point) or another superprocess. Sometimes ‘mutually’ or even ‘cyclically’ catalytic branching is considered [4].

Our model is simpler than most catalytic models as our catalytic/blocking areas are fixed, whereas in several catalytic models they are moving. On the other hand, while for catalytic settings studied so far, results were mostly only qualitative, we are aiming to get quite sharp *quantitative* results.

Another model of a critical spatial branching process was recently introduced and investigated by Véber in [16]. In that paper the author proves a convergence theorem for a sequence of super-Brownian motions moving among hard Poissonian obstacles, when the intensity of the obstacles grows to infinity but their diameters shrink to zero in an appropriate manner. The superprocesses are shown to converge in probability for the law \mathbf{P} of the obstacles, and \mathbf{P} -almost surely for a subsequence, towards a superprocess with underlying spatial motion given by Brownian motion and (inhomogeneous) branching mechanism $\psi(u, x)$ of the form $\psi(u, x) = u^2 + \kappa(x)u$, where $\kappa(x)$ depends on the density of the obstacles. This work draws on similar questions for a single Brownian motion.

A very recent manuscript by LeGall and Véber [6] derives asymptotics for the quenched probability that a critical branching Brownian motion killed at a small rate in Poissonian obstacles exits a large domain. Results are formulated in terms of the solution to a semilinear partial differential equation with singular boundary conditions. The proofs depend on a quenched homogenization theorem for branching Brownian motion among mild obstacles.

For the discrete setting there are much fewer results available. We give two examples here.¹ The first one is [9], where the branching particle system on \mathbb{Z}^d is so that its branching is catalyzed by another autonomous particle system on \mathbb{Z}^d . There are two types of particles, the A -particles (‘catalyst’) and the B -particles (‘reactant’). They move, branch and interact in the following way. Let $N_A(x, s)$ and $N_B(x, s)$ denote the number of A - (resp. B -) particles at $x \in \mathbb{Z}^d$ and at time $s \in [0, \infty)$. (All $N_A(x, 0)$ and $N_B(x, 0)$, $x \in \mathbb{Z}^d$ are independent Poisson variables with mean μ_A (μ_B).) Every A -particle (B -particle) performs independently a continuous-time random walk with jump rate D_A (D_B). In addition a B -particle dies at rate δ , and, when present at x at time s , it splits into two in the next ds time with probability $\beta N_A(x, s)ds + o(ds)$. Conditionally on the system of the A -particles, the jumps, deaths and splitting of the B -particles are independent. For large β the existence of a critical δ , separating local extinction regime from local survival regime, is shown.

The second example is [15]. In that paper a continuous time one-dimensional critical branching random walk on the lattice \mathbb{Z} is considered in which individuals may produce children at the origin only. Assuming that the underlying Markov random walk is homogeneous and symmetric, the authors describe the asymptotic behavior as $t \rightarrow \infty$ of the conditional distribution of the joint distribution of $\zeta(t)$ and $\mu(t)$ (scaled in an appropriate way), where $\zeta(t)$ and $\mu(t)$ are the numbers of individuals at the origin and outside the origin at moment t given $\zeta(t) > 0$.

¹A further example of the discrete setting is [1].

Remark 1.3. Finally, note that while the continuous equivalent of an IID trap configuration on the lattice is a Poisson trap configuration on \mathbb{R}^d , there is an important difference between the two. The discrete setting has the advantage that the difference between the sets K and K^c is no longer relevant. Indeed, in the discrete case the complement is also IID with a different parameter (self-duality), whereas in the continuous setting this nice duality is lost as the ‘Swiss cheese’ K^c is not the same type of geometric object as K ; the latter is the case, for example, in [5].

1.3.2. *Motivation in biological population models.* Spatial branching processes turn out to be very important processes for modeling purposes. They provide good models for studying the development of genetic diversity over a large number of generations — indeed, it was as models of genetic diversity that superprocesses (scaling limits of branching random walks) originally developed. Moreover, they are very useful when one studies for example

- the dispersion of mass via ocean currents;
- phenomena concerning ocean pollutants;
- the spatial structure of populations of small organisms living in the ocean, e.g. plankton.

In general, stochastic spatial models are very important from the point of view of applications in *theoretical biology* and *ecology*: they are used to study questions involving the pattern of fluctuations in density caused by demographic fluctuations in a spatially migrating population, predator-prey interactions, species coexistence and competition, epidemic and genetic diversity. See for example the recent monograph [3] which gives a very mathematical treatment of spatial ecology, though it treats the subject exclusively (except for a short exposition of certain interacting particle systems and their hydrodynamical limits) from a PDE/Dynamical Systems point of view.

Spatial branching processes have been used as models of the evolution of biological populations for several decades. In those models, the location of an individual can be interpreted in two different ways: it can be a spatial position in \mathbb{R}^d , or it can be her *genetic type*. (Working in \mathbb{R}^d means that each genetic type can be uniquely described by d quantitative characteristic properties.) Therefore, in the genetic setup, the ‘spatial motion’ describes *mutation* between types.

For example, a classical model of genetics is the *Wright-Fisher model*. In this model the number of individuals with a given gene evolves in time and undergo biological mutation, as well as Darwinian *selection*, leading to a Markov chain with a state space that has the size of the population. Then the method of diffusion approximation allows one to approximate the Markov chain by a continuous Markov process and use standard machinery from the theory of differential equations to analyze those processes.

It should be emphasized however, that in theoretical biology and ecology, probabilistic investigation is often more beneficial than an analysis based solely on differential equations. One of the reasons is that the genealogical structure may be of great significance even after passing to the diffusion approximation.

The first author has made some efforts to connect his mathematical work with biological models; talks have been initiated with Dr. R. Nisbet of the Ecology, Evolution and Marine Biology department, UC Santa Barbara and with C. Cosner, an applied mathematician at the University of Miami, whose research field is mathematical biology. C. Cosner pointed out that he had conversations with his collaborator, B. Fagan (an ecologist at the Department of Biology of the University of Maryland) about a phenomenon in marine systems where hypoxic patches form in estuaries because of stratification of the water. The patches affect different organisms in different ways but are detrimental to some of them. They appear and disappear in an effectively stochastic way². This is an actual system that has some features that correspond to the type of assumptions built into the models with mild obstacles.

²It might actually be deterministic but chaotic, but that is still a situation where a stochastic model is reasonable.

Correspondence has also been initiated with Dr. Fagan, whose work involves population dynamics with stochastic components. It appears³ that a very relevant existing ecological context in which to place the models with mild obstacles is the so-called “source-sink theory”. The basic idea is that some patches of habitat are good for a species (and growth rate is positive) whereas other patches are poor (and growth rate is zero or negative). Individuals can move between patches randomly or according to more detailed biological rules for behavior. (See [13] as the main reference; see also [8] and [2].)

