Abstract

In this paper we examine a number of models that generate random fractals. The models are studied using the tools of computational complexity theory from the perspective of parallel computation. Diffusion limited aggregation and several widely used algorithms for equilibrating the Ising model are shown to be highly sequential; it is unlikely they can be simulated efficiently in parallel. This is in contrast to Mandelbrot percolation that can be simulated in constant parallel time. Our research helps shed light on the intrinsic complexity of these models relative to each other and to different growth processes that have been recently studied using complexity theory. In addition, the results may serve as a guide to simulation physics.

Keywords: Cluster algorithms, computational complexity, diffusion limited aggregation, Ising model, Metropolis algorithm, P-completeness
1 Introduction

Random fractals are a major focus of investigation in statistical physics. Such patterns occur at equilibrium critical points and arise through a variety of non-equilibrium dynamical processes. A number of models generate random fractals including diffusion limited aggregation (DLA) and the Ising model at criticality. These models have been extensively studied by computer simulation methods and, in some sense, they are defined by the algorithms that are used to simulate them. In this paper we examine such defining algorithms from the viewpoint of the theory of computational complexity.

Computational complexity is the branch of theoretical computer science that seeks to quantify the resources required to solve problems. One of the main achievements of complexity theory is the identification of a hierarchy of complexity classes. The classes differ with respect to how the various resources, such as time, space and processors, scale in proportion to problem size. For example, does the running time increase as a logarithmic, power or exponential function of the problem size? Our emphasis is on parallel computational complexity. We seek to answer the following question: how do the number of processors and the amount of time required to simulate a system on a massively parallel computer increase with the system size?

The motivation for this work is two-fold. First, computational complexity may serve as a guide to simulation physics. With the growing availability of massively parallel computers, it is important to investigate models from the perspective of parallel complexity. Another, perhaps more significant, motivation is to provide an alternative characterization of these models. An enormous amount of effort has gone into characterizing the morphology of fractal patterns via critical exponents, fractal and multifractal dimensions, scaling functions, and so on. Such characterizations fail to adequately distinguish these models from the standpoint of what can be described intuitively as complexity. We believe that the intuitive notion of physical complexity is at least partially captured by the computational complexity measure of parallel time (with the number of processors appropriately restricted). This idea, in a slightly different form, has been previously proposed by Bennett [1].

In a nutshell, the idea is that simple objects can be generated quickly while complex objects require a long history for their formation. We illustrate this by comparing two random fractals. The first is Mandelbrot percolation [2]; an example of which is depicted in Fig. 1. We show Man-
delbrot patterns require only (non-uniform) constant parallel time to generate. Though they are fractals, there is very little interesting morphology; the structure on each length scale is independent of the structure on other length scales. Many properties of Mandelbrot percolation are susceptible to rigorous analysis [3]. The second example is DLA [4] that generates fractal patterns like those shown in Fig. 2. DLA patterns are produced by a highly sequential algorithm that seems to require polynomial (in the size of the aggregate) parallel time. DLA patterns reflect a subtle interplay of randomness and structure on many length scales. DLA has remained largely refractory to theoretical analysis. Whether or not one accepts a definition of physical complexity in terms of computational complexity, it is interesting that a variety of models in statistical physics can be sharply separated from one another by a fundamental new yardstick.

Our research extends a study of the complexity of a number of growth models. Ref. [5] is concerned with a fluid invasion model that generates clusters with the same statistics as DLA. This model is shown to be inherently sequential (technically, $P$-complete) and so it is unlikely that it can be efficiently simulated in parallel. Here we show that the original random walk dynamics for generating DLA clusters is also inherently sequential. In Ref. [6] we considered a number of other growth models—inv asion percolation, Eden growth, ballistic deposition and solid-on-solid growth—and showed that all of these models can be efficiently simulated in parallel. The fractal patterns associated with them can be generated on a parallel computer in a time that scales logarithmically in the system size while using a reasonable number of processors. Although each of these models is less complex than DLA, each is more complex than Mandelbrot percolation.

Other applications of computational complexity theory to statistical physics have focused mainly on the existence of polynomial time sequential algorithms. For example, the problems of finding the exact ground states of spin glasses [7] and computing self-avoiding walks in a random environments [8] have been shown to be computationally intractable (technically, $NP$-complete). On the other hand, a polynomial time algorithm exists for the random field Ising model [9]. There is also work that establishes the complexity of finding the partition function of the Ising model and related spin models on arbitrary lattices either exactly [10] or approximately using Monte Carlo methods [11].

In Section 2 we give an introduction to computational complexity theory. A reader familiar with this field may want to skim this section. In Section 3 we investigate the computational complexity of the following systems: Man-
delbrot percolation, DLA, Metropolis dynamics for the Ising model, Wolff dynamics for the Ising model, and Swendsen-Wang dynamics for the Ising model. Section 4 is devoted to a discussion of the results.

2 Computational Complexity Background

In this section we provide an introduction to computational complexity theory. The reader can find further information and details in a number of texts [12, 13] and monographs [14, 15, 16].

2.1 The parallel random access machine

The theoretical model we focus on is the parallel random access machine or P-RAM. It is the most commonly used model in parallel computation. We describe the P-RAM and then relate its resource usage to the corresponding measures for actual parallel computers.

The P-RAM consists of a number of processors each with local memory and having access to a common global random access memory. All processors run the same program but are distinguished by non-negative integer labels so that the processors may operate on their own data or skip instructions. Input to the machine is placed in designated, consecutive global memory locations as is output. The P-RAM is in the class of single-instruction multiple-data-stream (SIMD) models. The processors run synchronously and in each time step a single random access machine (RAM) instruction [17] or a global memory access instruction is executed by a subset of the processors. Examples of typical instructions are ‘write the contents of the accumulator to memory location a’ and ‘add the contents of the accumulator to the contents of register a, placing the sum in the accumulator.’

Although many processors may read the same memory location at a particular time, difficulties arise if multiple processors attempt to write to the same location. One frequently used arbitration scheme is the concurrent write model in which processors are assigned a write priority. When more than one processor attempts to write to a given location, the processor with the highest priority succeeds. This model is known as the PRIORITY CRCW P-RAM [18]. We adopt this model and simply refer to it as the P-RAM.

