

Parallel computation using active self-assembly*

Moya Chen

Doris Xin

Damien Woods

California Institute of Technology

Abstract

We study the computational complexity of the recently proposed nubot model of molecular-scale self-assembly. The model generalises asynchronous cellular automata to have non-local movement where large assemblies of molecules can be pushed and pulled around, analogous to millions of molecular motors in animal muscle effecting the rapid movement of macroscale arms and legs. We show that the nubot model is capable of simulating Boolean circuits of polylogarithmic depth and polynomial size, in only polylogarithmic expected time. In computational complexity terms, we show that any problem from the complexity class NC is solvable in polylogarithmic expected time and polynomial workspace using nubots.

Along the way, we give fast parallel nubot algorithms for a number of problems including line growth, sorting, Boolean matrix multiplication and space-bounded Turing machine simulation, all using a constant number of nubot states (monomer types). Circuit depth is a well-studied notion of parallel time, and our result implies that the nubot model is a highly parallel model of computation in a formal sense. Asynchronous cellular automata are not capable of this parallelism, and our result shows that adding a rigid-body movement primitive to such a model, to get the nubot model, drastically increases parallel processing abilities.

1 Introduction

We study the theory of molecular self-assembly, working within the recently-introduced *nubot* model by Woods, Chen, Goodfriend, Dabby, Winfree and Yin [51]. Do we really need another new model of self-assembly? Consider the biological process of embryonic development: a single cell growing into an organism of astounding complexity. Throughout this active, fast and robust process there is growth and movement. For example, at an early stage in the development of the fruit fly *Drosophila*, the embryo contains approximately 6,000 large cells arranged on its ellipsoid-shaped surface. Then, in just four minutes, the embryo rapidly changes shape to become invaginated, creating a large structure that becomes the mesoderm, and ultimately muscle. How does this fast rearrangement occur? A large fraction of these cells undergo a rapid, synchronised and highly parallel rearrangement of their internal structure where, in each cell, one end of the cell contracts and the other end expands. This is achieved by a mechanism that seems to crucially involve thousands of molecular-scale myosin motors pulling and pushing the cellular cytoskeleton to quickly effect this rearrangement [30]. At an abstract level one can imagine this as being analogous to how millions of molecular motors in a

*mpchen@caltech.edu, dorx@alumni.caltech.edu, woods@caltech.edu. Supported by National Science Foundation grants CCF-1219274, 0832824 (The Molecular Programming Project), and CCF-1162589. Preliminary version appeared at The 19th International Conference on DNA Computing and Molecular Programming (DNA 19).

muscle, each taking a tiny step but acting in a highly parallel fashion, effect rapid long-distance muscle contraction. This rapid parallel movement, combined with the constraint of a fixed cellular volume, as well as variations in the elasticity properties of the cell membrane, can explain this key step in embryonic morphogenesis. Indeed, molecular motors that together, in parallel, produce macro-scale movement are a ubiquitous phenomenon in biology.

We wish to understand, at a high level of abstraction, the ultimate limitations and capabilities of such molecular scale rearrangement and growth. We do this by studying the computational power of a theoretical model that includes these capabilities. As a first step towards such understanding, we show in this paper that large numbers of tiny motors (that can each pull or push a tiny amount) coupled with local state changes on a grid, are sufficient to quickly solve inherently parallelisable problems. This result, described formally below in Section 1.2, demonstrates that the nubot model is a highly parallel computer in a computational complexity-theoretic sense.

Another motivation, and potential test-bed for our theoretical model and results, is the fabrication of active molecular-scale structures. Examples include DNA-based walkers, DNA origami that reconfigure, and simple structures called molecular motors [53] that transition between a small number of discrete states (see [51] for references). In these systems the interplay between structure and dynamics leads to behaviours and capabilities that are not seen in static structures, nor in other unstructured but active, well-mixed chemical reaction network type systems. Our theoretical results here, and those in [51], provide a sound basis to motivate the experimental investigation of large-scale active DNA nanostructures.

There are a number of theoretical models of molecular-scale algorithmic self-assembly processes [39]. For example, the abstract Tile Assembly Model, where individual square DNA tiles attach to a growing assembly lattice one at a time [48, 42, 16], the two-handed (hierarchical) model where large multi-tile assemblies come together [1, 7, 11, 14], and the signal tile model where DNA origami tiles that form an “active” lattice with DNA strand displacement signals running along them [23, 36, 37]. Other models enable one to program tile geometry [12, 18], temperature [1, 25, 46], concentration [5, 8, 15, 26], mixing stages [11, 13] and connectivity/flexibility [24].

The well-studied abstract Tile Assembly Model [48] is an asynchronous, and nondeterministic, cellular automaton with the restriction that state changes are irreversible and happen only along a crystal-like growth frontier. The nubot model is a generalisation of an asynchronous and non-deterministic cellular automaton, where the generalisation is that we have a non-local movement primitive. Since the nubot model is intended to be a model of molecular-scale phenomena it ignores friction and gravity, allows for the creation/destruction of monomers (we assume an invisible “fuel” source) and has a notion of random uncontrolled motion (called agitation, but not used in this paper). Instances of the model evolve as continuous time Markov processes, and time is modelled as in stochastic chemical kinetics [19, 45]. The nubot style of rigid-body movement is analogous to that seen in reconfigurable robotics [6, 43, 32], and indeed results in these robotics models show that non-local movement can be used to effect fast global reconfiguration [4, 3, 41]. The nubot model includes features seen in cellular automata, Lindenmayer systems [40] and graph grammars [28]. See [51] for a more detailed comparison of the similarities and differences with these models.

1.1 Previous work on active self-assembly with movement

Previous work on the nubot model [51] showed that it is capable of building large shapes and patterns exponentially quickly: e.g. lines and squares in time logarithmic in their size. The same paper goes on to describe a general scheme to build arbitrary computable (connected, 2D) size- n

shapes in time and number of monomer states (types) that are polylogarithmic in n , plus the time and states required for Turing machine simulation due to the inherent algorithmic complexity of the shape. Furthermore, 2D patterns with at most n coloured pixels, where the colour choice for each pixel is computable in time $\log^{O(1)} n$ (i.e. polynomial in the length of the binary description of pixel indices), are nubot-computable in time and number of monomer types polylogarithmic in n [51]. The latter result is achieved without going outside the pattern boundary and in a completely asynchronous fashion. These results show that the nubot model is capable of parallelism not seen in many other models of self-assembly. The goal of the present paper is to characterise the kind of parallelism seen in the nubot model by formally relating it to the computational complexity of classical decision problems.

Dabby and Chen [10] study a 1D model, where monomers insert between, and push apart, other monomers. Their model is closely related to a 1D restriction of the nubot model without state changes, and they build length n lines in $O(\log^3 n)$ expected time and $O(\log^2 n)$ monomer types. They also show that the set of 1D polymers produced by any instance of their model is a context-free language, and give a design for implementation with DNA molecules. Malchik and Winslow [29] first show that any context-free language can be expressed as an instance of this model, and then give an asymptotically tight bound of $2^{\Theta(k^{3/2})}$ on the length of polymers produced using k monomer types (in merely $O(\log^{5/3} n)$ expected time), thus characterising two aspects of the model.

1.2 Main result

In the nubot model a program is specified as a finite set of nubot rules \mathcal{N} and is said to decide a language $L \subseteq \{0, 1\}^*$ if, beginning with a word $x \in \{0, 1\}^*$ encoded as a sequence of $|x|$ “binary monomers”, the system eventually reaches a configuration containing exactly the 1 monomer if $x \in L$, and 0 otherwise. Let NC denote the (well-known) class of problems solved by uniform polylogarithmic depth and polynomial size Boolean circuits.¹ Our main result is stated as follows.

Theorem 1. *For each language $L \in \text{NC}$, there is a set of nubot rules \mathcal{N}_L that decides L in polylogarithmic expected time, constant number of monomer states, and polynomial space in the input string length. Moreover, for $i \geq 1$, NC^i is contained in the class of languages decided by nubots running in $O(\log^{i+3} n)$ expected time, $O(1)$ monomer states, and polynomial space in input length n .*

NC problems are solved by circuits of shallow depth, hence they can be thought of as those problems that can be solved on a highly parallel architecture (simply run each layer of the circuit on a bunch of parallel processors, after polylogarithmic parallel steps we are done). NC is contained in P—problems solved by polynomial time Turing machines—and this follows from the fact that NC circuits are of polynomial size. Problems in NC, and the analogous function class, include sorting, Boolean matrix multiplication, various kinds of maze solving and graph reachability, and integer addition, multiplication and division. Besides its circuit depth definition, NC has been characterised by a large number of other parallel models of computation including parallel random access machines, vector machines, and optical computers [20, 52, 49]. It is widely conjectured, but unproven, that NC is strictly contained in P. In particular, problems complete for P (such as Turing machine and cellular automata [35] prediction, context-free grammar membership and many others [20]) are believed to be “inherently sequential”—it is conjectured that these problems are not solvable by parallel computers that run for polylogarithmic time on a polynomial number of processors [20, 9].

¹NC, or Nick’s class, is named after Nicholas Pippenger.

Thus our main result gives a formal sense in which the nubot model is highly parallel: for any highly parallelisable (NC) problem our proof gives a nubot algorithm to efficiently solve it in only polylogarithmic expected time and constant states. This stands in contrast to sequential machines like Turing machines, that cannot read all of an n -bit input string in polylogarithmic time, and “somewhat parallel” models like cellular automata and the abstract Tile Assembly Model, which can not have all of n bits influence a single bit decision in polylogarithmic time [27]. Thus, adding a movement primitive to an asynchronous non-deterministic cellular automation, as in the nubot model, drastically increases its parallel processing abilities.

