Large-Scale Simulations of Diffusion-Limited n-Species Annihilation

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We present results from computer simulations for diffusion-limited n-species annihilation, \( A_i + A_j \rightarrow 0 \) \((i, j = 1, 2, \ldots, n; i \neq j)\), on the line, for lattices of up to \(2^{28}\) sites, and where the process proceeds to completion (no further reactions possible), involving up to \(10^{15}\) time steps. These enormous simulations are made possible by the renormalized reaction-cell method (RRC). Our results suggest that the concentration decay exponent for \(n\) species is \(\alpha(n) = (n - 1)/2n\) instead of \((2n - 3)/(4n - 4)\), as previously believed, and are in agreement with recent theoretical arguments. 

We also propose a scaling relation for \(\Delta\), the correction-to-scaling exponent for the concentration decay: 
\[
c(t) \sim t^{-\alpha(A + B t^{-\Delta})}.
\]

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I. INTRODUCTION

Diffusion-limited reactions have attracted much interest in recent years. The kinetics of such systems is dominated by local fluctuations in the concentration of the reactants, thus posing a formidable problem which has not yet been solved: there exists no comprehensive theoretical approach for the analysis of diffusion-limited processes.

Few select models are amenable to exact analysis. These include one-species annihilation, \( A + A \rightarrow 0 \), and two-species annihilation \( A + B \rightarrow 0 \) (see references therein). In one dimension, the particle density for one-species annihilation decays as \(c(t) \sim t^{-1/2}\), while for two-species annihilation (with equal initial concentrations and same diffusion constants for the two species) \(c(t) \sim t^{-1/4}\). In either case the result is strikingly different from the mean-field kinetics of the corresponding reaction-limited process, \(c(t) \sim 1/t\). To bridge the gap between these disparate behaviors, ben-Avraham and Redner proposed the \(n\)-species annihilation model, where particles belonging to the \(n\) species \(A_1, A_2, \ldots, A_n\) diffuse on the line and react immediately upon encounter, according to the scheme:

\[
A_i + A_j \rightarrow 0, \quad i, j = 1, 2, \ldots, n, \quad i \neq j.
\]

For \(n = 2\) we recover two-species annihilation, while in the limit \(n \rightarrow \infty\) encounters between like-particles are improbable and the model is equivalent to one-species annihilation. For intermediate values of \(n\), one expects \(c(t) \sim t^{-\alpha(n)}\).

In [4] it was proposed, following a heuristic scaling argument and treating fluctuations via the Van Kampen \(\Omega\)-expansion, that \(\alpha(n) = (2n - 3)/(4n - 4)\). This was supported by numerical simulations of lattices of typically \(10^6\) sites, and up to \(10^6\) time steps. (In one time step, all of the particles in the system move one lattice spacing each, on average.) Recently, we have conducted extensive numerical simulations, following the method of Renormalized Reaction-Cells (RRC) [6]. The systems involved are up to \(2^{28} \approx 2.7 \times 10^8\) sites long, and the processes were simulated to completion (until no further reactions are possible), for up to \(10^{15}\) time steps. The new data leads us to the conjecture that \(\alpha(n) = (n - 1)/2n\). We also find a correction to the main decay mode, of the form \(c(t) \sim t^{-\alpha(n)}(A + Bt^{-\Delta(n)})\), \(\Delta(n) = 1/2n\). The same results were found, independently (and unbeknownst to us), by Deloubré et al. [10]. In their theoretical derivation, they consider a simplified version of \(n\)-species annihilation, where domains of alternating species loose particles to reactions at one and the same rate, in a synchronous fashion. The approximation is more than reasonable, yet it does not rigorously apply to the original model, and analysis of corrections is certainly beyond its scope. Moreover, the simulations in [10] are comparable in size to those in [4]. In what follows, we report the results of our large-scale simulations, which strongly support the conclusions of [10]. We also propose a scaling relation for the correction exponent \(\Delta\) for \(n\)-species annihilation, and possibly for other reaction models where particles segregate into distinct domains.