In the P-RAM model any processor can access any global memory location in one time step; the model allows unlimited parallelism. For this reason
the P-RAM serves as a convenient model for designing and analyzing parallel algorithms, for studying processor and time requirements and for proving lower bounds. Although the P-RAM is overly simplistic in its assumptions, it can nevertheless be simulated on models of parallel computation with more restricted connectivity such as the hypercube. These simulations usually have a slow down of a logarithmic factor and require roughly the same amount of hardware as the corresponding P-RAM computations, see [19] for additional details and references.

As an example of the utility of parallelism, consider the task of computing the parity of \( n \) bits. Parity is the problem of determining whether there is an even number of 1’s in the input. Initially, the \( n \) bits are stored in global memory locations \( 1, \ldots, n \). A P-RAM program that computes parity uses \( n/2 \) processors numbered \( 1, \ldots, n/2 \) to add \( n/2 \) pairs of bits (modulo 2) in parallel. That is, processor 1 adds the contents of locations 1 and 2 storing the result (modulo 2) in location 1, processor 2 adds locations 3 and 4 storing the result (modulo 2) in location 2, and so on. Similarly, the resulting \( n/2 \) values are added pairwise (modulo 2) by \( n/4 \) processors. This process is repeated until, after \( \lceil \log_2 n \rceil \) time steps, parity is computed. Notice, in this case the algorithm’s output is placed in global memory cell 1 by processor number 1. The algorithm runs in \( O(\log n) \) time using \( n/2 \) processors.

2.2 Complexity classes

The primary question addressed by computational complexity theory is how the difficulty of a computation scales with the size of the problem instance. Complexity theory usually focuses on decision problems. An instance of a decision problem is a string of bits encoding the problem; the solution is simply a 1 or 0. If the solution for input \( x \) is 1, we say that \( x \) is ‘accepted’ and otherwise \( x \) is ‘rejected.’ In this sense, a decision problem is defined by its set of accepted strings. The problem size, \( n = |x| \), is the length of the encoded input. A simple example involving the parity problem discussed above is as follows:

**Parity**

*Given:* \( b_1, \ldots, b_n \), where \( b_i \in \{0, 1\} \).

*Problem:* Do an even number of the \( b_i \)'s have value 1?

In this case the input is easily encoded using exactly \( n \) bits. The output is a 1 if there are an even number of \( b_i \)'s with value 1 and 0 otherwise. It is easy
to see that the answer may be found on a single processor computer (such as a RAM or more familiar desktop computer) with a running time that scales linearly in $n$ by simply scanning through the bits and maintaining their sum modulo 2.

We now define several important complexity classes for parallel computation.

**Definition 2.1**

- The class $AC^0$ consists of those decision problems that can be solved on a P-RAM in $O(1)$ (constant) time using $n^{O(1)}$ (polynomial) processors.

- The class $NC$ consists of those decision problems that can be solved on a P-RAM in $(\log n)^{O(1)}$ (poly-logarithmic) time using $n^{O(1)}$ processors.

- The class $P$ (polynomial time) consists of those decision problems that can be solved on a P-RAM in $n^{O(1)}$ time using $n^{O(1)}$ processors.

It is easy to see that $AC^0 \subseteq NC \subseteq P$. It is known that $AC^0 \neq NC$ and, while no proof yet exists, it is widely believed that $NC \neq P$. The classes in Def. 2.1 are robust in the sense that they may be equivalently defined with respect to several different computation models [15]; they are not tied to the P-RAM model of parallel computation.

All of the problems considered in this paper are in the class $P$. It is generally accepted that problems in the class $P$ have feasible sequential time solutions. The question we pose for the fractal models is whether a polynomial time problem can be qualitatively sped up via massive parallelism. For the parity example, the sequential solution mentioned takes $O(n)$ time. The parallel solution outlined previously shows a P-RAM can solve this problem in $O(\log n)$ time using $n/2$ processors. Thus, the parity problem is in the class $NC$ and a qualitative speed-up is achieved in the parallel setting. On the other hand, parity is *not* in $AC^0$, see [15] for example.

We will use the terminology that problems in $NC$ (and thus $AC^0$) are ‘efficiently solved in parallel,’ since we obtain a qualitative speed-up solving these problems in parallel. On the other hand, problems that are in $P$ but likely not in $NC$ are called ‘inherently sequential.’ The running time for solving an inherently sequential problem cannot be decreased from polynomial to poly-logarithmic using a polynomial number of processors. Below we identify a class of problems that are in $P$ but not $NC$ (unless it happens that $NC = P$).
In order to proceed we need to be able to relate problems to one another. This is accomplished via the notion of reduction. The idea is similar to a commonly used programming practice. To solve one problem, we often use a subroutine call to a different problem. In this sense we reduce our original problem to the one involved in the subroutine call. More formally,

**Definition 2.2** Let \( n = |x| \). (Throughout, \( |x| \) denotes the length of string \( x \) and a decision problem \( D \) is represented as a set of accepted strings.) Decision problem \( D_1 \) is NC many-one reducible or NC reducible \((≺)\) to decision problem \( D_2 \) if there exists a function \( f \) such that \( x \in D_1 \) if and only if \( f(x) \in D_2 \), and \( f \) can be computed on a P-RAM in \((\log n)^{O(1)}\) time using \( n^{O(1)} \) processors.

If \( D_1 \prec D_2 \) then \( D_1 \) is ‘no harder’ than \( D_2 \). This is because we could solve \( D_1 \) using an algorithm for \( D_2 \), where the input to \( D_2 \) is produced by an efficient calculation involving \( f \). We can also compare a given problem to an entire complexity class, via the concept of ‘completeness.’

**Definition 2.3** A decision problem \( D \) is P-complete if

1. \( D \in \mathbf{P} \) and
2. for all \( D' \in \mathbf{P} \), \( D' \prec D \).

The P-complete problems are therefore the hardest problems in \( \mathbf{P} \). Based on these definitions, the following theorem is straightforward to obtain.

**Theorem 2.4** If any P-complete problem is in NC then NC = P.

Thus, if the well-known conjecture in computer science that \( \text{NC} \neq \text{P} \) holds (and there is lots of evidence supporting this conjecture [14]), P-complete problems are inherently sequential.