C. Cosner has also pointed out another kind of scenario where models with mild obstacles would make sense: systems that are subject to periodic local disturbances. Those would include forests where trees sometimes fall creating gaps (which have various effects on different species but may harm some) or areas of grass or brush which are subject to occasional fires (becoming increasingly popular in Southern California and Southern Europe.) Again, the effects may be mixed, but the burned areas can be expected to be less suitable habitats for at least some organisms.

Finally, it appears that models with mild obstacles have biological merit, especially in the area of *biological invasions* before density dependence would occur⁴.

2. PRELIMINARIES

In this section we present two simple statements concerning our branching random walk model which are relatively easy to verify rigorously. Let S_n denote the event of *survival* for $n \geq 0$. That is, $S_n = \{Z_n \geq 1\}$, where Z_n is the population size at time n .

Theorem 2.1 (Monotonicity). *Let $0 \leq p < \hat{p} \leq 1$ and fix $n \geq 0$. Then*

$$\mathbf{P}_p(S_n) \leq \mathbf{P}_{\hat{p}}(S_n).$$

Proof. First notice that it suffices to prove the following statement:

Assume that we are given an environment with some ‘red’ cookies and some additional ‘blue’ cookies. Then the probability of S_n with the additional cookies is larger than or equal to the probability without them.

Indeed, one can argue by coupling as follows. Let $q := 1 - p$, $\delta := \hat{p} - p$. First let us consider the cookies that are coming with IID probabilities and parameter p . These will be the ‘red’ cookies. Now with probability δ/q at each site independently, add a blue cookie. Then the probability for any given site, that there is at least one cookie there is $p + \delta/q - p\delta/q = p + \delta = \hat{p}$. Now delete those blue cookies where there was a red cookie too. This way, the red cookies plus the additional blue cookies together correspond to parameter \hat{p} .

We are thus going to prove the statement in italics now, using an argument due to S. Kuznetsov. Consider the generating functions of no branching and critical branching: $\varphi_1(z) = z$ and $\varphi_2(z) = \frac{1}{2}(1 + z^2)$, and note that $\varphi_1 \leq \varphi_2$ on \mathbb{R} . Fix an environment and define

$$u(n, x, N) := P_{n,x}(S_N^c),$$

that is, the probability that the population emanating from a single particle which is at time n is located in x , becomes extinct at time N . Then, if the particle moves to the random location ξ_{n+1} , one has

$$\begin{aligned} u(n, x, N) &= E \sum_{i=0}^2 p_i(\xi_{n+1}) [P_{n+1, \xi_{n+1}}(S_N^c)]^i \\ &= E \sum_{i=0}^2 p_i(\xi_{n+1}) [u(n+1, \xi_{n+1}, N)]^i = E \varphi^{\xi_{n+1}}[u(n+1, \xi_{n+1}, N)], \end{aligned}$$

³Communication with Dr. Fagan.

⁴Communication with P. Chesson of the Section of Evolution and Ecology of the University of California at Davis.

where $p_i(\xi_{n+1})$ is the probability⁵ of producing i offspring at the location ξ_{n+1} , and $\varphi^{\xi_{n+1}}$ is either φ_1 or φ_2 .

Now consider two environments: one with only red cookies and another one where there are also some additional blue cookies, and let us denote the corresponding functions by u_1 and u_2 . We have

$$u_1(n, x, N) = E\varphi_1^{\xi_{n+1}}[u_1(n+1, \xi_{n+1}, N)]$$

and

$$u_2(n, x, N) = E\varphi_2^{\xi_{n+1}}[u_2(n+1, \xi_{n+1}, N)].$$

Clearly $u_1(N, x, N) = u_2(N, x, N) = 0$. Therefore, using that $\varphi_1 \leq \varphi_2$ and by (backward) induction, $u_2 \geq u_1$ for all $n = 0, 1, \dots, N-1$. In particular,

$$u_1(0, x, N) \leq u_2(0, x, N),$$

finishing the proof. \square

We now give a precise statement and a rigorous proof concerning the result about eventual extinction mentioned briefly above.

Theorem 2.2 (Extinction). *Let $0 \leq p < 1$ and let A denote the event that the population survives forever. Then, for \mathbb{P}_p -almost every environment, $P^\omega(A) = 0$.*

Proof. Let again Z_n denote the total population size at time n for $n \geq 1$. Then Z is a martingale with respect to the canonical filtration $\{\mathcal{F}_n; n \geq 1\}$. To see this, note that just like in the $p = 0$ case, $E(Z_{n+1} - Z_n | \mathcal{F}_n) = 0$, as the particles that do not branch (due to the presence of cookies) do not contribute to the increment. Being a nonnegative martingale, Z converges a.s. to a limit Z_∞ , and of course Z_∞ is nonnegative integer valued. We now show that for \mathbb{P}_p -almost every environment, $P^\omega(Z_\infty = 0) = 1$. Introduce the events

- $A_k := \{Z_\infty = k\}$ for $k \geq 1$,
- B : branching occurs at infinitely many times $0 < \sigma_1 < \sigma_2 < \dots$

Clearly, $A = \cup_{k \geq 1} A_k = \{Z_\infty \geq 1\}$. We first show that

$$(2.1) \quad \text{for } \mathbb{P}_p \text{ - a.e. environment, } P^\omega(B^c A) = 0.$$

Clearly, it is enough to show that $\mathbf{P}_p(B^c A) = 0$.

Now, $B^c A \subset C$, where C denotes the event that there exists a first time N such that for $n \geq N$, there is no branching and particles survive and stay in the region of cookies. On C , one can pick randomly a particle starting at N , and follow her path; this path visits infinitely many points P^ω -a.s. whatever ω is⁶. Since this path is independent of ω and $p < 1$, the \mathbb{P}_p -probability that it contains a cookie at each of its sites is zero. Hence $\mathbf{P}_p(C) = 0$, and (2.1) follows.

On the other hand, for each $k \geq 1$, there is a $p_k < 1$, such that the probability that the population size remains unchanged (it remains k) at σ_m is not more than p_k for every $m \geq 1$, uniformly in ω . Thus,

$$P^\omega(BA_k) = P^\omega(B \cap \{Z_{\sigma_m} = k \text{ for all large enough } m\}) = 0,$$

whatever ω is. Using this along with (2.1), we have that for \mathbb{P}_p -almost every ω , $P^\omega(A_k) = P^\omega(B^c A_k) + P^\omega(BA_k) = 0$, $k \geq 1$, and so $P^\omega(A) = 0$. \square

3. IMPLEMENTATION

The code for our simulations are written in C++ using the MPQueue parallel library [12]. We ran the code on 96 cores using a computing cluster containing Quad-Core AMD Opteron(tm) 2350 CPU's. We used an implementation [17] of the Mersenne Twister [11] to generate random numbers. The total running time for the simulations was several months.