We finish this discussion on a technical remark. Previous results [51] on the nubot model were of the form: for each $n \in \mathbb{N}$ there is a set of nubot rules \mathcal{N}_n (i.e. the number of rules is a function of n) to carry out some task parameterised by n (examples: quickly grow a line of length n or an $n \times n$ square, or grow some complicated computable pattern or shape whose size is parameterised by n , etc.). For each problem in NC our main result here gives a *single* set of rules (i.e. of constant size), that works for all problem instances.

1.3 Overview of results and paper structure

Section 1 contains the statement of our main result, the overall proof structure and some future work directions. Section 2 gives the full definition of the nubots model and relevant complexity classes. Section 3 serves as an introduction to the nubots model by giving a simple nubots algorithm to double the length of a length- n line in $O(\log n)$ expected time, we suggest the reader begins there.

1.3.1 New synchronization and line growth algorithms

In Section 4 we describe a fast signalling method for nubots from [51], here called *shift synchronization*, and give a new variant on this called *lift synchronization*. These signalling mechanisms are used throughout our constructions as a method to quickly send a bit, 0 or 1, distance n in $O(\log n)$ expected time, with the choice of 0 or 1 being encoded by the use of shift or lift synchronization respectively.

The line growth algorithm given in [51] grows a line of length $n \in \mathbb{N}$ in $O(\log n)$ time, using $O(\log n)$ monomer states and starting from a single monomer on the grid. Section 5 gives a new line-growth algorithm that completes in $O(\log^2 n)$ time, using $O(1)$ monomer states and starting from $O(\log n)$ monomers on the grid. A key feature of our algorithm is that it uses only a constant number of states. This helps us achieve our main result, which requires a *single* set of nubots rules that accept any word from some, possibly infinite, NC language: as part of our circuit simulation we need to build longer and longer lines to simulate larger and larger circuits, all with a single set of nubots rules.

1.3.2 Parallel sorting, Boolean matrix multiplication & space bounded Turing machine simulation

Section 6 shows that the nubots model is capable of fast parallel sorting: n numbers can be sorted in expected time polylogarithmic in n . More precisely, n distinct natural numbers, taken from the set $\{0, 1, \dots, n-1\}$ when presented as n unordered “strings” of binary (0 or 1) monomers on the grid, can be sorted in increasing numerical order in expected time $O(\log^3 n)$, space $O(n \log n) \times O(n)$, and $O(1)$ monomer states. Our sorting routine is used throughout our main construction and is

inspired by mechanisms, such as gel electrophoresis, that sort via spatial organization based on physical quantities, such as mass and charge [33].

Section 7 shows that two $n \times n$ Boolean matrices can be multiplied in $O(\log^3 n)$ expected time, $O(n^4 \log n) \times O(n^2 \log n)$ space and $O(1)$ monomer states. This immediately implies that problems reducible to Boolean matrix multiplication, such as directed graph reachability and indeed any problem in the complexity class NL, of languages accepted by nondeterministic logarithmic space bounded Turing machines, can be solved in polylogarithmic expected time on nubots.

Indeed in Section 8.1 we go on to generalise this result by showing that any nondeterministic logarithmic space bounded Turing machine that computes a function (as opposed to merely deciding a language) can also be simulated in polylogarithmic space. This involves modifying the usual matrix multiplication method to keep track of the contents of the output tape of the Turing machine, and correctly reassembling the encoded tape contents on the 2D grid.

These results show that the model is capable of fast parallel solution of many problems, in particular all of those in NL. Recall that $NL \subseteq NC$, so we are not done yet. Indeed these techniques form part of our more general result: polylogarithmic expected time solution of problems in NC via efficient simulation of uniform Boolean circuits, as described next.

1.3.3 Proof overview of main result: Theorem 1

Let $L \in NC$, in other words, L is decidable by a logspace-uniform family \mathcal{C}_L of Boolean circuits of polylogarithmic depth and polynomial size. To prove Theorem 1, we show that for each such L there exists a finite set of nubots rules \mathcal{N}_L that decides L . L being in logspace-uniform NC implies that there is a deterministic logarithmic space (in input size) Turing machine \mathcal{M}_L such that $\mathcal{M}_L(1^n) = c_n$, where c_n is a description of the unique Boolean circuit in \mathcal{C}_L that has n input gates. Our initial nubots configuration consists of a length- n line of binary nubots monomers denoted $[\tilde{x}]$, that represents some input word $x \in \{0, 1\}^*$ (as described in Definition 3). From this we create (copy) another length- n line of monomers that encode the unary string 1^n to be given as input to a nubots simulator of \mathcal{M}_L . The rule set \mathcal{N}_L includes a description of \mathcal{M}_L , and the system first generates a circuit by simulating the computation of \mathcal{M}_L on input 1^n , which produces a nubots configuration (collection of monomers in a connected component) that represents the circuit c_n . The circuit is then simulated on input x . Both of these tasks present a number of challenges.

Circuit Generation. Logspace Turing machines run in at most polynomial time in their input length (otherwise they repeat a configuration), but here we wish to generate the circuit in merely polylogarithmic time. To achieve this, our simulation of \mathcal{M}_L works in a highly parallel fashion. This uses a number of techniques. First, in nubots, we implement the (known) trick of space-bounded Turing machine simulation by fast iterated matrix multiplication, which in turn is used to solve reachability on the directed graph of all possible configurations of the Turing machine. One of the main challenges here is to carry out matrix multiplication on the 2D grid sufficiently fast but without monomers unintentionally colliding with each other. Second, although iterated matrix multiplication is sufficient to simulate a Turing machine that decides a language, here we wish to simulate a Turing machine that computes a *function*. To do this, our parallel matrix multiplication algorithm keeps track of any symbols written to the output tape by both valid (reachable) and invalid (unreachable) configurations, and at the end deletes those symbols written by invalid configurations leaving the valid output symbols only. These valid output symbols are then arranged into the correct order by our fast parallel sorting routine. This results in a string of monomers that encodes the circuit c_n .

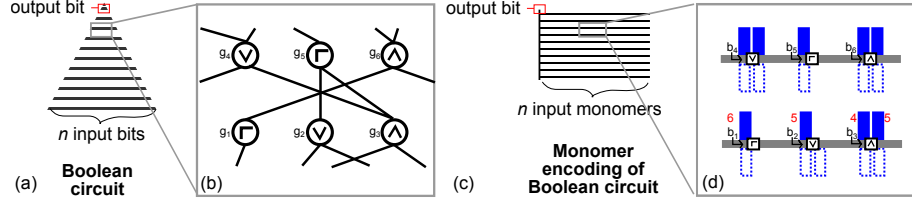


Figure 1: High-level overview of the encoding of a Boolean circuit as a nubots configuration (drawn on the square grid to save space). (a) Boolean circuit with (b) detailed zoom-in. (c) Nubots configuration encoding the circuit, with zoom-in shown in (d). A wire leading out of a gate in (b) has a destination gate number encoded in (d) as strips of $O(\log n)$ blue binary monomers (indices in red). After a gate computes some Boolean function (one of \vee , \wedge , \neg) the resulting bit is tagged onto the relevant blue strip of monomers that encode the destination addresses (red numbers). Circuits are not necessarily planar, so to handle wire crossovers these result bits are first sorted in parallel based on their wire address, and then pushed up to the next layer of gates.

These monomers then rearrange themselves in the plane, to lay out the circuit with each row of gates layered one on top of the next as shown in Figure 1. (Note that for convenience and to save space we sometimes draw figures on a square grid, although the nubots model is formally defined on the hexagonal grid.)

Circuit Simulation. As already described, the input x is encoded as the binary monomers $[\tilde{x}]$, and the entire circuit c_n is “grown” from $[\tilde{x}]$. The monomers $[\tilde{x}]$ now move to the first (bottom) row of the encoded circuit (Figure 1(c)) and position themselves so that each gate can “read” its 1 or 2 input bit monomers from $[\tilde{x}]$. After each gate computes a “result” bit, layer 1 “synchronizes” via a $O(\log n)$ expected time synchronization routine.

Next, we wish to send the “result” bits from layer 1 to layer 2. Circuits are not necessarily planar, so we need to handle wire crossings. We use our fast parallel sorting routine: the outputs from the first circuit layer are sorted, from left to right in increasing order, using their “to” address as a key. For example, a layer 1 result bit that is destined for gate 5 in layer 2 will be placed to the left of a layer 1 result bit that is destined for gate 6 in layer 2. Using this sorting routine, the blue “wire address” regions in the circuit (Figure 1(d)) are sorted in increasing order from left to right, then appropriately padded with empty space in between (using counters), and are passed up to the next level. Layer 1 then destroys itself. The entire circuit is simulated, level by level, from bottom to top, in this manner. After the “output gate” monomer computes its output bit it destroys itself, leaving a single monomer in state output_0 or output_1 . No more rules are applicable and so the system has halted with its answer. This completes the overview of the simulation.

This overview ignores many details. In particular the nubots model is asynchronous, that is, rule updates happen independently as discrete events in continuous time with no two events happening at the same time (as in stochastic chemical kinetics). The construction includes a large number of synchronization steps and signal passing to ensure that all parts of the construction are appropriately staged, but yet the construction is free to carry out many fast, asynchronous, parallel steps between these “sequential” synchronization steps.