\( \prec \) is a transitive relation. That is, if \( D_1 \prec D_2 \) and \( D_2 \prec D_3 \), then \( D_1 \prec D_3 \). Therefore, if some problem \( D' \) is shown to be P-complete and \( D' \prec D \) then \( D \) must be as difficult to solve as \( D' \). In this case we say \( D \) is P-hard. If \( D \) is also in \( \mathbf{P} \), it is P-complete as well. Using transitive reductions, a large number of P-complete problems have been identified and no efficient parallel solution has been found for any of them, providing evidence for the conjecture. In this paper we will prove that several problems from statistical physics are P-complete by showing that known P-complete problems reduce to them.
The fundamental P-complete problem is the circuit value problem (CVP); it is phrased in terms of Boolean circuits. Before describing CVP, we give an informal description of circuits. A Boolean circuit is a collection of connected NOT, AND and OR gates. NOT gates have one input and multiple outputs; AND and OR gates have multiple inputs and multiple outputs. The fan-in (fan-out) is the number of inputs (outputs) of a gate. The connection of the gates is ‘feedforward.’ That is, it must be possible to number the gates so that the outputs of a gate are connected to the inputs of gates with higher numbers. Such a numbering is called a topological numbering and we say the gates are in topological order. In calculating outputs from inputs each gate computes its Boolean function just once. Sometimes gates other than NOT, AND and OR are considered. The size of a circuit is defined as the number of gates. The depth is the longest path from an input to an output.

**Circuit value problem (CVP)**

**Given:** A compact encoding $\pi$ of a Boolean circuit together with its inputs $x_1, \ldots, x_n$, and a designated output gate $g$.

**Problem:** Does $g$ evaluate to 1 on input $x_1, \ldots, x_n$?

**Theorem 2.5** The circuit value problem is P-complete \[20\].

Numerous variants of CVP are P-complete \[14\]. In NOR CVP the circuit consists entirely of NOR gates with fan-in and fan-out two. NOR CVP without fan-out restrictions is also P-complete for planar circuits; this version is called planar NOR CVP. In monotone CVP the circuit is composed of AND and OR gates (NOT gates are absent) having fan-in and fan-out two. This problem is P-complete for arbitrary circuits but it is in NC for planar circuits. We shall make use of planar NOR, monotone and other restricted variants of CVP in Section 3.

A proof that a problem $D$ is P-complete via a reduction from CVP is tantamount to what in other contexts has been called ‘computational universality.’ The dynamics of hard spheres in classical mechanics \[21\] and some cellular automata rules \[14, 22\] have been shown to be computationally universal. Our proofs that DLA and various Ising Monte Carlo dynamics are P-complete depend on showing that arbitrary logical calculations can be embedded in these dynamics.

\footnote{A compact encoding of a circuit is polynomial in the circuit size.}
In addition to the resources of parallel time and number of processors, the notion of uniformity plays an important role in computations. Roughly speaking, a uniform solution to a problem uses the “same” program for each problem size whereas a non-uniform solution may use a different program for each size. For example, simulating the Ising critical point in three dimensions using conventional Monte Carlo methods is a non-uniform problem because the critical temperature is required as a parameter in the algorithm. As the system size increases, the program must contain an increasingly accurate value of the critical temperature. On the other hand, simulating DLA clusters is a uniform task since no fine tuning of parameters is required. The same can be said for other ‘self-organized’ critical points such as invasion percolation. Recently, a uniform algorithm for sampling Ising critical points has been developed \[23\].

2.3 Parallel time and logical depth

The P-RAM model and the complexity measures that are built from it are in some sense unphysical because unit time is assigned to a single read or write step. Eventually, as such a device is scaled up, the communication time between processors and memory dominates the running time and the unit time assumption fails. Indeed, in the limit of large systems all of the models discussed here require polynomial time to simulate on any real world device because all are capable of generating random patterns with correlations on the scale of the system size. These correlations cannot be set up without communication across the system and this requires polynomial time.

Nonetheless, parallel time correctly identifies an important aspect of the problem which can be called ‘logical depth.’ The logical depth is the minimum number of logical operations that must be carried out in sequence before a problem is solved. This concept can be made rigorous by considering families of Boolean circuits \[15\]. A family of Boolean circuits, one for each problem size, can simulate a P-RAM programmed to solve a given problem and vice versa. The definitions of the complexity classes \(\text{AC}^0\), \(\text{NC}\) and \(\text{P}\) can be stated in terms of families of Boolean circuits: the number of processors corresponds roughly to the size of the circuit (number of gates) and the parallel time roughly to the depth of the circuit (length of the longest path from input to output). Thus for example, a problem is in the class \(\text{NC}\)
if it can be solved by a uniform family of Boolean circuits having polynomial size and poly-logarithmic depth in the number of inputs (the problem size).

A few comments are in order regarding the number of processors. If only one processor is allowed, then all the problems treated here require polynomial time. If on the other hand, the number of processors is unrestricted it can be shown [24] that all the problems discussed here are solvable in constant P-RAM time using exponentially many processors (or equivalently by circuit families with exponential size and constant depth) and again the interesting distinctions based upon parallel time disappear. Interesting results are found when polynomial parallelism is permitted.

2.4 Complexity of sampling methods

Computer scientists study decision problems whereas computational statistical physicists are usually concerned with sampling problems—generating states from some equilibrium or nonequilibrium distribution. Sampling algorithms require a supply of random numbers and produce as output a system configuration. This configuration is described by $m$ bits representing the degrees of freedom of the system expressed in binary. One can extend the ideas of complexity theory to sampling methods by introducing probabilistic P-RAM’s in which each processor is equipped with a register for generating random bits.

Instead of producing random bits dynamically one could equivalently produce the required random bits in advance and include them as inputs to a deterministic calculation. In this way a sampling method is reduced to $m$ decision problems, one for each binary degree of freedom. An example of such a decision problem is ‘Does Ising spin $s_j$ ($1 \leq j \leq m$) have value +1 after $M$ iterations of the Monte Carlo procedure using random numbers $x_i$?’ Note that these $m$ decision problems may be run in parallel with, in the worst case, a factor of $m$ increase in the number of processors. Therefore, the sampling algorithm has the same parallel time requirement up to a constant factor as the associated decision problem.