⁵So either $p_1 = 1$ or $p_0 = p_2 = 1/2$, according to whether there is no cookie there or there is one.

⁶Because for every ω , it is true P^ω -a.s., that every particle that does not branch, has a path that visits infinitely many points

Algorithm 1 The annealed simulation function of the code.

```

1  typedef struct {
2     Tcell cell;           // location
3     int iter;           // number of iterations
4 } Tparticle;
5  typedef vector < Tparticle > Tparticles;
6
7  void simulate (
8     int dim,           // dimension
9     double p,         // cookie probability
10    int maxiter,       // maximum number of iteration steps
11    int &iter         // longest survival
12 ) {
13    board.clear();     // remove cookies
14    iter = 0;          // iteration counter
15    Tparticles particles; // all the particles
16    particles.reserve (maxiter/10); // reserve room
17    Tparticle particle;
18    particle.cell=Tcell(dim, 0); // cell at origin
19    particle.iter = 0;
20    particles.push_back (particle); // particle at center
21    while (!particles.empty () && iter < maxiter) { // alive particles
22        int coord = mtr.randInt (dim - 1); // random coordinate
23        int dir = 2 * mtr.randInt (1) - 1; // -1 or +1
24        particles.back ().cell[coord] += dir; // move
25        if (cookie (p, particles.back ().cell)) { // there is a cookie
26            particles.back ().iter++; // survived iteration
27            iter = max (iter , particles.back ().iter); // longest survival
28        }
29        else // no cookie
30            if (mtr2.randInt (1)) { // probability 0.5
31                particles.back ().iter++; // survived iteration
32                iter = max (iter , particles.back ().iter); // longest survival
33                particles.push_back (particles.back ()); // duplicate particle
34            }
35        else // die
36            particles.pop_back (); // remove the particle
37    }
38 }

```

3.1. Annealed simulation. Algorithm 1 shows the C++ function that runs a single annealed simulation. We are essentially implementing a ‘depth-first search.’ Below we give a detailed description of the code.

- line 1: We define a data type to store particles.
- line 2: The location of the particle is stored in the *cell* field, that is a vector with the appropriate dimensions.
- line 3: The *iter* field stores the number of iterations survived by the particle.
- line 5: We define a data type to store all the living particles.
- line 7: The simulation function takes three input variables and one output variable.
- line 8: The dimension of the space is the first input.
- line 9: The probability of a cookie at any given location is the second input.
- line 10: The maximum number of allowed iterations is the third input.
- line 11: The output of the function is the maximum number of iterations any particle survived.
- line 13: We erase all the cookies from the board. Every run of the simulation uses a new cookie placement.
- line 14: The initial value of the output must be zero.
- line 15: We define a variable to store all our alive particles.

- line 16: We reserve some space to store the particles. Making the reserved space too small results in unnecessary reallocation of the variable which degrades performance. On the other hand, reserving too much space can be a problem too since different CPU's compete with each other for RAM.
- lines 18–19: The initial particle starts at the origin before the iterations start.
- line 20: At the beginning we only have the initial particle.
- line 21: We run the simulation while we have alive particles and none of them stayed alive for the maximum allowed number of iterations.
- lines 22–23: We generate a random direction.
- line 24: We move the last of our alive particles in this random direction.
- line 25: We call the cookie function to check if there is a cookie at the new location of the particle. The cookie function checks in the global variable *board* if any particle already visited this location and as a result we know already whether there is a cookie there. If no particle visited this location before, then the function uses the cookie probability to decide whether to place a cookie there or leave the location empty. This information is then stored for future visitors.
- line 26: If there is a cookie at the new location, then the particle survived one more iteration, so we increment the *iter* variable.
- line 27: It is possible that this is the longest surviving particle so far, so we update the output variable.
- line 29: If there is no cookie at the new location, then the particle splits or dies.
- line 30: We generate a random number to decide what happens.
- lines 31–32: If the particle splits, then it survives so we update information about the number of iterations.
- line 33: The particle splits, so we place a copy of it into our collection of particles as the last particle.
- lines 35–36: If the particle dies, then we remove it from our collection of particles.

The rest of our code takes care of the parallelization, data collection and the calculation of survival probabilities. The program splits the available nodes into a boss node and several worker nodes. The boss assigns simulation jobs to the workers. The workers call the simulation function several times. The boss node collects the results of these jobs and calculates the survival probabilities using all the available simulation runs. More precisely, $\mathbf{P}_p(S_n)$ is estimated as the number of simulation runs with longest survival value not smaller than n divided by the total number of runs.

3.2. Quenched simulation. The code for quenched simulation is essentially the same with only minor modifications. In this version, line 13 of the simulation function is missing, since we do not want to replace the board at every simulation.

The other change in the simulation function is at line 25. In the annealed case, every worker node has a local version of the board and the cookie function can create the board on the fly. In the quenched case, the worker nodes need to use the same board, so the cookie function cannot generate the board locally. The new version of the cookie function still stores information about the already visited locations. On the other hand, if a location is not visited yet, then the worker node asks the boss node whether this new location has a cookie. The boss node first checks whether the location was visited by any other particle at any other worker node. If the location was visited, then the boss already has a record of this location. Otherwise, the boss node uses the cookie probability to decide whether the location should have a cookie. Essentially, the boss node has the ultimate information about the board, but the worker nodes keep partial versions of the board and only consult the boss node when it is necessary.

Remark 3.1. In the quenched case, note that if $\hat{\rho}_n$ denotes the relative frequency of survivals (up to n) after r runs for a fixed environment ω , that is,