1.4 Future work and open questions

The line growth algorithm in [51] runs in expected time $O(\log n)$, uses $O(\log n)$ states and space $O(n) \times O(1)$. In Section 5 we give another line growth algorithm that runs in expected time $O(\log^2 n)$, uses $O(1)$ states and space $O(n) \times O(1)$. Is there a line-growth algorithm that does better than $\text{time} \times \text{space} \times \text{states} = \Omega(n \log^2 n)$? To keep the game fair, the input should be a collection of monomers with $\text{space} \times \text{states} = O(\log n)$.

Theorem 1 gives a lower bound on the power of the nubot model. What are the best lower and the upper bounds on the power of confluent² polylogarithmic expected time nubots? One challenge involves finding better Turing machine space, or circuit depth, bounds on computing multiple applications of the movable set (see Section 2) on a polynomial size (or larger) nubot grid.

Synchronization is a signalling method we use to quickly send signals in a non-local fashion. In this paper it is used extensively to compose nubot algorithms. What conditions are necessary and sufficient for composition of arbitrary nubot algorithms that do not use synchronization? Theorem 7.1 in [51] shows that a wide class of patterns can be grown without synchronization, and its proof of this gives examples of composition without synchronization. It would be interesting to formalise this notion of composition in our distributed systems without the long-range fast signalling that synchronization gives.

Agitation is a kind of undirected, or random, movement that was defined for the nubot model in [51] and is intended to model a nanoscale environment where there are uncontrolled movements and turbulent fluid flows in all directions interacting with each monomer. Is it possible to simulate nubot-style movement using agitation? As motivation, note that every self-assembled molecular-scale structure was made under conditions where agitation is a dominant source of movement! Our question asks if we can *programmably exploit* this random molecular motion to build structures quicker than without it.

Is the nubot model intrinsically universal? More precisely, does there exist a set of monomer rules U , such that any nubot system \mathcal{N} can be simulated by “seeding” U with a suitable initial configuration? The notion of intrinsic universality is giving rise to interesting characterisations, and separations, in a variety of tile assembly models [16, 17, 12, 14, 31, 21, 22], for an overview see the survey [50]. Our hope would be that intrinsic universality, with its tight notion of simulation, could be used to tease apart the power of different notions of movement (for example to understand if nubot-style movement is weaker or stronger than other notions of movement).

Other open problems and further directions can be found in [51].

2 The nubot model and other definitions

In this section we formally define the nubot model. Figure 2 gives an overview of the model and rules, and Figure 3 gives an example of the movement rule. An example nubot construction for “line-doubling” is given in Section 3 which may aid the reader at this point. Let $\mathbb{N} = \{0, 1, 2, \dots\}$.

The model uses a two-dimensional triangular grid with a coordinate system using axes x and y as shown in Figure 2(a). A third axis, w , is defined as running through the origin and $\vec{w} = -\vec{x} + \vec{y} = (-1, 1)$, but we use only the x and y coordinates to define position. The *axial directions* $\mathcal{D} = \{\pm\vec{x}, \pm\vec{y}, \pm\vec{w}\}$ are the unit vectors along axes x, y, w . A grid point $\vec{p} \in \mathbb{Z}^2$ has the set of six *neighbours* $\{\vec{p} + \vec{u} \mid \vec{u} \in \mathcal{D}\}$. Let S be a finite set of monomer states. A nubot

²By confluent we mean a kind of determinism where the system (rules with the input) is assumed to always make a unique single terminal assembly.

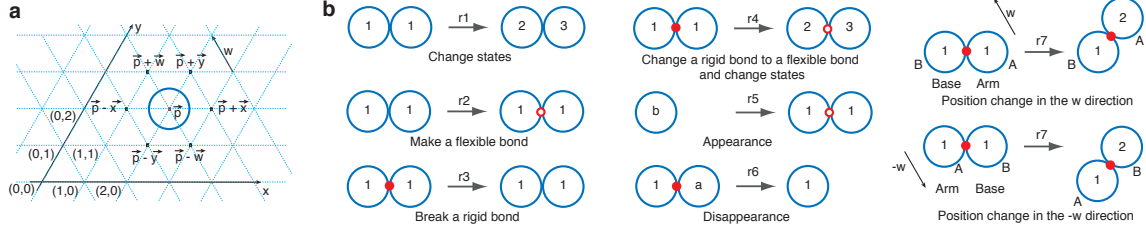


Figure 2: Overview of the nubot model. (a) A nubot configuration showing a single nubot monomer on the triangular grid. (b) Examples of nubot monomer rules. Rules r1-r6 are local cellular automaton-like rules, whereas r7 effects a non-local movement that may translate other monomers as shown in Figure 3. A flexible bond is depicted as an empty red circle and a rigid bond is depicted as a solid red disk.

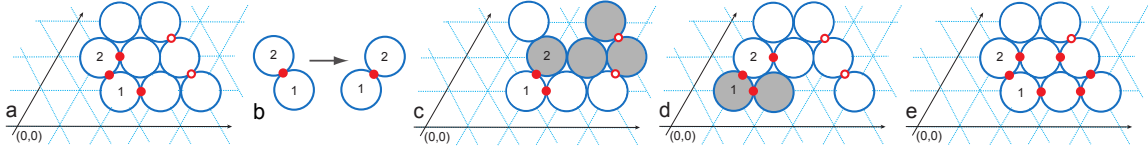


Figure 3: Movement rule. (a) Initial configuration. (b) Movement rule with one of two results depending on the (nondeterministic) choice of arm or base. (c) Result if the monomer with state 1 is the base, or (d) if the monomer with state 2 is the base. The shaded monomers are the moveable set. We can think of (c) as pushing and (d) as pulling. The affect on rigid (filled red disks), flexible (hollow red circles) and null bonds is shown. (e) A configuration for which the movement rule is blocked: movement of 1 or 2 would force the other to move hence the rule is not applicable.

monomer is a pair $X = (s_i, p(X))$ where $s_i \in S$ is a state and $p(X) \in \mathbb{Z}^2$ is a grid point. Two monomers on neighbouring grid points are either connected by a *flexible* or *rigid* bond, or else have no bond (called a *null* bond). Bonds are described in more detail below. A *configuration* C is a finite set of monomers along with the bonds between them.

One configuration *transitions* to another via the application of a single *rule*, $r = (s1, s2, b, \vec{u}) \rightarrow (s1', s2', b', \vec{u}')$ that acts on one or two monomers.³ The left and right sides of the arrow respectively represent the contents of two monomer positions before and after the application of rule r . Here $s1, s2 \in S \cup \{\text{empty}\}$ are monomer states where at most one of $s1, s2$ is *empty* (denotes lack of a monomer), $b \in \{\text{flexible}, \text{rigid}, \text{null}\}$ is the bond type between them, and $\vec{u} \in \mathcal{D}$ is the relative position of the $s2$ monomer to the $s1$ monomer. If either of $s1$ or $s2$ (respectively $s1'$ or $s2'$) is *empty* then b (respectively b') is null (monomers can not be bonded to empty space). The right is defined similarly, although there are some further restrictions on valid rules (involving \vec{u}') described below. A rule is only applicable in the orientation specified by \vec{u} , and so rules are not rotationally invariant.

³In reference [51] the nubot model includes “agitation”: each monomer is repeatedly subjected to random movements intended to model a nano-scale environment where there is Brownian motion, uncontrolled movements and turbulent fluid flows in all directions. Our constructions in this paper work with or without agitation, hence they are robust to random uncontrolled movements, but we choose to ignore this issue and not formally define agitation for ease of presentation.

A rule may involve a movement (translation), or not. First, in the case of no movement: $\vec{u} = \vec{u}'$. Thus we have a rule of the form $r = (s1, s2, b, \vec{u}) \rightarrow (s1', s2', b', \vec{u})$. From above, at most one of $s1, s2$ is **empty**, hence we disallow spontaneous generation of monomers from empty space. *State change* ($s1 \neq s1'$ and/or $s2 \neq s2'$) and *bond change* ($b \neq b'$) occur in a straightforward way, examples are shown in Figure 2(b). If $s_i \in \{s1, s2\}$ is **empty** and s'_i is not, then the rule induces the *appearance* of a new monomer at the empty location specified by \vec{u} if $s2 = \text{empty}$, or $-\vec{u}$ if $s1 = \text{empty}$. If one or both monomer states go from non-empty to **empty**, the rule induces the *disappearance* of monomer(s) at the orientation(s) given by \vec{u} .

For a *movement* rule, $\vec{u} \neq \vec{u}'$. Also, it must be the case that $d(\vec{u}, \vec{u}') = 1$, where $d(u, v)$ is Manhattan distance on the triangular grid, and $s1, s2, s1', s2' \in S \setminus \{\text{empty}\}$. If we fix $\vec{u} \in \mathcal{D}$, then there are two $\vec{u}' \in \mathcal{D}$ that satisfy $d(\vec{u}, \vec{u}') = 1$. A movement rule is applicable if it can be applied both (i) locally and (ii) globally, as follows.

(i) Locally, the pair of monomers should be in state $s1, s2$, share bond b and have orientation \vec{u} of $s2$ relative to $s1$. Then, one of the two monomers is chosen nondeterministically to be the *base* (that remains stationary), the other is the *arm* (that moves). If the $s2$ monomer, denoted X , is chosen as the arm then X moves from its current position $p(X)$ to a new position $p(X) - \vec{u} + \vec{u}'$. After this movement \vec{u}' is the relative position of the $s2'$ monomer to the $s1'$ monomer, as illustrated in Figure 2(b). Analogously, if the $s1$ monomer, Y , is chosen as the arm then Y moves from $p(Y)$ to $p(Y) + \vec{u} - \vec{u}'$. Again, \vec{u}' is the relative position of the $s2'$ monomer to the $s1'$ monomer. Bonds and states may change during the movement.