In statistical physics, the problem size is conventionally identified with the system size; the number of bits, $m$ required to specify a system configuration. This differs from complexity theory where it is the number of bits required to state the problem that is identified as the problem size. The following definition insures that the two notions of problem size are compatible. For a given sampling method with $r$ random inputs, $o$ ordinary inputs and $m$ outputs, we define the associated natural decision problem as follows.
The input is of length $m + o + r$. The first $m$ bits represent the degrees of freedom of the system. Of these bits exactly one is a 1. The position of the 1 specifies which degree of freedom of the system (e.g. which Ising spin) is to be evaluated. Since the selected degree of freedom is expressed in unary, the decision problem size is at least as great as the system size. For example, to represent the fifth out of ten degrees of freedom our unary expression would be ‘0000100000.’ The next $o$ bits are the ordinary inputs to the problem expressed in a suitably compact form. These inputs might include the size of the lattice, the temperature, the number of iterations of an elementary Monte Carlo step and other relevant parameters expressed in binary notation. The final $r$ bits are the random bits needed for the sampling method. So that the answer or other potentially useful information is not built into these bits, we require that they be interpreted as independent random variables that take the value 1 with probability 1/2. We restrict our attention to ‘reasonable’ sampling methods where $r$ is bounded by a polynomial in $m$.

The decision problem for a sampling method can now be studied using conventional computational complexity theory. It must be emphasized that the complexity of the decision problem is only an upper bound on the complexity of sampling a given distribution. The reason is that the decision problem is associated with a particular sampling method. It may be that an alternative method leads to a less complex decision problem. In principle we would like to know how the time, number of processors and number of random bits scale with $m$ for the optimal sampling method. Unfortunately, tools for studying this question have yet to be developed. Instead, we focus on the complexity of several known sampling methods. Nonetheless, if the best known sampling methods are investigated and their complexity is established, it is plausible that the complexity of sampling has also been found. (Note, proving that a particular sampling method is optimal seems to be a very difficult task.)

3 Complexity of Random Fractals

In this section we consider the following models: Mandelbrot percolation, diffusion limited aggregation and the Ising model. We discuss sampling methods for these systems and then study the parallel computational complexity of the associated decision problems. Each model generates random

\footnote{This helps insure that the problems considered are in P and that the number of processors used will be polynomial in the input size.}
mass fractals—sets of ‘occupied’ sites whose number scales as a noninteger power of the lattice size.

3.1 Mandelbrot percolation

This random fractal was first described by Mandelbrot \cite{2}. It was analyzed by rigorous methods in \cite{3}, and was later generalized and applied as a model of a fractal porous medium \cite{25, 26, 27}. Mandelbrot percolation is defined on a $d$-dimensional lattice. It is parameterized by a rational retention factor $Q$ ($0 \leq Q < 1$), a positive integer rescaling factor $N$ and iteration number $k$. System configurations are described by a bit at each lattice site. If the bit is a 1, we say the site is ‘occupied.’ For purposes of illustration, we consider the two dimensional version on an $N^k \times N^k$ square lattice. A configuration is generated in the following way: at the $i$th step ($0 \leq i \leq k-1$) the lattice is completely divided into, $N^i \times N^i$ non-overlapping squares and each square is independently ‘retained’ with probability $Q$. If a square is retained, the site(s) in it are not changed. If a square is not retained then all of the site(s) in it are changed to unoccupied. After $k$ steps unoccupied regions with a wide range of sizes are typically created. The resulting set of occupied sites is a random fractal with limiting Hausdorff dimension, $D_H = 2 + (\log Q)/(\log N)$ if $D_H > 0$. A realization of Mandelbrot percolation with $N = 2$, $Q = .9$ and $k = 7$ is shown in Fig. 1.

A natural decision problem associated with Mandelbrot percolation takes as input random numbers, $x_i$ with $0 \leq x_i < 1$. These numbers are used to generate ‘retention bits’ that are 1 if $x_i < Q$ and 0 otherwise. Each retention bits determines whether a particular square of a given size is retained.

**Mandelbrot percolation** (dimension $d$, scale factor $N$, precision $b$)

**Given:** A non-negative integer $k$, a designated lattice site $s$ expressed in unary with $|s| = N^{dk}$, a retention factor $Q$ ($0 \leq Q < 1$) with $Q$ represented by a $b$-bit binary number and a list of $(N^{dk} - 1)/(1 - N^{-d})$ random numbers $x_i$ with $0 \leq x_i < 1$ expressed as a $b$-bit number.

**Problem:** Is site $s$ occupied by the Mandelbrot percolation process?

The instances of Mandelbrot percolation require that the dimension, scale factor and precision are all fixed inputs. In terms of the discussion of

\footnote{Our method of producing random numbers via coin tossing suggests this coding choice. Such a scheme does not allow all possible rationals in the interval $[0, 1)$ to be represented.}
A constant time P-RAM algorithm for Mandelbrot percolation is sketched below. First, retention bits for every square of each size are computed in parallel by comparing the $x_i$ to $Q$. Since $b$ is a constant, this can be done in constant time. The $m$ retention bits for the individual sites are placed in memory cells 1 to $m$. For each site $j$ ($1 \leq j \leq m$) the occupancy of $j$ is determined by taking the AND of all the retention bits of the $k$ squares containing $j$. To compute the AND, all processors reading a retention bit 0 write a value of 0 into global memory cell $j$. This step uses $km$ processors. Note, cell $j$ is 1 if site $j$ is occupied and 0 otherwise. Next, the AND of cell $j$ and the $j^{th}$ place in the unary expression of $s$ is computed; the result placed in cell $j$. Now, cell $j$ is 1 if and only if site $j$ is occupied and is the selected site. Finally, the OR of cells 1 through $m$ is taken (by having any processor reading a 1 write to memory cell 1) to determine if the selected site is occupied.

This P-RAM algorithm uses constant time and polynomial ($km$) processors. For problems in constant time a very strong notion of uniformity (meaning highly uniform), DLOGTIME, is typically imposed. The algorithm described does not seem to be DLOGTIME uniform. This is because for each lattice size, a different program may be needed to quickly incorporate information about which retention bits belong to a given site. Note that in general proving a problem does not meet a given uniformity condition is very difficult. Our algorithm is $\text{NC}^2$ uniform. We have the following:

**Theorem 3.1** Mandelbrot percolation is in non-uniform $\text{AC}^0$.  

### 3.2 Diffusion limited aggregation

Diffusion limited aggregation \footnote{This is not precise as delimiters are also used in the encoding to make decoding easier.} is a cluster growth model where new occupied sites are added to the growing cluster one at a time. Here we illustrate DLA for a two dimensional lattice with growth initiated along a line. A random walker is started at a random position along the top edge of an $L \times L$ square lattice. The walker moves until it is a nearest neighbor of an existing occupied site at which point it joins the cluster. Initially, the bottom edge of the lattice is considered occupied. If a walk fails to join the cluster, hits the top boundary of the lattice or is unable to move (goes off the lattice or encounters a site that is occupied in its first move), it is discarded. A new
random walk is started as soon as the previous walk has joined the cluster or been discarded; the process continues until a cluster of the desired size is grown.