$$\hat{\rho}_n = \tilde{\rho}_n^\omega := \frac{|\text{survivals}|}{r},$$

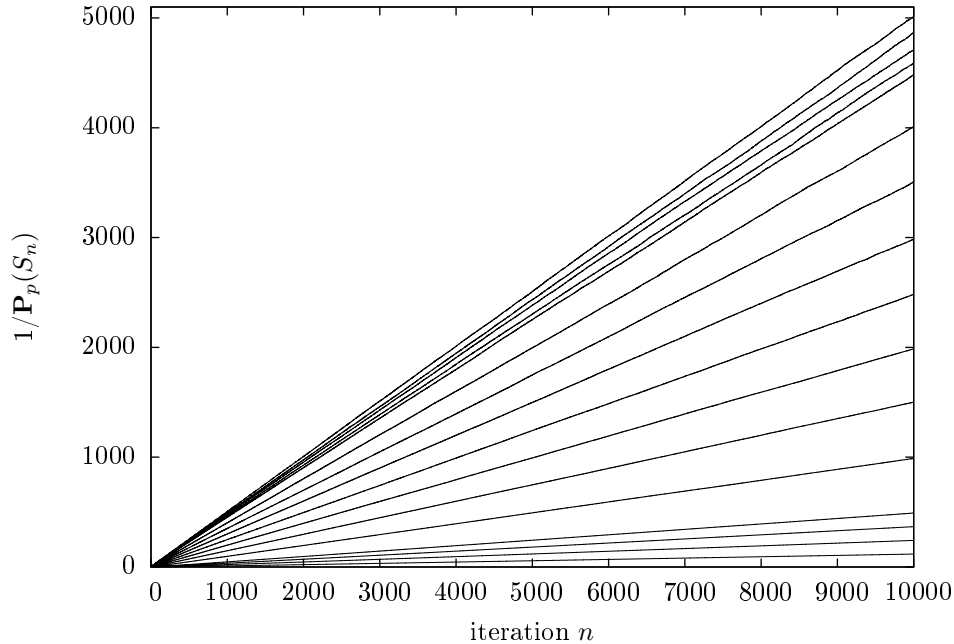


FIGURE 4.1. Results for an annealed 2-dimensional simulation. The graph shows the reciprocal of the survival probability as a function of the number of iterations. Each line represents a different cookie probability. One such line is the result of 10^8 runs of the simulation with a newly generated cookie landscape.

then using our method of simulation, the random variables $\hat{\rho}_n$ and $\hat{\rho}_m$ are not independent for $n \neq m$, because the data are coming from the same r runs.

Similarly, in the annealed case, for a fixed environment and a fixed run, the random variables $\mathbf{1}_{S_n}$ and $\mathbf{1}_{S_m}$ are not independent for $n \neq m$.

4. SIMULATION RESULTS

We ran our annealed simulation on \mathbb{Z}^d with $d \in \{1, 2, 3\}$. The 1-dimensional case turned out to be the most challenging. So we start the description of our results with the 2-dimensional case. The 3-dimensional case produced essentially the same output as the 2-dimensional case.

4.1. Annealed simulation on \mathbb{Z}^2 . We executed 10^8 runs allowing a maximum of $n_{\max} = 10000$ iterations with $p \in \{0, 0.025, 0.05, 0.075, 0.1, 0.2, \dots, 0.9, 0.925, 0.95, 0.975\}$. For $p = 1$ we used the known survival probability of 1. Preliminary results made it clear that the simulation is more sensitive for small and large values of p . This is why we picked more of these values instead of a uniformly placed set of values. The reciprocal of the calculated survival probabilities are shown in Figure 4.1. The figure suggests that $n \mapsto 1/\mathbf{P}_p(S_n)$ is asymptotically linear. We calculated the slopes for these curves from the values at $4n_{\max}/5$ and n_{\max} . These slope values are shown in Figure 4.2.

To verify the correctness of our simulation we computed the exact theoretical survival probabilities after the first two iterations. It is easy to see that $\mathbf{P}_p(S_1) = 1/2 + p/2$ and

$$\mathbf{P}_p(S_2) = 3/8 + 11p/32 + 3p^2/16 + 3p^3/32.$$

Table 1 compares some of the exact and simulated values.

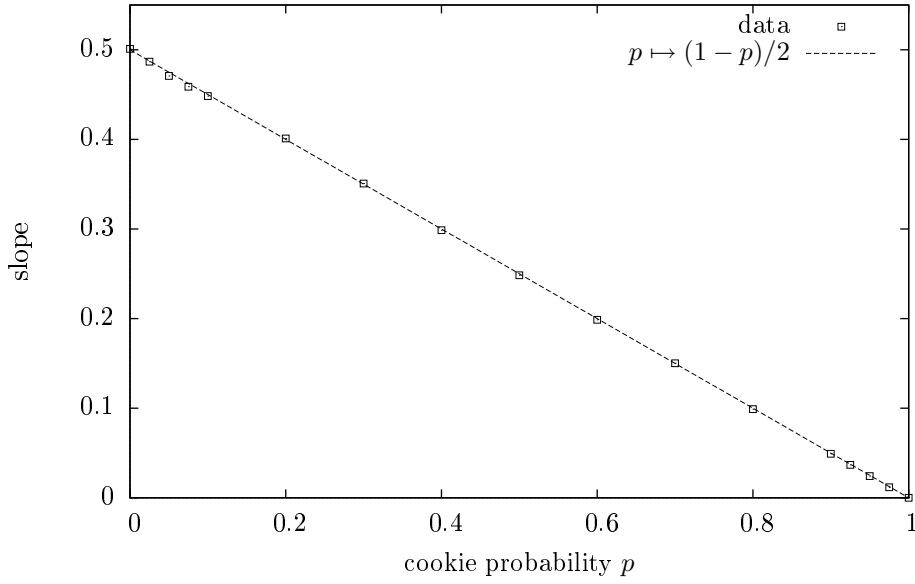


FIGURE 4.2. Results for an annealed 2-dimensional simulation. The graph shows the apparent slopes in Figure 4.1 (i.e. the limits of the functions of Figure 5.1) as a function of the cookie probability together with the graph of $p \mapsto (1-p)/2$.

p	0	.5	.975	0	.5	.975
n	1	1	1	2	2	2
exact	.5	.75	.9875	.375	.605469	.975292
simulated	0.50005	.74998	.98749	.37501	.605465	.97528

TABLE 1. Exact and simulated survival probability values $\mathbf{P}_p(S_n)$ on \mathbb{Z}^2 . The simulated values are calculated from the data used in Figure 4.1.

4.2. Annealed simulation on \mathbb{Z}^1 . A 1-dimensional simulation with 10^8 runs and $n_{\max} = 10000$ produces less satisfactory results as shown in Figure 4.3. The reasons behind this are explained in Subsection 5.2 below, where a discussion is given concerning the fluctuations of the empirical curves in the figures. Essentially, in the annealed case, small values of $\mathbf{P}_p(S_n)$ result in large errors (see Subsection 5.2 for more explanation) and therefore we modified the original algorithm by introducing a *stopping rule*: when the estimated value of $\mathbf{P}_p(S_n)$ reaches a certain small threshold value, we stop and do not simulate more iterations. Fortunately, when larger threshold values needed, they are actually large: we obtained slower convergence for large values of p , and, clearly, for those values, the probability $\mathbf{P}_p(S_n)$ is large. The threshold value was set $1/4000$, based on trial and error. This way, we stopped the iteration at $n_{\text{stop}}(p)$; the slopes were then calculated from the values at $4n_{\text{stop}}(p)/5$ and $n_{\text{stop}}(p)$. See Figure 4.4.