(ii) Globally, the movement rule may push and/or pull other monomers, or if it can not then it is not applicable. This is formalised as follows, and an example is shown in Figure 3. Let $\vec{v} \in \mathcal{D}$ be a unit vector. The \vec{v} -boundary of a set of monomers Q is defined to be the set of grid points outside Q that are unit distance in the \vec{v} direction from monomers in Q . Let C be a configuration containing adjacent monomers A and B , and let C' be C except that the bond between A and B is null in C' if not null in C . The *movable set* $M = \mathcal{M}(C, A, B, \vec{v})$ is the smallest subset of C' that contains A but not B and can be translated by \vec{v} to give the set $M_{+\vec{v}}$ where the new configuration $C'' = (C' \setminus M) \cup M_{+\vec{v}}$ is such that: (a) monomer pairs in C' that are joined by rigid bonds have the same relative position in C' and C'' , (b) monomer pairs in C' that are joined by flexible bonds are neighbours in C'' , and (c) the \vec{v} -boundary of M contains no monomers. If there is no such set, then we define $M = \mathcal{M}(C, A, B, \vec{v}) = \{\}$. If $\mathcal{M}(C, A, B, \vec{v}) \neq \{\}$, then the movement where A is the arm (which should be translated by \vec{v}) and B is the base (which should not be translated) is applied as follows: (1) the movable set $\mathcal{M}(C, A, B, \vec{v})$ moves unit distance along \vec{v} ; (2) the states of, and the bond between, A and B are updated according to the rule; (3) the states of all the monomers besides A and B remain unchanged and pairwise bonds remain intact (although monomer positions and flexible/null bond orientations may change). If $\mathcal{M}(C, A, B, \vec{v}) = \{\}$, the movement rule is inapplicable (the rule is “blocked” and in particular A is prevented from translating).

A *nubot system* $\mathcal{N} = (C_0, \mathcal{R})$ is a pair where C_0 is the initial configuration, and \mathcal{R} is the set of rules. If configuration C_i transitions to C_j by some rule $r \in \mathcal{R}$ we write $C_i \vdash C_j$. A *trajectory* is a finite sequence of configurations C_1, C_2, \dots, C_ℓ where $C_i \vdash C_{i+1}$ and $1 \leq i \leq \ell - 1$. A nubot system is said to *assemble* a target configuration C_t if, starting from the initial configuration C_0 , every trajectory evolves to a translation of C_t .

A nubot system evolves as a continuous time Markov process. The rate for each rule application is 1. If there are k applicable transitions for a configuration C_i (i.e. k is the sum of the number of rule and agitation steps that can be applied to all monomers), then the probability of any given transition being applied is $1/k$, and the time until the next transition is applied is an exponential

random variable with rate k (i.e. the expected time is $1/k$). The probability of a trajectory is then the product of the probabilities of each of the transitions along the trajectory, and the expected time of a trajectory is the sum of the expected times of each transition in the trajectory. Thus, $\sum_{t \in \mathcal{T}} \Pr[t] \cdot \text{time}(t)$ is the expected time for the system to evolve from configuration C_i to configuration C_j , where \mathcal{T} is the set of all trajectories from C_i to any translation of C_j , and $\text{time}(t)$ is the expected time for trajectory t .

The complexity measure *number of monomers* is the maximum number of monomers that appears in any configuration. The *number of states* is the total number of distinct monomer states that appear in the rule set. *Space* is the maximum area, over the set of all reachable configurations, of the minimum area $l \times w$ rectangle (on the triangular grid) that, up to translation, contains all monomers in the configuration.

The following lemma is used to analyse some of our constructions and was proven in [51].

Lemma 2 ([51]). *In a nubot system, if there are m rule applications a_1, a_2, \dots, a_m that must happen, and (1) the desired configuration is reached as soon as all m rule applications happen, (2) for any specific rule application a_i among those m rule applications, there exist at most k rule applications r_1, r_2, \dots, r_k such that $a_i = r_k$ and for all j , r_j can be applied directly after r_1, r_2, \dots, r_{j-1} have been applied, regardless of whether other rule applications have happened or not, (3) $m \leq c^k$ for some constant c , then the expected time to reach the desired configuration is $O(k)$.*

2.1 Nubots and decision problems

Let $\mathbb{N} = \{0, 1, 2, \dots\}$. $[y]$ denotes a finite length *line segment* of nubot monomers. Given a binary string $x \in \{0, 1\}^*$, written $x = x_0 x_1 \dots x_{k-1}$, we let $[\tilde{x}]$ denote a line segment of k nubot monomers that represent x using one of two “binary” monomer states. $||[\tilde{x}]|| \in \mathbb{N}$ denotes the number of monomers in $[\tilde{x}]$. Given a line of monomers A composed of m line segments, the notation $[A, i]$ means segment i of A , and $[A, i]_j$ means monomer j (or sometimes the bit encoded by monomer j) of segment i of A . We next define what it means to decide a language (or problem) using nubots.

Definition 3. *A finite set of nubot rules \mathcal{N}_L decides a language $L \subseteq \{0, 1\}^*$ if for all $x \in \{0, 1\}^*$ there is an initial configuration C_0 consisting only of the horizontal line $[\tilde{x}]$ of monomers, where by applying the rule set \mathcal{N}_L , the system always eventually reaches a configuration containing only a single “answer” monomer which is in one of two states: (a) “accept” if $x \in L$, or (b) “reject” if $x \notin L$. Further, from the time it first appears, the answer monomer never changes its state.*

2.2 Boolean circuits and the class NC

We define a Boolean circuit to be a directed acyclic graph, where the nodes are called gates and each node has a label that is one of: input (with in-degree 0), constant 0 (in-degree 0), constant 1 (in-degree 0), \vee (OR, in-degree 1 or 2), \wedge (AND, in-degree 1 or 2), \neg (NOT, in-degree 1). One of the gates is identified as the output gate, which has out-degree 0. The *depth* of a circuit is the length of the longest path from an input gate to the output gate. The *size* of a circuit is the number of gates it contains. Besides the output gate, all other gates have out-degree bounded by the circuit size. We work with *layered* circuits: gates on layer i feed into gates on layer $i + 1$. A circuit computes a Boolean (no/yes) function on a fixed number of Boolean variables, by the inputs and constants defining the output gate value in the standard way. In order to compute functions over an arbitrary number of variables, we define (usually, infinite) families of circuits. We say that a

family of circuits $\mathcal{C}_L = \{c_n \mid c_n \text{ is a circuit with } n \in \mathbb{N} \text{ input gates}\}$ decides a language $L \subseteq \{0, 1\}^*$ if for each $x \in \{0, 1\}^*$ circuit $c_{|x|} \in \mathcal{C}_L$ on input x outputs 1 if $w \in L$ and 0 if $w \notin L$.

In a *non-uniform* family of circuits there is no required similarity, or relationship, between family members. In order to specify such a requirement we use a *uniformity function* that algorithmically specifies some similarity between members of a circuit family. Roughly speaking, a *uniform circuit family* \mathcal{C} is an infinite sequence of circuits with an associated function $f : \{1\}^* \rightarrow \mathcal{C}$ that generates members of the family and is computable within some resource bound. Here we care about logspace-uniform circuit families:

Definition 4 (logspace-uniform circuit family). *A circuit family \mathcal{C} is logspace-uniform, if there is a function $f : \{1\}^* \rightarrow \mathcal{C}$ that is computable on a deterministic logarithmic space Turing machine, and where $f(1^n) = c_n$ for all $n \in \mathbb{N}$, and $c_n \in \mathcal{C}$ is a description of a circuit with n input gates.*

Without going into details, we assume reasonable descriptions (encodings) of circuits as strings. We note that there are stricter, but more technical to state, notions of uniformity in the literature, such as AC^0 and $DLOGTIME$ uniformity [2, 20, 34]. We do not require anything less powerful than logspace uniformity here as our main result is a lower bound on nubots' power, hence the more expressive the uniformity condition on circuits, the better (although most of the common circuit classes are reasonably robust under these more restrictive definitions anyway).

Define NC^i to be the class of all languages $L \subseteq \{0, 1\}^*$ that are decided by $O(\log^i n)$ depth, polynomial size logspace-uniform Boolean circuit families. Define $NC = \bigcup_{i=0}^{\infty} NC^i$, in other words NC is the class of languages decided by polylogarithmic depth and polynomial size logspace-uniform Boolean circuit families. Since NC circuits are of polynomial size, they can be simulated by polynomial time Turing machines, and so $NC \subseteq P$. It remains open whether this containment is strict [20]. See [47] for more on circuits.

The complexity class NL is the set of languages accepted by nondeterministic Turing machines that have a read-only input tape and a single workspace of length logarithmic in the input length.

3 Example: A nubots line doubling routine

This section describes a simple construction with the goal of familiarising the reader with the nubot model. We give an algorithm for doubling the length of a line of l monomers in $O(\log l)$ expected time. This algorithm is essentially a simplification of the line growth algorithm in [51], and it will be used in later sections of the paper. We first describe the algorithm then provide a proof for correctness and a time and space analysis.

We require that the input line be comprised of monomers of alternating states, i.e. every monomer in the input line is in one of two unique states with the property that no two adjacent monomers are in the same state. This property of the line is preserved at the end of the line doubling routine.