A natural decision problem associated with the dynamics of diffusion limited aggregation is defined below.

**Diffusion limited aggregation** (dimension $d$)

**Given:** Three positive integers $L$, $M_1$ and $M_2$, a designated site $s$ expressed in unary with $|s| = L^d$ and a list of random bits specifying $M_1$ walk trajectories each of length $M_2$ defined by a starting point on the top edge of the lattice together with a list of directions of motion (e.g. N, S, E and W for two dimensions).

**Problem:** Is site $s$ occupied by the aggregation process?

The proof that DLA is P-complete proceeds by a reduction from a variant of the planar NOR circuit value problem. The reduction has a similar flavor to the proof that a closely related fluid invasion problem is P-complete [5], although there seems to be no way to make use of that proof directly.

**Theorem 3.2** Diffusion limited aggregation is P-complete.

**Proof sketch:** The idea is to prescribe a sequence of walks capable of carrying out the evaluation of a modified (but still P-complete) version of the planar NOR circuit value problem. In this version of CVP the NOR gates have a fan-in and fan-out of two. We also allow single input OR gates with fan-out restricted to at most two. The circuit encoding requires that the gates are numbered in topological order. The encoding specifies a planar layout of the circuit with gates being located at grid points. Finally, the circuit is required to be synchronous. That is, each gate receives its inputs only from gates on the immediately preceding level. Gates at level one are the only gates that are allowed to have direct circuit inputs. It can be shown that this version of CVP is P-complete.

The walks to simulate the circuit are chosen so that the cluster grows along linear paths of sites and bonds that play the role of wires connecting gates. A wire carries the value TRUE if the cluster grows along it. Wires that remain unoccupied carry the value FALSE. The gates themselves are represented by locations where wires meet and several parts of the growing cluster interact. Below we describe how logical values are propagated along wires and how NOR and OR gates are implemented.
Logical values are propagated as follows. Each wire is realized by a preassigned sequence of walks. These walks move to successive locations along the wire. Each walk moves to its assigned site along the wire and, if the value of the wire is true, it sticks there. Each walk reverses its path after reaching its assigned site. In this way if the wire carries the value false, the walk returns to the upper boundary and is discarded. For example, the first walk creating the output wire for the gate shown in Fig. 3 arrives at site ‘d’ from above. If ‘c’ is occupied this walk sticks at ‘d’ and the cluster begins to grow along the output wire. If ‘c’ is not occupied, the walk turns around and retraces its steps back to the upper boundary where it is discarded. Thus the cluster grows along the output if site ‘c’ is occupied. It is straightforward to have the output wire split into two separate wires. These then become the inputs to other gates. Note, a larger fan-out could be supported, however, the details of the proof become more involved.

A nor gadget is shown in Fig. 3. The solid lines and circles represent the input wires, the output wire and the ‘power’ wire. The role of the power wire is to provide a growing tip for the output if needed. The dashed line represents a walk that will stick at one of the three open circles labeled ‘a,’ ‘b’ or ‘c.’ The dashed walk evaluates the gate and so must not occur until the input and power wires have been grown to completion. Suppose that input 1 is true so that the corresponding segment of wire is occupied. Then the trajectory sticks at ‘a.’ If input 1 is false but input 2 is true, the dashed walk sticks at ‘b.’ Finally, if both inputs are false then the walk sticks at ‘c.’ The occupancy of site ‘c’ records the output of the gate.

Fig. 4 shows the implementation of a single input or gate. Effectively, it shows how to cross the power wire (running toward the left) over a logical wire (running vertically). First, the logical wire is grown to site ‘1.’ The walk represented by the dotted line on the right sticks at ‘a’ if the logical wire is true and sticks at ‘b’ if the logical wire is false. Similarly, the walk represented by the dotted line on the left sticks at ‘c’ if the logical wire is true and sticks at ‘d’ if the logical wire is false. Finally, the logical wire may continue to grow vertically and the power wire may continue to grow to the left without interfering with one another. In this way a single input or is simulated.

For three and higher dimensional lattices each gate may be separately supplied with its own power wire. For example, each gate has a ‘column’ dedicated to its power wire. When the wire reaches the appropriate height, it is routed horizontally to the desired gate. For two dimensional DLA there is an additional complication in arranging to have the power wire arrive
at each gate without interfering with the wires that carry truth values. To accomplish this we use a single power wire for the entire circuit. It is ‘snaked’ through the gates level by level. See the power wire in the example shown in Fig. 3; this example is discussed further later.

The discussion above shows how to pass the power wire through an OR gate. It is also necessary to have the power wire cross through a NOR gate. This can be done as shown in Fig. 5. In this figure the path of the power wire is numbered and the walks that bring particles to the wire are shown as dashed lines. Recall that exactly one of the sites ‘a,’ ‘b’ or ‘c’ is occupied during the evaluation of the gate. If ‘c’ is occupied, the power wire grows along the full path ‘1–8.’ If instead ‘b’ is occupied, sites ‘1’ and ‘2’ are skipped and growth starts at ‘3.’ In this case the walks that go to ‘1’ and ‘2’ turn around there and return to the upper boundary where they are absorbed. Finally, if ‘a’ is occupied then growth of the power wire starts at ‘6.’ Thus after helping with the evaluation of a NOR gate, the power wire may be passed through.

A single power wire traverses all the gates in the sequence in which they would be evaluated in topological order. This requires that the gates be arranged in levels as shown in the example in Fig. 6. The thick lines are circuit wires, the filled circles are NOR gates and the open circles are single input OR gates. The power wire traverses the gates one level at a time. Gates are evaluated from bottom to top and level by level along the path of the power wire. The lower edge of the lattice is used as a source for TRUE inputs to gates. At each gate the power and input wires arrive first, then the gate is evaluated and finally the power wire is continued to the next gate. After an entire level has been simulated, outputs are grown to the succeeding level. The routing between levels can be accomplished by ‘spreading’ the circuit out on the lattice and then allocating a couple of horizontal channels to each output of a gate. An output will be grown upward to its designated channel, grown horizontally underneath its appropriate gate and then grown upward to serve as an input. The planarity of the original circuit guarantees that there will be no interference of walks during this routing.