After adjusting the algorithm by using the above stopping rule, the curve indeed straightened out and the picture became very similar to the 2-dimensional one in Figure 4.2.

To verify the correctness of our simulation we computed the exact theoretical survival probabilities after the first two iterations. It is easy to see that $\mathbf{P}_p(S_1) = 1/2 + p/2$ and

$$\mathbf{P}_p(S_2) = 3/8 + 5p/16 + p^2/4 + p^3/16.$$

Table 2 compares some of the exact and simulated values.

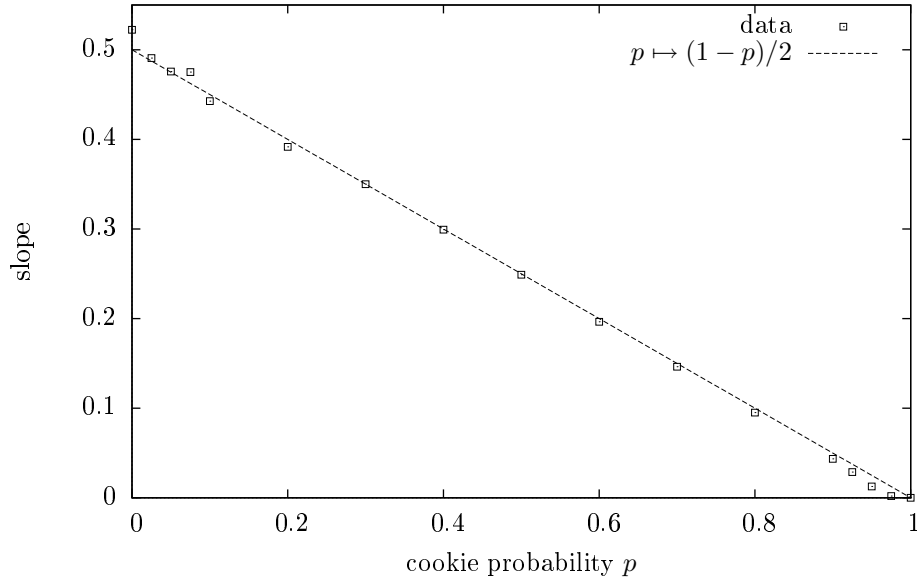


FIGURE 4.3. Results for an annealed 1-dimensional simulation. Every parameter for this simulation is chosen to be the same as that of Figure 4.2 except the dimension.

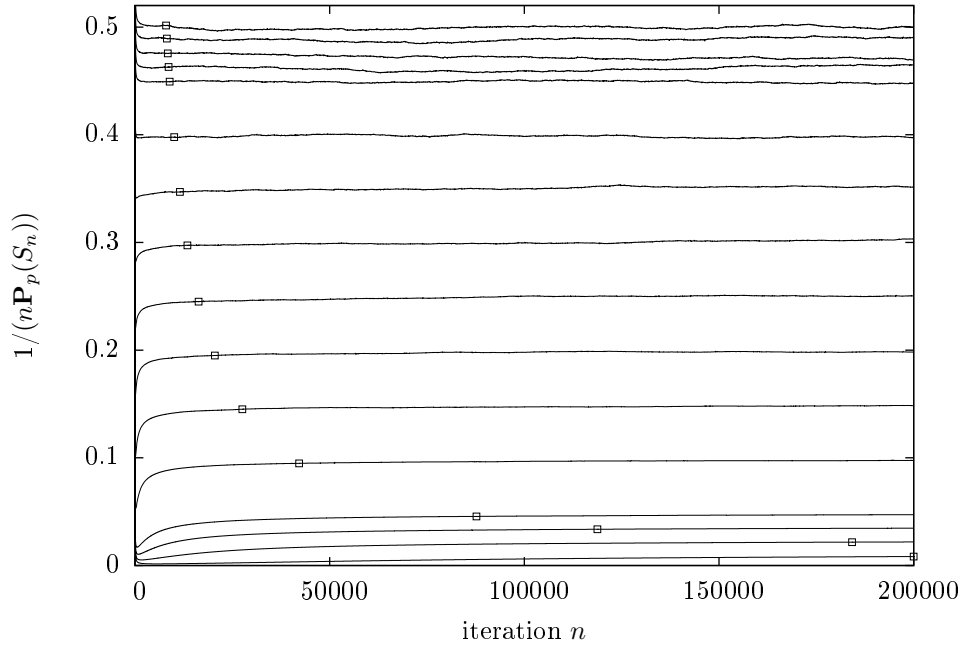


FIGURE 4.4. Annealed 1-dimensional simulation with 959965800 runs. For small values of p (lines at the top), a small iteration number would actually give better results, because otherwise the survival probability $\mathbf{P}_p(S_n)$ becomes too small, even with a huge number of runs. On the other hand, for large p values (lines at the bottom) one needs large iteration numbers because the convergence is apparently slow. The squares represent the iteration thresholds after which $\hat{\rho}_n < \frac{1}{4000}$.

p	0	.5	.975	0	.5	.975
n	1	1	1	2	2	2
exact	.5	.75	.9875	.375	.601563	.975272
simulated	0.50002	.7499992	.98749	.37502	.601569	.975269

TABLE 2. Exact and simulated survival probability values $\mathbf{P}_p(S_n)$ on \mathbb{Z}^1 . The simulated values are calculated from the data used in Figure 4.4.