Lemma 5. *A length l line of monomers can be doubled to length $2l$ in $O(\log l)$ expected time, $O(1)$ states $O(l) \times O(1)$ space.*

Proof. Algorithm description. The algorithm uses concurrent applications of the pair doubling subroutine (PDS) described in Figure 4. As described in more detail below, the algorithm treats the input line of l monomers as a line of $l/2$ monomer pairs that can double in length independently of each other, for even l . After the execution of the subroutine, a monomer pair is transformed into two monomer pairs in alternating states different from the original pair. This ensures that each

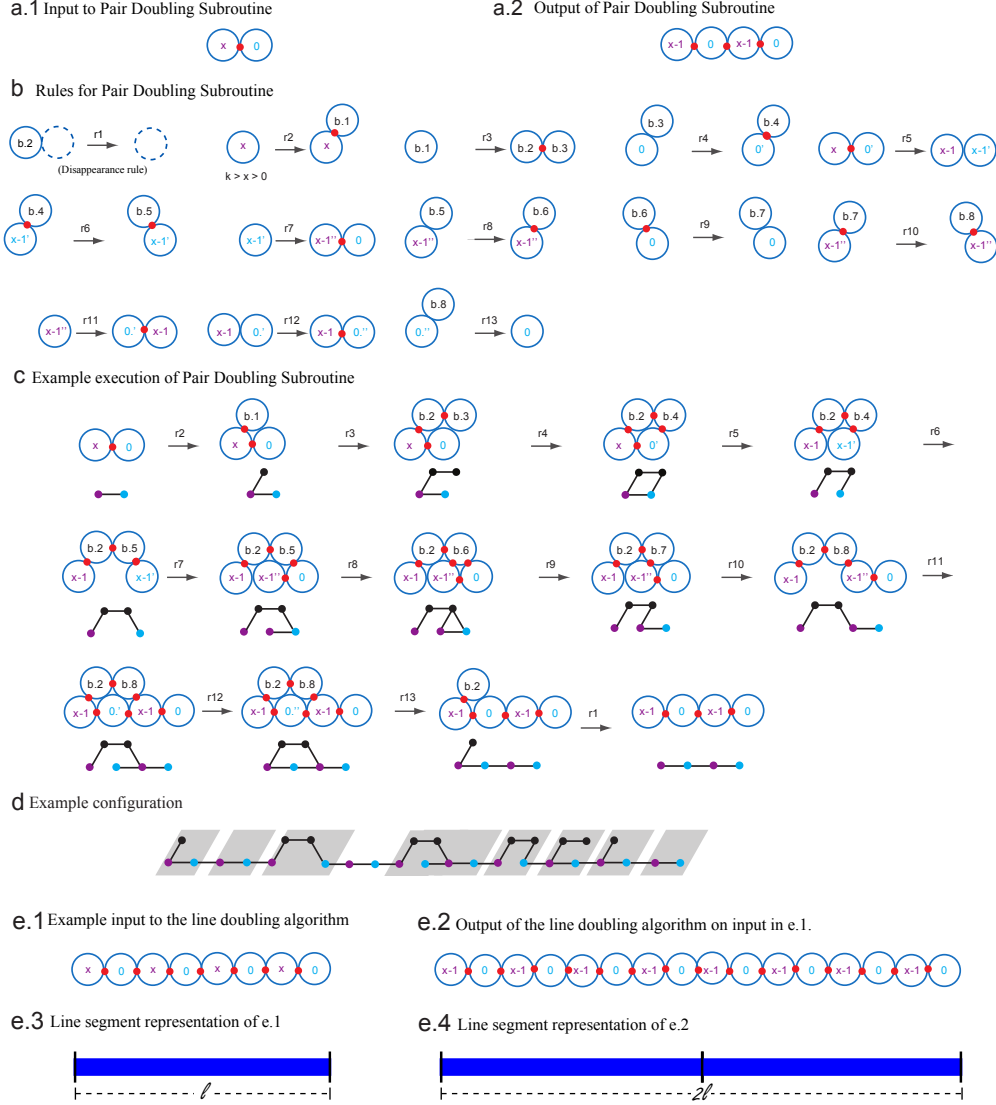


Figure 4: Line doubling nubot algorithm for a line of l monomers that uses a technique from [51]. (a.1) Input, (a.2) output and (b) rule set for the pair doubling subroutine (PDS). The input and output monomers have alternating blue/purple states (with numbers as shown). Since the LHS of Rule r_i is the RHS of Rule r_{i-1} for $i > 2$, the rules must be applied sequentially. (c) Example execution of PDS. (d) Example configuration of a line undergoing length doubling with concurrent applications of PDS to demonstrate the asynchronous nature of the algorithm. (e.1) Example input for the line doubling algorithm. (e.2) Example output for the line double algorithm. (e.3) & (3.4) A simplified “line segment” representation of (e.1) & (e.2) used throughout the paper.

pair of monomers in the input line can only double in length once during the course of the entire algorithm execution. Thus, the length of the input line is doubled by the end of the algorithm, which terminates when every monomer pair in the input has been doubled in length via the subroutine. For odd l , the same thing happens for $\lfloor l/2 \rfloor$ monomer pairs, and the rightmost monomer simply adds a single new monomer to its right.

PDS begins with a pair of monomers with states $x, 0$ and ends with four monomers in states $x-1, 0, x-1, 0$. Figure 4a provides an example input and output of the line doubling algorithm, where monomers are shown as left (purple), right (blue) pairs. The rules for PDS are given in Figure 4b and an example execution is shown in Figure 4c. Each monomer on the line assumes either the “left” or the “right” state: left is colored purple, right is colored blue. The initial $x_{\text{left}}, 0_{\text{right}}$ monomers send themselves to state $(x-1)_{\text{left}}, 0_{\text{right}}$ while inserting two new monomers to give the pattern $(x-1)_{\text{left}}, 0_{\text{right}}, (x-1)_{\text{left}}, 0_{\text{right}}$. To achieve this, the initial pair of monomers create a “bridge” of 2 monomers on top and, by using movement and appearance rules, two new monomers are inserted. The bridge monomers are then deleted and we are left with four monomers. Throughout the execution, all monomers are connected by rigid bonds so the entire structure is connected. PDS completes in constant expected time 13 as shown in Figure 4c since there are a total of 13 rules for PDS that must be applied sequentially, as shown in Figure 4b.

PDS has the following properties: (i) during the application of its rules to an initial pair of monomers $x_{\text{left}}, 0_{\text{right}}$ it does not interact with any monomers outside of this pair, and (ii) a left-right pair creates two adjacent left-right pairs. These properties imply that along a partially formed line, multiple subroutines can execute asynchronously and in parallel, on disjoint left-right pairs, without interfering with each other.

Correctness. To demonstrate that the algorithm doubles the length of the line correctly, it is sufficient to demonstrate that the following invariant holds throughout the algorithm execution and that the algorithm terminates. Every left/right pair of monomers in states $x_{\text{left}}, 0_{\text{right}}$ the input becomes replaced by two left/right monomer pair in states $(x-1)_{\text{left}}, 0_{\text{right}}, (x-1)_{\text{left}}, 0_{\text{right}}$. Locally, the invariant holds from the fact that PDS takes a pair of left/right monomers in states $x_{\text{left}}, 0_{\text{right}}$ as shown in Figure 4a.1 and outputs four monomers in states $(x-1)_{\text{left}}, 0_{\text{right}}, (x-1)_{\text{left}}, 0_{\text{right}}$ as shown in Figure 4a.2, with Figure 4c demonstrating that PDS does this correctly. Since PDS can be applied to each monomer pair independently of any other pair, adjacent concurrent applications of PDS will not block each other. To see that the algorithm terminates, we note that since the input and the output of PDS assume different states and PDS can only double monomer pairs in the input states, each pair of monomers in the original input line can undergo PDS exactly once.

Time and space analysis. As shown in Figure 4c, the space complexity of PDS is 4×2 . Since PDS only attaches monomers on top of the input monomers as per the rules, adjacent monomer pairs in the input of the line doubling algorithm will remain on the same axis (i.e. maintain their y -coordinates on the triangle grid shown in Figure 2a). Thus, the space complexity of the line doubling algorithm is $O(l) \times O(1)$. We have established above that the expected time for PDS is 13. The event in which an application of PDS takes place is a Poisson process; therefore, the expected time for a single occurrence of this event to take place is $1/k$, where k is the total possible positions for PDS to be applied. Let T be the time it takes for the line doubling algorithm to terminate on an input of length l , then the expected value of T is $E[T] = 13 \sum_{i=1}^{l/2} 1/i = O(\log l)$. \square

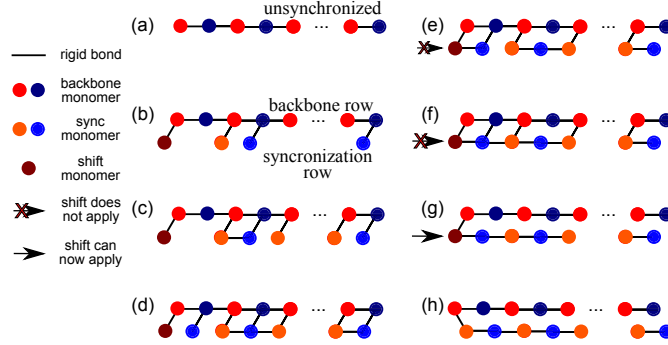


Figure 5: Shift synchronization, see reference [51] for more details. (a) Initial state. (b) Monomers, randomly and in parallel, each grow a new *synchronization monomer* below. The leftmost new monomer (in brown) is denoted the “shift monomer”. (c)–(d) When synchronization monomers detect neighbouring horizontal synchronization monomers to the left and right, they bond. When a synchronization monomer has bonded to both horizontal neighbours, its bond to its parent monomer is removed. (e) When the shift monomer detects a synchronization monomer neighbour to the right, it changes state, permitting a movement rule to be applied, although the connectivity prevents this movement from occurring yet. (f) Synchronization monomers continue to appear and update their bond structure. (g)–(h) All of the vertical rigid bonds are gone and the movement rule can now be applied in one step. All monomers on the original horizontal line detect the change in state (parity) of their neighbour below.