II. SCALING

As is well known, local fluctuations in the concentrations of the different species drive the kinetics of \(n\)-species annihilation [3, 4]. An initially random homogeneous distribution of the particles evolves into a continuously growing mosaic of domains of alternating surviving species. Two length scales characterize the emerging distribution and dominate the system: the inter-domain distance — the distance between the last particle in a domain and the first particle in the domain next to it — \(\ell_{AB}(t)\), and the domain length, \(\ell(t)\) [11]. These quantities grow with time as

\[
\ell(t) \sim t^\beta, \quad \ell_{AB}(t) \sim t^\gamma.
\]
Once domains form, reactions might take place only at the domain boundaries, and particles have to diffuse across the domain gap $\ell_{AB}$ to react with other species. This takes a typical time of $\Delta t \sim \ell_{AB}^2/D$, where $D$ is the diffusion constant. The change in particle concentration during time $\Delta t$ equals the total number of domain boundaries divided by the lattice size $L$; $\Delta c \sim -(L/\ell)/L = -1/\ell$. Thus,

$$\frac{\Delta c}{\Delta t} \sim -\frac{D}{\ell \ell_{AB}}. \quad (3)$$

On substituting the relations (2) and $c(t) \sim t^{-\alpha}$, we derive the scaling rule

$$2\gamma + \beta - \alpha = 1. \quad (4)$$

Due to the underlying transport mechanism, we expect that domains grow diffusively, as $\ell \sim t^{1/2}$, so $\beta = 1/2$, and in effect there is only one independent exponent: $2\alpha - \gamma = 1/2$. The general scaling form holds also for two-species annihilation in the presence of drift (and with hard-core repulsion between like species), where $\alpha = 1/3$, $\beta = 7/12$, and $\gamma = 3/8$. \footnote{The $n$-species annihilation process is simulated as follows. The sites of a one-dimensional lattice are either empty or occupied by a particle (of one of the $n$ species). Periodic boundary conditions are imposed, so the lattice is effectively a ring. At each Monte Carlo step a particle is chosen randomly and is moved to the nearest site to its right or left, with equal probabilities. If the target site is occupied by a particle of a different species, then both particles are removed from the system, mimicking the reaction $1$. If the target site is occupied by a particle of the same species, then the move is disallowed and it does not take place. Regardless of the outcome, time is incremented by $1/N(t)$, where $N(t)$ is the total number of extant particles.}

\section{Simulation Results}

The $n$-species annihilation process is simulated as follows. The sites of a one-dimensional lattice are either empty or occupied by a particle (of one of the $n$ species). Periodic boundary conditions are imposed, so the lattice is effectively a ring. At each Monte Carlo step a particle is chosen randomly and is moved to the nearest site to its right or left, with equal probabilities. If the target site is occupied by a particle of a different species, then both particles are removed from the system, mimicking the reaction $1$. If the target site is occupied by a particle of the same species, then the move is disallowed and it does not take place. Regardless of the outcome, time is incremented by $1/N(t)$, where $N(t)$ is the total number of extant particles.

As the simulation proceeds, the particle concentration declines and the typical distance between particles increases. The time spent on simulating the diffusive motion of the particles until they interact grows even faster, as the square of the distance between them. Because of that, computer simulations are limited to relatively short times. This problem is overcome by the RRC method \footnote{Simulations were performed on DEC Alpha processors running Linux. Since each lattice site requires 6 bytes (for species, number of particles, and a pointer to a list of populated sites that is used for fast selection at each Monte Carlo step), with 2 Gigabytes ($2^{31}$ bytes) memory we were able to simulate lattices of up to $2^{28}$ sites. The compiler was given special #pragma pack(1) instructions to circumvent word alignment (which would allocate 32 bytes for our 6-byte site).}

To test the technique, we have simulated the cases of $n = 2$ and $n = 3$ on lattices of $2^{16} = 65,536$ sites, both in the RRC and the traditional simulation method. These lattices are small enough to enable the simulation of the process by the traditional method to completion. On the other hand, the system is large enough to let us examine the effect of the renormalizations: with $2^{16}$ sites and $c(0) = 1/16$ the RRC method requires 12 renormalizations. In the second, more stringent test, but the overall agreement is excellent. Similar results were obtained for the domain size and the distance between domains.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{fig1.png}
\caption{Comparison between the traditional (circles) and RRC (solid line) simulation methods. Plotted is the number of surviving particles $N(t)$ (top) and the local slope $\alpha(t) = -d \ln N(t)/d \ln t$ (bottom) for $2^{16}$-site lattices.}
\end{figure}
Having gained some confidence in the RRC method, we proceed to larger simulations. In Fig. 2 we show the surviving number of particles, \( N(t) \), at time \( t \), for \( n = 3, 4, \) and 5, and several lattice sizes. In Fig. 3 we plot the local decay exponent \( \alpha(t) \) for our largest simulations of \( n = 3 \). The maximum of the curve at \( t \approx 10^4 \) agrees with the earlier prediction that \( \alpha = 3/8 \). (Indeed, simulations in \( 4 \) yielded a somewhat smaller value than the theoretical \( 3/8 \), in perfect agreement with current results.) However, \( \alpha(t) \) is seen to diminish with time, suggesting a long-time asymptotic limit of \( \alpha \approx 1/3 \). This limiting value is confirmed in the data collapse (especially at long times) of Fig. 4 where we plot \( t^\alpha c(t) \) vs. \( t^\beta / L \) for various system sizes, and \( \alpha = 1/3, \beta = 1/2 \). Independent measurements show that \( \beta = 1/2 \), as assumed, to within \( 2\% \), and the data collapse of Fig. 4 deteriorates with other choices for the values of \( \alpha \) and \( \beta \).