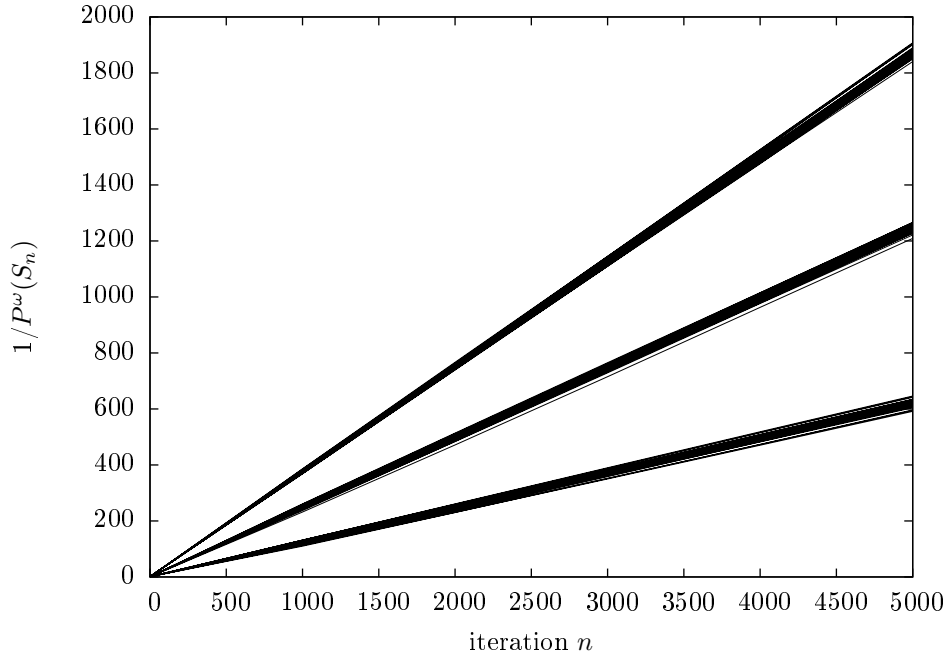


FIGURE 4.5. Results for a quenched 2-dimensional simulation with 10^8 runs. Each line represents a different cookie landscape. One such line is the result of 10^8 runs of the simulation. The lines are in three groups corresponding to three different cookie probability. Each group has 50 lines. The cookie probabilities from top to bottom are 0.25, 0.5 and 0.75. The total number of simulations required for this graph is $3 \cdot 50 \cdot 10^9$. The total running time was about 29 hours.

4.3. Quenched simulation. From the annealed simulation it has been clear that convergence is much faster in two dimensions than in one dimension. Therefore, in the quenched case we chose to present our results in the $d = 2$ case. In fact, we got qualitatively similar results for $d = 1$.

In Figure 4.5 we see three bundles corresponding to three values of p . Those bundles are very thin, so essentially the same thing happens for every realization, and the slopes of the lines are roughly $3/8$, $1/4$ and $1/8$ from top to bottom, corresponding to $p = 0.25$, $p = 0.5$ and $p = 0.75$, respectively. That is, for each one of these values of p , the slope is the same as in the annealed case.

Although Figure 4.5 is about the $d = 2$ case, we have a similar simulation result for $d = 1$; in fact *we conjecture that this qualitative behavior* (that is, the coincidence of the first order asymptotics of the quenched and annealed survival probability) *will hold for all $d \geq 1$.*

5. INTERPRETATION OF THE SIMULATION RESULTS

5.1. Main finding. Recall first the classic result due to Kolmogorov [7, Formula 10.8], that for critical unit time branching with generating function φ , as $n \rightarrow \infty$,

$$P(\text{survival up to } n) \sim \frac{2}{n\varphi''(1)}.$$

As a particular case, let us consider now a non-spatial toy model as follows. Suppose that branching occurs with probability $q \in (0, 1)$, and then it is critical binary, that is, consider the generating function

$$\varphi(z) = (1 - q)z + \frac{1}{2}q(1 + z^2).$$

It then follows that, as $n \rightarrow \infty$,

$$(5.1) \quad P(\text{survival up to } n) \sim \frac{2}{qn}.$$

Turning back to our spatial model, the simulations suggest (Figures 4.1 and 4.5) the *self averaging* property of the model: as explained in the previous section, the asymptotics for the annealed and the quenched case are the same. In fact, this asymptotics is *the same as the one in (5.1)*, where $p = 1 - q$ is the probability that a site has a cookie. In other words, despite our model being spatial, in an asymptotic sense, the parameter q simply plays the role of the branching probability of the above non-spatial toy model. To put it yet another way, q only introduces a ‘time-change.’

The intuitive picture behind this asymptotics is that *there is nothing either the environment or the BRW could do to increase the chance of survival*, at least as far as the leading order term is concerned (as opposed to well known models, for example when a single Brownian motion is placed into random medium [14]). Hence, given any environment the particles move freely and experience branching at q proportion of the time elapsed (quenched case), and the asymptotics agrees with the one obtained in the non-spatial setting as in (5.1). Furthermore, creating a ‘good environment’ (annealed case) and staying in the part of the lattice with cookies for very long would be ‘too expensive.’

Note that whenever the total population size reduces to one, the probability of that particle staying in the region of cookies is known to be much less than $\mathcal{O}(1/n)$ (hard obstacle problem for random walk). So the optimal strategy for this particle to survive is obviously not to try to stay completely in that region and thus avoid branching. Rather, survival will mostly be possible because of the potentially large family tree stemming from that particle. In fact, the formula $\mathbf{P}_p(S_n) \sim \frac{2}{qn}$, together with the martingale property of $|Z_n|$, implies that $\mathbf{E}_p(|Z_n| \mid S_n) \sim \frac{2}{q} \cdot n$.

Notice that the straight lines on Figures 4.2 and 4.3 start at the value $1/2$, that is, as $p \downarrow 0$, one gets the well known non-spatial asymptotics $2/n$, which is a particular case of Kolmogorov’s result above. We conclude that there is apparently *no discontinuity* at $p = 0$ (no cookies) for the quantity $\lim_{n \rightarrow \infty} nP(\text{survival up to } n)$.

5.2. Interpretation of the fluctuations in the diagrams. Since we estimated the reciprocal of the survival probabilities and not the probabilities themselves both in the annealed and the quenched case (Figures 4.1 and 4.5), one cannot expect good approximation results when those probabilities are small. Indeed, in the annealed case, if $\rho_n := \mathbf{P}_p(S_n)$ (with p being fixed) and $\hat{\rho}_n$ denotes the relative frequency obtained from simulations, then the Law of Large Numbers (LLN) only says, that if the number of runs is large, then the difference $|\rho_n - \hat{\rho}_n|$ is small. However, looking at the difference of the reciprocals

$$\left| \frac{1}{\rho_n} - \frac{1}{\hat{\rho}_n} \right| = \frac{|\rho_n - \hat{\rho}_n|}{\rho_n \hat{\rho}_n},$$

it is clear that a small ρ_n value magnifies the error; in fact the effect is squared as $\hat{\rho}_n$ is close to ρ_n , exactly because of the LLN. This effect is the reason of the ‘zigzagging’ of the line on Figure 4.3 for small p values. In fact, small p values result in small ρ_n values in light of