The reduction described above shows that the special instance of CVP we constructed is faithfully evaluated by the growth of the DLA cluster. Furthermore, it is an NC reduction. The key point is that the choice of paths for the walks is independent of the evaluation of the circuit. The full layout of the walks is given globally by the planar layout of the original circuit as outlined above and locally by Figs. 3 through 5. All calculations required to compute these walks can be performed in NC. □
3.3 Metropolis dynamics for the Ising model

Configurations of the Ising model are defined by spin variables, $\sigma_i$, on a lattice where each spin may take the value $-1$ or $+1$. The conventional way to obtain equilibrium states of the Ising model is via the Metropolis Monte Carlo algorithm. At each step of the algorithm a site $i$ is chosen at random and the energy change, $\Delta E_i$, for flipping the spin at this site is computed. The energy change is given by

$$\Delta E_i = 2J\sigma_i \sum_{<i,j>} \sigma_j$$

where the summation is over nearest neighbors of site $i$ and $J$ is the coupling energy. If $\Delta E_i \leq 0$ the spin is ‘flipped’ ($\sigma_i \rightarrow -\sigma_i$), whereas if $\Delta E_i > 0$ the spin is flipped with probability $e^{-\Delta E_i/T}$, where $T$ is the temperature. After this procedure has been iterated sufficiently many times, the resulting probability distribution for the spin configurations is close to the equilibrium state.

Metropolis dynamics is governed by a random list of sites and, for each site in the list, a random number $x_i$ with $0 \leq x_i < 1$ such that the site is flipped if $x_i \leq e^{-\Delta E_i/T}$. We can define the following natural decision problem for Metropolis dynamics.

**Metropolis dynamics** (dimension $d$)

**Given:** A positive integer $L$, an initial configuration of $L^d$ spins $\{\sigma_i\}$ with $\sigma_i \in \{-1, +1\}$, a temperature variable $Q = e^{-4J/T}$ where $Q$ is expressed as a $b$-bit binary number, a designated site $s$ expressed in unary with $|s| = L^d$, a list of $M$ sites and a list of $M$ random numbers $x_i$ with $0 \leq x_i < 1$ expressed as a $db$-bit number.

**Problem:** Is $\sigma_s = +1$ after running the Metropolis algorithm?

Given the random numbers $x_i$ we can assign flip variables, $g_i \in \{0, \ldots, d\}$, to each site $i$. For example, in three dimensions the flip variables are defined by the inequalities

- $g_i = 0$ if $0 \leq x_i \leq Q^3$,
- $g_i = 1$ if $Q^3 < x_i \leq Q^2$,
- $g_i = 2$ if $Q^2 < x_i \leq Q$ and
- $g_i = 3$ if $Q < x_i < 1$.

If a site $k$ is chosen for a possible flip at step $i$ and $\Delta E_k/4J \leq 3 - g_i$, then the flip is carried out; otherwise, the spin is not changed. In other words, a
Theorem 3.3 Metropolis dynamics is P-complete for $d$ greater than or equal to 3.

Proof sketch: The Metropolis problem is proved P-complete by a reduction from monotone CVP. The circuit is first `embedded’ in a three dimensional lattice. The AND and OR gates are represented by sites and wires connecting gates by chains of sites and bonds. For a circuit having $N$ edges, it can be shown that such an embedding may be carried out in NC. Initially, all spins on the lattice are $-1$. Logical values are represented by spin values with $+1$ ($-1$) meaning true (false). Logical values are propagated along wires by the following device: spins along the wire are sequentially chosen for flipping and assigned the flip variable 1. If the predecessor spin along the wire is $+1$, the current spin will flip to $+1$ but if the predecessor spin along the wire is $-1$, the flip is rejected. Thus, once initiated, logical values propagate along wires. Wires must always be separated by one or more lattice spacings except where they meet at gates. Sites representing gates have two input wires and two output wires. After all sites along the input wires have taken their logical values, the gate is ready for evaluation. Gates are assigned the flip variable 2 (1) for an AND (OR) gate. Thus, if at least one input is true an OR gate registers true, while both inputs must be true for an AND gate to register true. This NC reduction shows that we can simulate an arbitrary monotone circuit using Metropolis dynamics in three or more dimensions. Therefore, the Metropolis dynamics problem is P-complete.

Note that the planar monotone circuit value problem is in NC, see [14] for a list of references regarding this problem. So, our proof does not show that the two-dimensional Metropolis problem is P-complete. We have been unsuccessful in our attempts to implement a NOT gate within the framework of Metropolis dynamics.

The construction in Theorem 3.3 relies on a special ordering of the sites chosen for flipping. However, we can easily extend the proof to updating schemes in which sweeps through the lattice are performed in a fixed order. For example, consider the case of parallel updating where first the odd sublattice is flipped all at once and then the even sublattice. The problem...
statement is slightly different here since now at each time step flip variables are assigned to half the sites in the lattice. It is easy to keep sites inactive by assigning them flip variables. Sites are assigned flip variables 1 or 2 as in the above construction at the times they are to be evaluated.

### 3.4 Cluster dynamics for the Ising model

Cluster flipping algorithms due to Wolff [28] and Swendsen and Wang [29] are very efficient methods for generating equilibrium states of the Ising model near criticality. In this section we show that natural decision problems associated with the Wolff and Swendsen-Wang algorithms are P-complete.

We illustrate the Wolff algorithm on an $L \times L$ square lattice. The starting point is a configuration of spins, \{\sigma_j\}. Next the bonds of the lattice are independently occupied with probability $p$ as in bond percolation. The occupation parameter is related to the temperature, $T$, according to $p = 1 - Q$ with $Q = e^{-2J/T}$ and $J$ the coupling energy between neighboring spins. A site $u$ on the lattice is chosen at random and a cluster is grown from this site. A site $v$ is in the cluster grown from $u$ if there is a path from $u$ to $v$ such that all the bonds along the path are occupied and all the spins along the path including $\sigma_v$ are equal to $\sigma_u$. The cluster of spins defined in this way is ‘flipped’ ($\sigma \rightarrow -\sigma$ for each $\sigma$ in the cluster) which yields a new spin configuration. The procedure is iterated $M$ times. If the temperature $T$ is chosen to be the critical temperature and if $M$ is sufficiently large the final configuration of spins is close to the equilibrium Ising critical point. At the Ising critical temperature, the clusters defined by the algorithm are critical droplets [30, 31, 32] with Hausdorff dimension, $D_H$ equal to $15/8$.