4 Using synchronization to communicate quickly

In previous work [51] a fast signalling method, called synchronization, was introduced for the nubot model. Here, we use the term “shift synchronization” for this technique, and introduce another kind of synchronization called “lift synchronization”. With these two synchronization mechanisms, we can send one of two distinct messages (bits) to all monomers on a line in expected time that is merely logarithmic in the line length.

Lemma 6 (Communication via synchronization). *Let ℓ be a length n line of monomers, where each monomer in ℓ is in one of two distinct states $\{s_0, s_1\}$, with each adjacent pair distinct from each other. A bit $b \in \{0, 1\}$ can be communicated to all monomers on the line in $O(\log n)$ expected time, $O(1)$ monomer states and $O(n) \times O(1)$ space.*

Proof. We first give a brief overview of shift synchronization using Figure 5, more details can be found in [51]. Each monomer on the line, in state $s \in \{s_0, s_1\}$, attaches a new *synchronization monomer* below itself with state s' and with a rigid bond. When a synchronization monomer with state s' senses a new horizontally adjacent neighbouring synchronization monomer it forms a rigid (horizontal) bond with this monomer. After connecting to both neighbouring synchronization monomers, the monomer removes the bond between it and its parent monomer (with state s) above.

The rightmost and leftmost synchronization monomers are treated differently. At the rightmost end of the line, the new monomer requires only one bonded neighbour (to the left) before removing its bond to its parent monomer. The leftmost synchronization monomer is called the “shift monomer”.

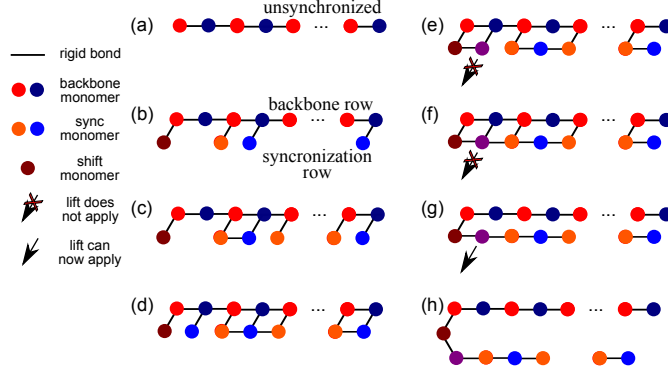


Figure 6: Lift synchronization. Lift synchronization uses similar ideas to shift synchronization, except instead of pushing the entire synchronization row horizontally, the synchronization row is moved vertically below, and away, from the original line. The monomers on the original line are then free to detect the disappearance of synchronization monomers, signalling the completion of the lift synchronization.

This shift monomer attempts to push the (new) synchronization row to the right. However, by definition of the movement rule, the shift monomer can move only after all of vertical rigid bonds between the synchronization row and the original line have been removed. Also, due to the order in which bonds are formed and removed, this can only happen after the entire synchronization row has grown. At some point, we are guaranteed to get to the configuration in Figure 5(g), where the shift monomer is free to push right. After the move (Figure 5(h)), the relative position of synchronization monomers to their generating monomers has changed. Thus, the original line of monomers are free to detect that synchronization has occurred, and a 0 bit has been communicated to all of them.

To send a 1 bit we use a similar method, called lift synchronization, shown in Figure 6. In lift synchronization the synchronization row is lifted vertically down, and away, from the original line, rather than being shifted right. As with shift synchronization this can only occur after the entire synchronization row has been built and all bonds are in their final form. After the move (Figure 6(h)), the monomers on the original line detect the new empty space below, and thus detect that a 1 bit has been communicated to them.

In this way, for a line in any of the 6 rotations, it is possible to communicate a 0 or 1 bit, depending on whether shift or lift synchronization is used. The expected time to send the bit is $O(\log n)$, as (a) all new monomers are created independently and in parallel, and (b) each monomer needs only to wait on a constant number of neighbours in order to get its bond structure to the final configuration. The space and states bounds are straightforward to see. \square

5 Fast line growth using $O(1)$ states

The line growth algorithm given in [51] grows a line of length $n \in \mathbb{N}$ in $O(\log n)$ time, using $O(\log n)$ monomer states and starting from a single monomer on the grid. Here, we provide an alternative line growth algorithm that completes in $O(\log^2 n)$ time, using $O(1)$ monomer states and starting from $O(\log n)$ monomers on the grid. Although our construction is an $O(\log n)$ factor slower than

```

1 line_growth( $b = b_{\lfloor \log_2 n \rfloor} \dots b_2 b_1 b_0$  : binary string that encodes  $n \in \mathbb{N}$ )
2 Initialise:  $k = 0$ ,  $\text{mask} = 1$ ,  $\text{generator} = 1$ ,  $\text{line} = 0$ 
3 while  $k \leq \lfloor \log_2 n \rfloor$  do
4      $\text{mask} := 2 \cdot \text{mask}$                                      /*  $\text{mask} := 2 \cdot 2^k = 2^{k+1}$  */
5     if  $b_k = 0$  then
6          $\text{generator} := 2 \cdot \text{generator}$                      /*  $\text{generator} := 2 \cdot 2^k = 2^{k+1}$  */
7     else if  $b_k = 1$  then
8          $\text{generator} := 3 \cdot \text{generator}$                      /*  $\text{generator} := 3 \cdot 2^k$  */
9          $\text{line} := \text{line} + \text{generator} - \text{mask}$                  /*  $\text{line} := \text{line} + 2^k$  */
10         $\text{generator} := \text{generator} - (\text{generator} - \text{mask})$  /*  $\text{generator} := 3 \cdot 2^k - 2^k = 2^{k+1}$  */
11        Delete  $b_k$ 
12         $k++$ 
13 return line

```

Figure 7: Algorithm that takes a binary string b as input, that encodes $n \in \mathbb{N}$, and returns the integer n . This algorithm describes the control flow for the nutbots construction that builds a line of length n in the proof of Theorem 8.

that in [51], it uses only $O(1)$ states while maintaining the property that all growth is contained within an $O(n) \times O(1)$ region. The latter two properties are both requirements in achieving our main theorem via the other constructions in this paper, which extensively use this line growth algorithm.

Problem 7 (Binary Line Growth problem). *Input: A line of $\lfloor \log_2 n \rfloor + 1$ monomers each in one of two binary states from $\{s_0, s_1\}$, that encode the binary string $b = b_{\lfloor \log_2 n \rfloor} \dots b_1 b_0$ in the standard way, where $n = \sum_{i=0}^{\lfloor \log_2 n \rfloor} b_i \cdot 2^i$.*
Output: A line of n monomers.

Theorem 8 (Binary Line Growth). *There is a nubot algorithm to solve the Binary Line Growth problem in expected time $O(\log^2 n)$, space $O(n) \times O(1)$, and with $O(1)$ states.*

Proof. As described in the problem statement, the input n is encoded as a line of $\lfloor \log_2 n \rfloor + 1$ monomers where the i th monomer encodes bit b_i of the binary string $b = b_{\lfloor \log_2 n \rfloor} \dots b_1 b_0$, and where b encodes $n \in \mathbb{N}$ in the usual way. The construction proceeds iteratively: at iteration k , where $0 \leq k \leq \lfloor \log_2 n \rfloor$, bit b_k is read from the input and if $b_k = 1$ the partially grown line is increased in length by the value 2^k , otherwise the length of the line remains unchanged. The idea is described at a high-level in the algorithm in Figure 7, below we show that the integer variables in that algorithm can be implemented as lines of the corresponding integer lengths, and these can be acted upon in a way that quickly builds the length n line.

Construction details. During construction, the line-growing configuration is composed of three main regions. The first is the “input”, as described above; at iteration k of the algorithm the least significant bit b_k of the input is read (stored), and deleted. Then we have a working region containing two lines, respectively called the “generator” and the “mask”, each of which have length 2^k at iteration k . Finally we have the “line” under construction: at iteration k , the line length is given by the binary number $b_{k-1} \dots b_1 b_0$ encoded by the first k bits (LSBs) of the input.

The construction begins with the rightmost of the input monomers growing a small, constant-size, hardcoded structure containing both the generator and mask, both initialised to be of length 1.

Figure 7 describes a (seemingly overcomplicated, but analogous to our construction) algorithm for generating the integer n from a bit string b . Our construction implements this algorithm, but

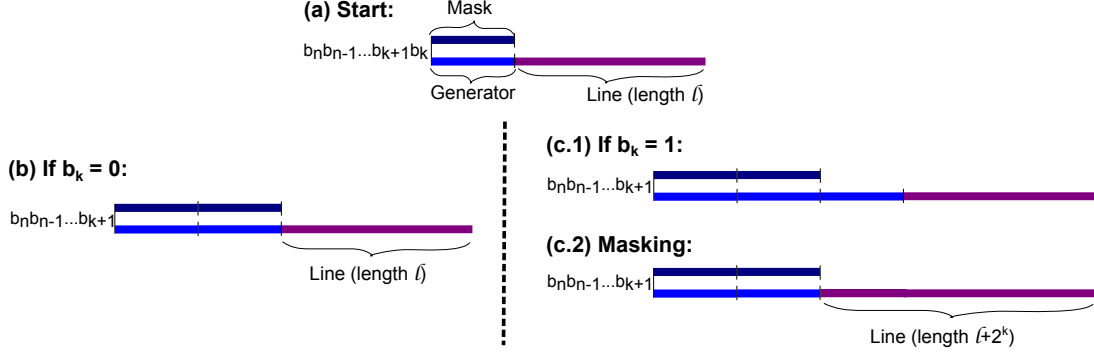


Figure 8: Reading a single input bit, and growing the line accordingly. (a) From the start state, depending on if the least significant bit remaining in the string (bit b_k , the k^{th} bit of the original string) is a 0 or 1, the system will end up in one of two different configurations, shown in (b) or (c.2). More details for the $b_k = 1$ case are shown in Figure 9.