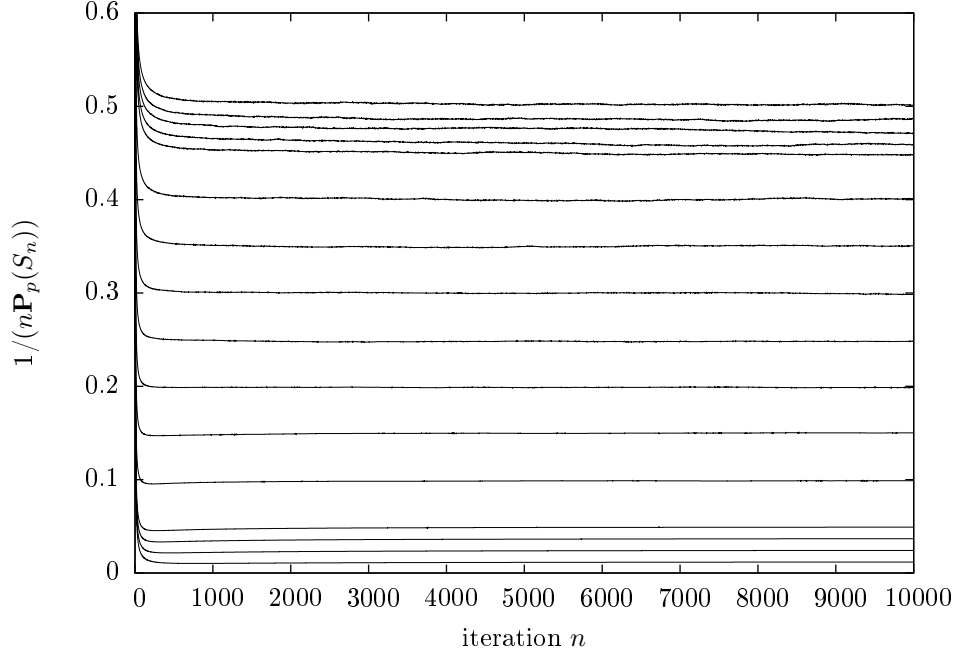


FIGURE 5.1. Results for an annealed 2-dimensional simulation with 10^8 runs. The graph shows the reciprocal of the survival probability divided by the number of iterations as a function of the number of iterations. The data used to create the graph is the same as that of Figure 4.1.

Theorem 2.1 and the continuity property mentioned at the end of the previous subsection. Clearly, there is a competition between ρ_n being small (as a result of p being small and n being large) on the one hand, and the large number of runs on the other. The first makes the denominator small in $\frac{|\rho_n - \hat{\rho}_n|}{\rho_n \hat{\rho}_n}$, while the second makes the numerator small according to LLN.

Looking at Figure 4.3, one notices another peculiarity in the 1-dimensional setting. For large values of p , the empirical curve is slightly under the straight line. The explanation for the relatively poor fit is simply that the iteration number is not large enough for the asymptotics to ‘kick in.’

These arguments are bolstered by the experimental findings that increasing the number of runs helps for small p values, whereas increasing the number of iterations helps for large ones. For example, in Figure 4.4 we increased the maximal iteration number n_{\max} to 200000 and plotted $n \mapsto (n\mathbf{P}_p(S_n))^{-1}$. One can see that for small p values it is beneficial to stop the iterations earlier, but for large values large iteration numbers give better results.

We do not have an explanation, however, for the deviation *downward* from the straight line (for large p values) in Figure 4.3. We suggest, as an open problem, to find at least a heuristic explanation for this phenomenon.

Interestingly, for higher dimensions there is apparently a perfect fit for large values of p , indicating that for higher dimensions the asymptotics is much quicker than for $d = 1$. Figure 5.1 checks the assumption (for $d = 2$, annealed) that the reciprocal of the survival probability is $\frac{qn}{2} + o(n)$ as $n \rightarrow \infty$, by dividing the reciprocal of the survival probability by n . The graphs convincingly show the existence of a limit, depending on p .

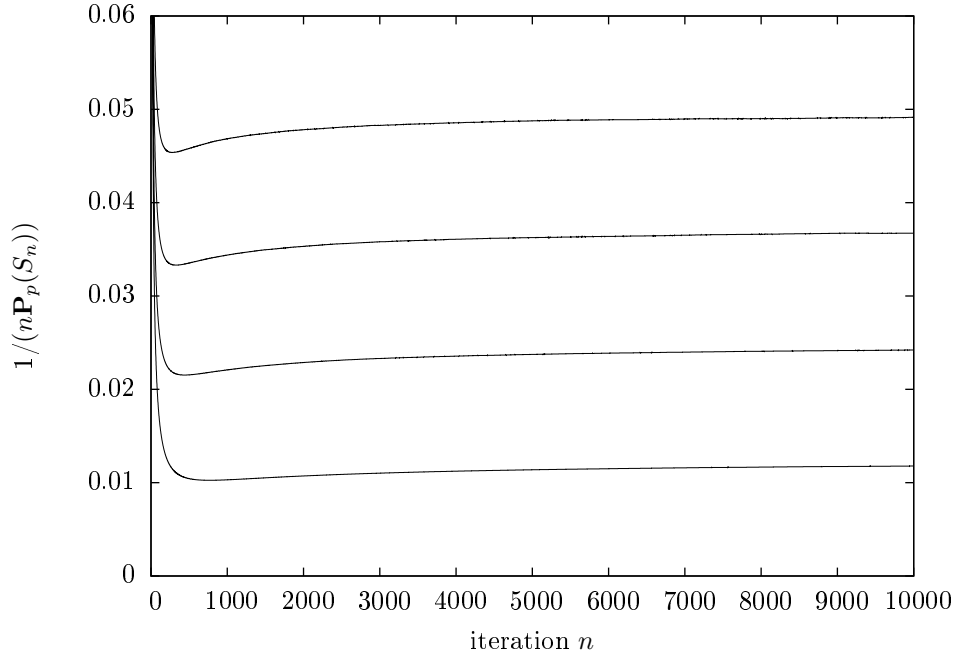


FIGURE 5.2. Zooming in at the bottom part of Figure 5.1 (i.e. large p values): the convergence is apparently from below.

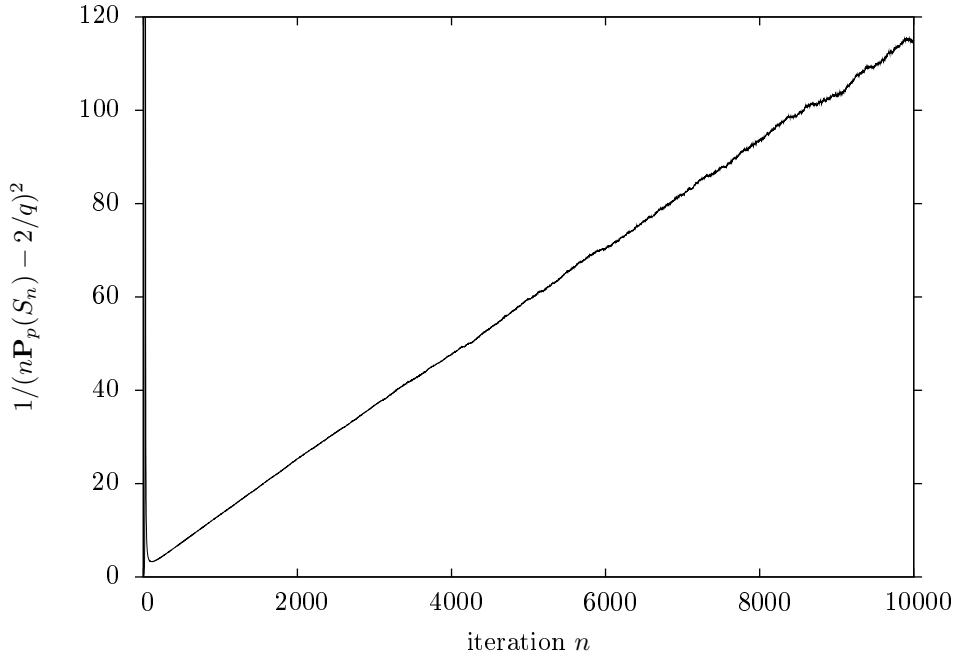


FIGURE 6.1. Annealed 1-dimensional simulation with 7,259,965,800 runs and $p = 0.5$.