The Swendsen-Wang algorithm is very similar to the Wolff algorithm except that in each step of the algorithm all connected clusters defined by the occupied bonds are identified. All sites of each cluster are assigned the same spin value. The spin values for each cluster are determined independently by a fair coin toss.

For each iteration of the Wolff or Swendsen-Wang algorithm, every bond of the lattice is occupied with probability $p$ equal to $1 - Q$. To implement this we utilize random numbers $x_{ij}$ with $0 \leq x_{ij} < 1$ for each nearest neighbor pair $(ij)$. The bond $(ij)$ is occupied if $x_{ij}$ is greater than $Q$. At each time step a cluster is grown from the starting point according to the occupation variables and the current spin configuration as described above. This cluster is flipped and the procedure repeated $M$ times. We can define the following natural problem based on Wolff dynamics.
Wolff dynamics (dimension \(d\))

**Given:** A positive integer \(L\), an initial configuration of \(L^d\) spins \(\{\sigma_i\}\) with \(\sigma_i \in \{-1, +1\}\), a temperature variable \(Q = e^{-2J/T}\) where \(Q\) is expressed as a \(b\)-bit binary number, a designated site \(s\) expressed in unary with \(|s| = L^d\), a list of \(M\) sites and \(dML^d\) random numbers \(x_{ij}\) with \(0 \leq x_{ij} < 1\) expressed as a \(b\)-bit number.

**Problem:** Is \(\sigma_s = +1\) after running the Wolff algorithm?

Given the random numbers \(x_{ij}\) we can assign bond occupation variables \(b_{ij}\) such that \(b_{ij} = 0\) if \(x_{ij} \leq Q\) and \(b_{ij} = 1\) otherwise. Bonds are counted as occupied if \(b_{ij} = 1\). It is clear that the Wolff decision problem can be NC reduced to a version in which the random input is given as the \(b_{ij}\) instead of the \(x_{ij}\). It is this version that we show is \(P\)-complete using a reduction from the planar NOR circuit value problem.

**Theorem 3.4** Wolff dynamics is \(P\)-complete.

**Proof idea:** The reduction is best illustrated by a simple example. We sketch it for the case \(d\) equals two. Consider the planar circuit shown in Fig. 7 with three inputs and three NOR gates. The evaluation of this circuit can be reduced to the Wolff problem shown in Fig. 8. The lower case letters indicate occupied bonds and time steps that are arranged in alphabetical order. All bonds labeled ‘a’ are occupied only during step 1, all bonds labeled ‘b’ are occupied only during step 2 and so on. Bonds that are not explicitly shown are never occupied. All spins on the lattice originally have the value +1 except those that are labeled FALSE. A +1 spin represents TRUE and vice versa. Numbers label initiation sites for cluster growth. Cluster growth is initiated at gates and logical constants. Site ‘1’ initiates the cluster growth at time step 1 and represents the TRUE input to the circuit; site ‘2’ initiates growth at time step ‘2’ and so on. The first cluster propagates as far as site ‘4’ and both sites ‘1’ and ‘4’ (and the intermediate site) are flipped to −1. Site ‘2’ initiates the next cluster which does not propagate. Site ‘2’ is flipped to +1. The cluster initiated at ‘3’ does not propagate so that at the beginning of time step 4 site ‘4’ is in the FALSE state; the first gate has been properly evaluated. At time step 4 site ‘4’ flips but nothing else happens; the second gate properly evaluates to TRUE. At time step 5 a cluster propagates from site ‘5’ to the output gate that is flipped to −1. The output of the circuit is FALSE as it should be.
More generally, a NOR gate is represented by spins, and wires connecting gates are represented by paths of bonds and spins. All of the bonds in a wire are occupied at the time step during which the wire transmits its logical value. If the logical value is TRUE, a cluster of up spins is propagated along the wire and the output end of the wire is flipped to FALSE if this has not yet occurred. A gate transmits its value by initiation of a cluster and the output of a gate can be read off as soon as all of its predecessors have transmitted their values. Note that a fan-out higher than two may easily be supported. Sites representing gates and logical constants must not be nearest neighbors. It is clear that the reduction of the circuit to Wolff dynamics can be carried out locally and is an NC reduction. Since planar NOR CVP is P-complete, so is the Wolff dynamics problem.

Next we turn our attention to a natural decision problem associated with the Swendsen-Wang algorithm. The problem statement requires random ‘bits,’ \( c_i \) equal to \( \pm 1 \), to be used to determine the spins in the clusters. Sites are given a conventional ordering. Connected clusters defined by \( Q \) and the variables \( x_{ij} \) are labeled by the lowest ordered site, \( l \), in the cluster and all the spins in the cluster are assigned the value \( c_l \).

**Swendsen-Wang dynamics (dimension \( d \))**

**Given:** A positive integer \( L \), an initial configuration of \( L^d \) spins \( \{\sigma_i\} \) with \( \sigma_i \in \{-1, +1\} \), a temperature variable \( Q = e^{-2J/T} \) where \( Q \) is expressed as a \( b \)-bit binary number, a designated site \( s \) expressed in unary with \( |s| = L^d \), a number of iterations \( M \), a list of \( dML^d \) random numbers \( x_{ij} \) with \( 0 \leq x_{ij} < 1 \) expressed as a \( b \)-bit number and a list of \( ML^d \) random bits, \( c_i \).

**Problem:** Is \( \sigma_s = +1 \) after running the Swendsen-Wang algorithm?

**Theorem 3.5** Swendsen-Wang dynamics is P-complete.

**Proof hint:** The proof is similar to that for the Wolff problem and consists of a reduction from planar NOR CVP. Here again a value of \( d \) equal to two suffices for the reduction. Consider a two-dimensional lattice and suppose that the conventional ordering of sites on the lattice is from left to right and then from bottom to top. The occupation variables \( b_{ij} \) are chosen the same as for the reduction to the Wolff problem. The cluster spin variables, \( c_i \), are \(-1\) for gates and TRUE inputs at the time they transmit their values and \(+1\) for all other sites and all other times.