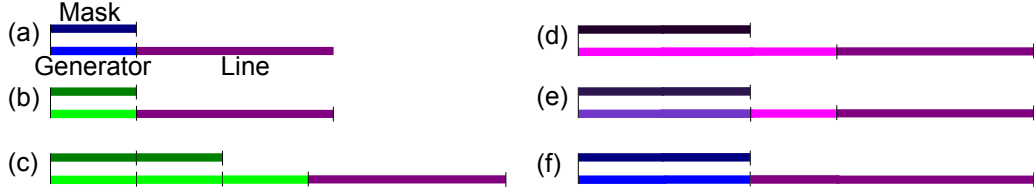


Figure 9: Reading a 1 bit. (a) Initial configuration. (b) Synchronization message sent to the mask (dark blue) and generator (light blue) lines to initiate tripling of the generator. (c) Masks doubles in length, generator triples. (d) Synchronization. (e) Masking: monomers in the generator look immediately above for a corresponding monomer in the mask line. If none exists, the generator monomer changes its state to that of a line monomer, if one exists it stays part of the generator. (f) Masking finished, synchronization.

where the integer variables “mask”, “generator”, “line” are encoded in unary as lines of monomers of that length. It is straightforward to verify, via induction on k , that upon input of the string $b \in \{0,1\}^*$, that encodes $n \in \mathbb{N}$, the algorithm in Figure 7 returns the integer n . Our nubot implementation of one iteration of this algorithm is shown in Figure 8. Figure 8 uses a high-level notation where lines of nubot monomers are represented as colored lines drawn on the square grid. We describe the construction by describing the main primitives it uses to implement the algorithm in Figure 8: line doubling or tripling implement multiplying by 2 or 3; synchronization implements bit communication—and thus which instructions to implement next—to all monomers; and masking implements taking differences.

Line doubling and tripling. Line doubling takes a line of length ℓ and generates a line of length 2ℓ , as described in Section 3. Line tripling takes a line of length ℓ and generates a line of length 3ℓ , using a similar technique (rather than inserting 2 monomers, we insert 1, synchronize, then insert 1 again), hence we omit the details.

Synchronization & communicating a bit. We use a synchronization algorithm to simultaneously switch a line of monomers into a single shared state. As described in Section 4, we have the two methods of lift and shift synchronization: we use one to communicate a 0 bit and the other to communicate a 1 bit to monomers in the generator and mask.

Masking. For two lines of different lengths, $\ell_1 > \ell_2$, *masking* communicates their difference $\ell_1 - \ell_2$ to the line of greater length ℓ_1 . The lines are assumed to be orientated parallel, touching, and horizontal with their leftmost extent at the same x position. Assume the shorter line is on top: it synchronizes (by growing a new synchronization row on top), then the longer line synchronizes (by growing a new synchronization row on bottom). Then the monomers in the longer line detect the presence or absence of monomers on the shorter line above: if there is a monomer above then the longer line monomer goes to state s_1 , if not it goes to state s_2 . See Figure 9(d)-(f) for an outline.

Final steps. The final bit of the input to be read is $b_{\lfloor \log_2 n \rfloor} = 1$ (the MSB of a binary number is always 1) and just before reading it the line length is $n - 2^{\lfloor \log_2 n \rfloor}$. Upon reading the final bit $b_{\lfloor \log_2 n \rfloor}$ some message passing occurs (via synchronizations) to trigger the deletion of the mask and to cause the generator monomers to change state so that they are now part of the line. This latter step adds $2^{\lfloor \log_2 n \rfloor}$ (generator length) to the line, giving the desired line length of n .

Time, space, and states analysis. Line doubling/tripling of a length n line happens in expected time $O(\log n)$, as does synchronization. There are $O(\log n)$ iterations each with a constant number of doublings/triplings and synchronizations, hence the total expected time is $O(\log^2 n)$. The three lines (mask, generator, line) are of length $\leq n$ and with their synchronization rows the height needed is 4, giving a space bound of $O(n) \times O(1)$. A straightforward analysis of the algorithm shows that $O(1)$ states are sufficient. \square

6 Fast parallel sorting

In this section we show how, on the nubot model, to sort n binary numbers, taken from the set $\{0, 1, \dots, n-1\}$, in polylogarithmic expected time and a constant number of states. Our sorting algorithm is loosely inspired by the work of Murphy et al. [33] who show that physical techniques can be used to sort numbers that are represented as the magnitude of some physical quantity. They show that a variety of physical mechanisms can be thought of as an implementation of fast parallel sorting, including gel electrophoresis and chromatography (molecular weight), rainbow sort [44] (frequency), and mass spectrometry (mass to charge ratio). However, our construction needs to take care of the fact that ours is a robotic-style geometric model that needs to implement fast growth while handling blocking and other geometric constraints. A similar algorithm works for variations on this problem, such as sorting $< n$ such numbers, but we omit the details of that.

We first define the distinct element sorting problem and then formally state the result.

Problem 9 (Distinct element sorting problem). *Input: A line of monomers, denoted $A = [A, 1][A, 2] \dots [A, n]$, composed of $n \in \mathbb{N}$ contiguous line segments where for each $i \in \{1, 2, \dots, n\}$ line segment $[A, i]$ is of length $|[A, i]| = \lfloor \log_2 n \rfloor + 1$ and encodes a distinct binary number from $\{0, 1, \dots, n-1\}$, where specifically, for all $j \in \{1, 2, \dots, \lfloor \log_2 n \rfloor\}$ it is the case that monomer $[A, i]_j$ is in one of two binary states from $\{s_0, s_1\}$ and the end-of-segment monomer $[A, i]_{\lfloor \log_2 n \rfloor + 1}$ is in*

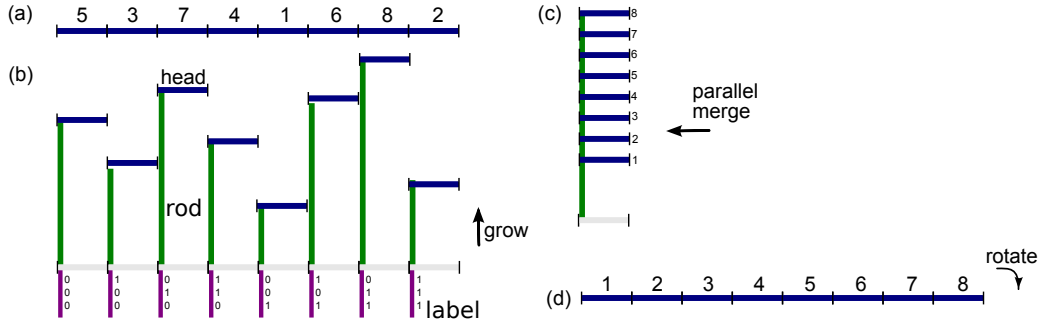


Figure 10: High-level overview of the sorting algorithm. (a) A line of $n(\lfloor \log_2 n \rfloor + 1)$ monomers, with n blue line segments (“heads”) each is the binary representation of a natural number $i \leq n$. (b) A blue head that encodes value i is grown to height i by a green rod, in time polylog in i . Purple “labels” are also grown at the bottom. (c) The heads are horizontally merged, using the labels to synchronize, to be vertically aligned. (d) Merged heads rotate down into a line configuration, giving the sorted list. Each stage occurs in expected time polylogarithmic in n , more details appear in subsequent figures.

one of two binary end-of-segment states $\{s\#_0, s\#_1\}$.

Output: A line A' consisting of the n binary line segments sorted in increasing order of the standard lexicographical ordering of their binary sequences.

Theorem 10 (Distinct element sorting). *Any instance A of the distinct element sorting problem is solvable on the nubot model in expected time $O(\log^3 n)$, space $O(n \log n) \times O(n)$, and $O(1)$ monomer states.*

Proof. The general idea is as follows. For each element i (encoded as a “head”) to be sorted, we grow a line of monomers (a “rod”) to length i as shown in Figure 10(b). After doing so, the relative heights of the heads gives their order. We then move each head horizontally left, through a sequence of $O(\log n)$ parallel merging steps, so that all heads are vertically aligned (Figure 10(c)). Finally, the heads are rotated and translated so that they lay along a vertical line as shown in Figure 10(d), in increasing order. The details are described next.

6.1 Sorting details: rod growth and labeling

We begin with an instance of the distinct element sorting problem, an example of which is shown in Figure 11(a).