![FIG. 2: Concentration decay for \( n = 3, 4, \) and 5-species annihilation. Plotted is the number of surviving particles, \( N(t) \), for system sizes \( L = 2^{16}, 2^{20}, 2^{24}, \) and \( 2^{28} \) (bottom to top).](image-url)

![FIG. 3: Local decay exponent for 3-species annihilation.](image-url)

We have analyzed in this fashion \( n = 3, 4, \) and 5-species annihilation, and measured the exponents \( \alpha, \beta, \) and \( \gamma \). Our results are summarized in Table I. In all cases, the scaling relation (4) seems to hold, and \( \beta = 1/2 \) to within numerical errors. Looking for a simple expression of these results, that would have the appropriate limits for the known cases of \( n = 2 \) (two-species annihilation) and \( n \to \infty \) (one-species annihilation), we were led to the conjecture (6) for the region \( n = 3 \) on a lattice, where our data is most reliable. The results are most compatible with \( \Delta = 1/6 \) (for \( n = 3 \)). Similar tests for other values of \( n \) lead us to the conjecture that \( \Delta(n) = 1/2n \).

Finally, let us address the issue of corrections to scaling of the concentration decay. We look for corrections of the form

\[
c(t) \sim t^{-\alpha}(A + Bt^{-\Delta})^\gamma,
\]

where \( A \) and \( B \) are constants. Our strategy consists of performing a least squares linear fit of \( A + Bt^{-\Delta} \) to \( t^{\alpha}c(t) \), for different powers \( \Delta \), and searching for the value of \( \Delta \) which minimizes the error. The scaling form (6) is expected to work only after the asymptotic regime sets in, and before finite-size effects begin, and the sticky part of our procedure is deciding which times demarcate this region. Experimenting with different choices gives us a feel for the errors involved. In Fig. 4 we show best fits for the region \( t = 10^8 - 10^{12} \), for \( n = 3 \) on a \( L = 2^{28} \) lattice, where our data is most reliable. The results are most compatible with \( \Delta = 1/6 \) (for \( n = 3 \)). Similar tests for other values of \( n \) lead us to the conjecture that \( \Delta(n) = 1/2n \).

The correction exponent can be understood by a simple-minded argument. In deriving (8) we have assumed that the typical distance between reacting particles, at the edges of adjacent domains, is \( \xi_{AB} \). While this is correct, we note that, had the distribution of particles been homogeneous, the distance between reacting pairs would
where $\ell_{AA} \sim L/c \sim t^\alpha$, quite different from the assumed $\ell_{AB} \sim t^\gamma$. Using $\Delta t \sim \ell_{AA}^2/D$ instead of $\ell_{AB}^2/D$, yields a faster decay; $c \sim t^{-(1-\beta)}$. Diffusion provides a natural drive toward a homogeneous distribution, and so it is conceivable that this faster mode of decay is manifested as a correction to the main behavior, $c \sim t^{-\alpha}$. It follows from (6) that the correction exponent is

$$\Delta = 1 - \beta - \alpha = \frac{1}{2n},$$

where the last equality applies to $n$-species annihilation, provided that the conjecture holds. The more general relation works well for two-species annihilation with drift, where $\alpha = 1/3$, $\beta = 7/12$, and $\Delta = 1/12$.

IV. SUMMARY AND DISCUSSION

We have presented large scale simulation results for diffusion-limited $n$-species annihilation, in one dimension, using the RRC method. Our simulations contradict previous work and are in favor of new theoretical arguments advanced by Deloubrière et al. We have also provided a new scaling relation for the correction-to-scaling exponent $\Delta$, valid for diffusion-limited reactions in one dimension, where the particles segregate into alternating domains. The corrections to the main decay mode are large, and explain the failure of to obtain the correct asymptotic behavior with the size of simulations employed at that time. An important conclusion to be drawn is that predicting asymptotic behavior from the typical size of simulations used commonly in the field is dangerous. More advanced techniques and larger simulations seem to be imperative.

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