6. BEYOND THE FIRST ORDER ASYMPTOTICS

In this section we will attempt to draw conclusions about more delicate phenomena beyond the first order asymptotics, and those conclusions will necessarily be less reliable than the ones in the previous sections.

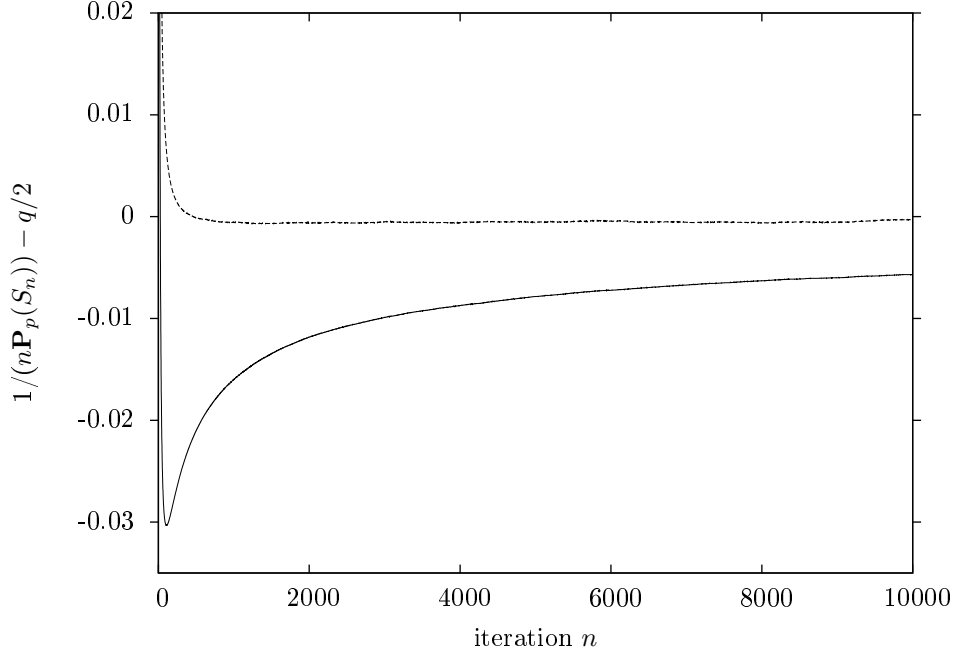


FIGURE 6.2. Annealed simulation with $p = 0.5$. The solid curve shows the 1-dimensional result, the dashed curve shows the 2-dimensional result.

6.1. Two dimensions. Consider again Figure 5.1. Zooming in gives Figure 5.2. Looking at Figure 4.4, Figure 5.1 and Figure 5.2, for small p values (top lines) the convergence seems to be from above, and for large p values it seems to be from below.

6.2. One dimension. For $d = 1$ we obtained figures somewhat similar to the 2-dimensional ones, which we summarize below without actually providing them. Simulation seems to suggest that for not too small values of p , the convergence is also from below; this is in line with the fact that, as we have already discussed, on Figure 4.3 the 1-dimensional empirical curve is *below* the straight line for large p 's. For very small p 's, the direction of the convergence is not clear from the pictures. Although the convergence is apparently quicker, the effects are 'blurred' due to the magnification of error explained earlier.

The following conjecture is based on Figure 6.1.

Conjecture 6.1 (Second order asymptotics). *The 1-dimensional annealed survival probability obeys the following second order asymptotics:*

$$\mathbf{P}_p(S_n) = \frac{2}{nq} + f(n),$$

where $\lim_{n \rightarrow \infty} f(n) \cdot n^{3/2} = C > 0$, and C may depend on p . That is, loosely speaking,

$$n\mathbf{P}_p(S_n) \approx \frac{2}{q} + c/\sqrt{n}.$$

6.3. Comparison between one and two dimensions. The annealed convergence to the limit $2/q$ seems to be quite different for $d = 1$ and $d = 2$. Figure 6.2 shows this difference, and in particular, it illustrates that in 1-dimension, the convergence is slower and it is apparently from below for $p = 0.5$.

7. CONCLUSION

In this paper we have investigated a critical branching random walk in d dimensions, $d \geq 1$, both in the annealed and the quenched setting. The environment is given by an IID collection of ‘cookies’ which suppress the branching at their locations. Each site contains a cookie with probability $p \in [0, 1]$. (The $p = 0$ and $p = 1$ cases are trivial.) We ran several types of simulations using a high performance computer cluster.

The simulations suggest *self averaging*: the asymptotic survival probability in n steps is the same in the annealed and the quenched case; it is $\frac{2}{qn}$, where $q := 1 - p$. This particular asymptotics indicates a non-trivial phenomenon both in the annealed and the quenched case: the tail of the survival probability is the same as in the case of *non-spatial* unit time critical branching, where branching only takes place with probability q for every particle at every iteration. Intuitively: there is no ‘good spatial survival strategy,’ given the environment (quenched setting), and establishing a ‘good environment’ is too costly (annealed setting).

We also obtained a second order asymptotics by the simulations. Namely, it appears that the 1-dimensional annealed survival probability satisfies that

$$\mathbf{P}_p(S_n) = \frac{2}{nq} + f(n),$$

where $\lim_{n \rightarrow \infty} f(n) \cdot n^{3/2} = C > 0$, and C may depend on p . (Here S_n denotes survival up to $n \geq 1$.)

These results are very clear from the simulations, yet their theoretical explanation is still to be achieved.

We have also found some interesting aspects of the simulation techniques, such as the dependence of the convergence speed on the dimension and on p . (For example convergence is much faster in two dimensions than in one dimension.)

Finally, simulations suggest that the asymptotics is sometimes achieved from one direction (for example, annealed convergence to the limit $2/q$ for one dimension and for $p = 0.5$ is apparently from below), the theoretical explanation of which, again, is a challenge.

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