For both Wolff and Swendsen-Wang dynamics, a single iteration of the algorithm can be accomplished in poly-logarithmic parallel time using a
polynomial number of processors. This is because the most complex step is
the identification of a connected component(s), which can be carried out by
a standard NC algorithm [33]. More specifically, if the number of iterations
$M$ is set to a constant in the statement of either the Wolff problem or the
Swendsen-Wang problem, the resulting decision problem is in NC. For the
Metropolis algorithm, an even stronger result holds. If $M$ equals a constant,
the Metropolis decision problem is in $\text{AC}^0$. These conclusions are not in
conflict with the P-completeness proofs that rely on setting $M$ comparable
to the size of the Boolean circuit being simulated. The P-completeness
results show that a polynomial (in the system size) number of iterations
of these algorithms cannot be compressed into poly-logarithmic number of
parallel steps, unless NC = P.

4 Discussion

4.1 Summary of results

We have studied the computational complexity of natural decision problems
associated with several models in statistical physics. Our results can be
summarized as follows:

1. Mandelbrot percolation is in (non-uniform) AC$^0$ (Theorem 3.1).
2. Diffusion limited aggregation is P-complete (Theorem 3.2).
3. Metropolis dynamics for the Ising model is P-complete (Theorem 3.3).
4. Wolff dynamics for the Ising model is P-complete (Theorem 3.4).
5. Swendsen-Wang dynamics for the Ising model is P-complete (Theo-
rem 3.5).

4.2 Scope of the results

It is important to understand the limitations of the P-completeness results
for DLA and the variants of Ising dynamics. Suppose we accept the hy-
pothesis from complexity theory that NC $\neq$ P. In this case the particular
dynamics discussed here for generating DLA clusters or equilibrium Ising
configurations are inherently sequential and cannot be efficiently simulated
in parallel. There are other ways to generate (approximate) equilibrium
states of the Ising model or DLA clusters; our results do not imply that
these ways are associated with \textbf{P}-complete problems. However, it was previously shown \cite{5} that a second method of producing DLA clusters is also \textbf{P}-complete. It seems plausible that there are no poly-logarithmic time methods for sampling the DLA distribution. On the other hand, the jury remains out on whether it is possible to sample from an approximation to the equilibrium critical distribution for spin models in poly-logarithmic time. It would be of great interest to obtain results on the difficulty of sampling physically interesting distributions.

A second limitation of the \textbf{P}-completeness statements is that they are worst case rather than average case results. For example, assuming that \textbf{NC} \neq \textbf{P}, we know that there exist instances of the DLA problem that cannot be solved in poly-logarithmic time although we do not know whether these ‘hard’ instances are typical or very rare. Indeed, the instances used in the \textbf{P}-completeness proofs are atypical. If the ‘hard’ instances are sufficiently rare, we may be able to sample the distribution in poly-logarithmic time on average. The theory of average case complexity \cite{34,35} addresses questions of this kind. Unfortunately, it is not easy to see how to apply this theory to the present problems.

\section*{4.3 Parallel complexity and critical slowing down}

Away from a critical point, the equilibration time of real systems without macroscopic inhomogeneities is independent of system size. Similarly, the Metropolis algorithm can generate good approximations to equilibrium configurations of the Ising model away from the critical point in constant parallel time since each sweep can be done in constant time and the number of sweeps is independent of the system size. The associated decision problem is in \textbf{AC}^0.

Equilibration of many real systems becomes increasingly slow near critical points. Typically the equilibration time at a critical point scales as \( L^z \), where \( L \) is the system size and \( z \) is the dynamic exponent. This phenomena, known as critical slowing down, also afflicts most Monte Carlo methods used to sample spin configurations at critical points. The dynamic exponent \( z \) is customarily defined for Monte Carlo dynamics if relaxation to equilibrium requires flipping \( L^{z+d} \) spins. It is often said that an algorithm suffers no critical slowing down if \( z = 0 \) (with possible logarithmic corrections). This is not a satisfactory general definition of ‘absence of critical slowing down.’ For example, imagine an algorithm for which each spin is flipped only once but an enormous computation is required to decide whether or not to effect
the flip. Alternatively, one might propose that ‘absence of critical slowing down’ means that the sequential time (computational work) is $o(L^{d+\epsilon})$ for any $\epsilon > 0$. This definition is both machine dependent and unnecessarily stringent.

We propose that ‘absence of critical slowing down’ should be identified with the class NC. A sampling method suffers no critical slowing down if it can be run in poly-logarithmic time on a P-RAM with polynomially many processors. This definition is, for the most part, in agreement with the $z = 0$ definition. If $z > 0$ for the Monte Carlo methods studied here, the P-completeness results show that there is critical slowing down according to the new definition. On the other hand, if $z = 0$ for either the Metropolis or Swendsen-Wang algorithms, there is no critical slowing down since a single sweep through the lattice for either of these algorithms can be done efficiently in parallel and $z = 0$ implies a poly-logarithmic number of sweeps. In contrast, the Wolff algorithm suffers critical slowing down even for $z = 0$ according to the new definition. The reason is that the average size of Wolff clusters scales as $L^{\gamma/\nu}$, where $\gamma$ is the susceptibility exponent and $\nu$ the correlation length exponent. Thus, even if $z = 0$ one typically requires $L^{d-\gamma/\nu}$ iterations of the algorithm to reach equilibrium. The P-completeness result shows that carrying out these iterations can almost certainly not be done in poly-logarithmic time using a polynomial number of processors.

4.4 Final remarks

In this and two previous papers we have investigated the parallel computational complexity of a variety of models in statistical physics. We have claimed that parallel complexity provides statistical physics with a robust and sharply defined measure that reflects some of our more intuitive notions of complexity. We have classified a wide variety of models into three broad classes: those that require constant parallel time to simulate, those that require poly-logarithmic time and those that require polynomial time. In each case we allow a polynomial number of processors. Even among models that generate random fractal patterns, we find representatives in each of these classes. Comparisons between members of different classes reveal that models in the higher classes generally pose a more difficult theoretical challenge. It would be extremely interesting to find more precise correlations between computational complexity and the quantities conventionally studied in statistical physics.
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