Initialization. The monomers begin in binary states as described in Definition 9. Growth begins at each of the n blue heads: the head is copied and rotated down to vertical as shown in Figure 11(b). This rotation of a $O(\log n)$ length line takes $O(\log \log n)$ expected time to complete using the parallel “arm rotation” method in [51]—that is, each monomer independently rotates by one position, relative to its leftmost neighbour. After rotation (Figure 11(b)), each blue-green line independently synchronizes, then makes a copy of itself which is in turn rotated down to become

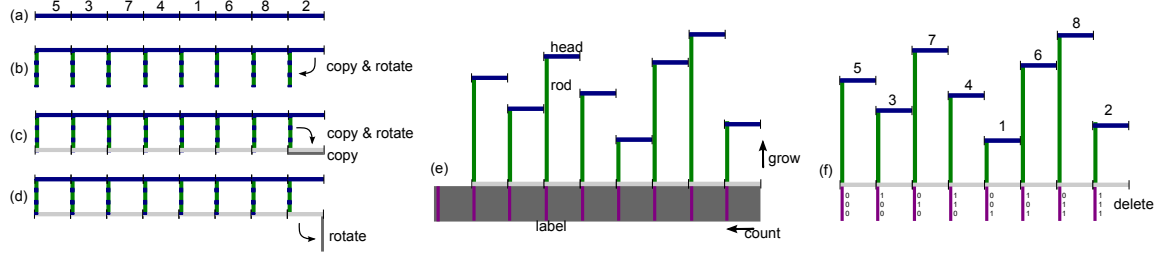


Figure 11: The beginning of our sorting algorithm, this gives the details for the overview in Figure 10(a)–10(b). (a) Initial configuration with head monomers in blue. (b) Head monomers are copied and rotated down to vertical (dashed blue-green), and then (c) are copied and rotated down to horizontal to form the light-grey label region. The light-grey region synchronizes after being copied. On the right, the dark grey region is copied and in it rotates down to vertical, as shown in (d). In (d) the heads have received a “synchronization done” message from the light-grey region and (e) they grow a vertical green rod (line) of length equal to the value encoded in the head (note the heads are not connected to each other, except through the green rods and light-grey region). Also in (e), the dark grey region grows a purple counter, from right to left, that counts from $2^{\lfloor \log_2 n \rfloor + 1} - 1$ down to 0 (see main text for details) and is padded with $\lfloor \log_2 n \rfloor + 1$ monomers between each counter column (thus growing a $(\lfloor \log_2 n \rfloor + 1) \times (\lfloor \log_2 n \rfloor + 1) 2^{\lfloor \log_2 n \rfloor + 1}$ rectangle, in dark grey and purple). (f) All parts of the dark grey region delete themselves, except those directly below a green rod (purple). The purple regions that remain encode n distinct binary strings.

one of the n horizontal light-grey line segments shown in Figure 11(c). After all light-grey segments are horizontal, they bond to each other and synchronize. This entire process completes in expected time $O(\log n)$, using Lemma 2, and is dominated by the synchronization process.

Grow rods. After this synchronization step, as shown in Figure 11(c), the rightmost grey line segment is copied to form a dark grey segment that is copied down to vertical in Figure 11(d). Also in Figure 11(d), and triggered by the previous synchronization, the blue-green rods, in $O(\log n)$ expected time, signal the heads to disconnect from each other, and the blue-green rods then begin “growing upwards”. This vertical growth of the rods implements a form of counting: we want the rods to grow to the height encoded by their blue head. This is carried out by using the line-growth algorithm in Section 5 which takes time $O(\log^2 n)$ (an alternative method would be to use a suitable counter, such as the one described below). After a rod has grown to the value encoded in its head, shown in Figure 11(e), the rod synchronizes, this latter step taking expected time $O(\log n)$. After expected time $O(\log^3 n)$ all n rods have synchronized.

Label growth. Rod growth occurs above the light-grey line. Below that line another process takes place, the purpose of which is to label each rod with its position (from right), as a binary number in purple. Here the dark-grey line (Figure 11(d), on right) grows a “padded” counter, from right to left. The result of this counter is shown in Figure 11(e) and is a $(\lfloor \log_2 n \rfloor + 1) \times (\lfloor \log_2 n \rfloor + 1) 2^{\lfloor \log_2 n \rfloor + 1}$ rectangle where each of the purple columns, from right to left, encodes a distinct value from $2^{\lfloor \log_2 n \rfloor + 1}$ down to 1, with the grey regions in between being there for padding purposes only.

This counter works as follows. The counter is a modified version of the one used in Section 6.2 of [51]; the counter in [51] used $O(\log n)$ states, here we use $O(1)$ states. First note that the dark grey strip is of height $\lfloor \log_2 n \rfloor + 1$, it begins counter growth by converting each of its monomers to a state that represents the bit 1, giving the binary representation of the number $2^{\lfloor \log_2 n \rfloor + 1} - 1$. Let $j = \lfloor \log_2 n \rfloor + 1$, and we begin from the single dark grey column, applying the following procedure iteratively to each new column until $j = 1$. Each column copies itself to the left and in the new column the j th bit is flipped. Both columns then decrement their value of j , and *both* iterate the copy and bit-flip procedure. As is the case in [51], this process happens asynchronously and independently to all columns. After this happens we have a $(\lfloor \log_2 n \rfloor + 1) \times 2^{\lfloor \log_2 n \rfloor + 1}$ rectangle containing all of the purple columns. We are not done yet: we wish for the purple counter columns to align themselves with the n green rods which are distance $\lfloor \log_2 n \rfloor + 1$ apart, as in Figure 11(e). To achieve this, another round of column insertion (i.e. counting) begins, so that between each pair of counter columns, exactly $k = \lfloor \log_2 n \rfloor$ new columns are inserted (between each pair of purple columns we are implementing a counter that counts from k down to 1; note that the integer k is available since the purple counter columns are of height k). Now the purple counter rows are exactly distance $k + 1 = \lfloor \log_2 n \rfloor + 1$ apart. When the process is complete the bottom row of the entire rectangle synchronizes, to give the structure illustrated in Figure 11(e) (although the grey rectangle extends further to the left than shown).

To give a straightforward time analysis, we assume that the copying and decrementing for an individual column happens sequentially and so takes expected time $O(\log n)$. Then in the completed counter, each counter column is the result of no more than $O(\log n)$ column-copying operations, hence any monomer in the final rectangle depends on the application of $O(\log^2 n)$ rules. Applying Lemma 2 gives an expected time of $O(\log^2 n)$. The final synchronization step costs $O(\log n)$ expected time, giving a total expected time of $O(\log^2 n)$.

Deletion and synchronization. All columns of the dark grey region then delete themselves, *except* those that are directly below a green rod. The deletion events happen in time $O(\log n)$. The purple regions that remain are n counter columns that encode n distinct binary numbers. After each green rod (above) has synchronized it signals to the light-grey line. As each counter row below completes deletion, it too signals to the light-grey line. The light-grey line undergoes a lift synchronization. The system is now in the configuration shown in Figure 11(f).

Analysis. As already discussed, rod growth and the subsequent synchronization of all n rods takes expected time $O(\log^3 n)$, and label growth takes expected time $O(\log^2 n)$.

6.2 Sorting details: merging

Now that all rods have grown, and are labeled, we will now merge them as shown in in Figure 10(c).

Main idea. Intuitively, we would like to simply shift all of the heads to the left, deleting any rods that get in the way. However, if we are not careful, rods can block each other and significantly slow down the process so that it no longer runs in time polylogarithmic in n (consider the worst case, where the shortest rod is the rightmost one, and we wish to move all heads to the left). Our merging algorithm gets around this issue by merging in a pairwise fashion. Every second pair of heads merge, deleting one of the rods and then the light-grey line synchronizes. We are left with $\lceil n/2 \rceil$ rods, each having two heads. Then every second pair of those merge, and so on for $O(\log n)$ iterations. To

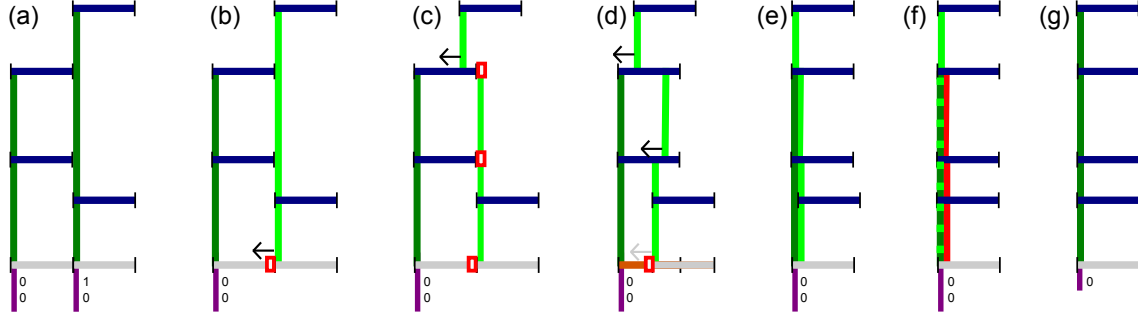


Figure 12: Merging the heads on two adjacent rods, this gives the details for the overview in Figure 10(c). Exactly one round of parallel (pairwise) head-merging has already occurred, and so each green rod has two blue heads. (a) Two rods, with two heads each: the goal is to merge all 4 heads onto the left rod. (b) Due to how they were generated the LSBs (top bits) on the two purple labels are distinct. If the LSB bit is 1, the rod moves left, and deletes the purple label monomers. The rod tries to move left by having the light-grey line sequentially delete its $O(\log n)$ monomers one at a time, although here the rigid rod is immediately blocked due to collisions with the heads on the left. (c) Collisions are marked in red. The rod monomers at the collision locations delete themselves. The new shorter rods can continue moving to the left, by “walking” along the blue heads as shown in (c) and (d). (e) The rods on the right make contact with the rods on the left. (f) The contact triggers a “done” state to be reached by the rod on the left. It also signals for the rod on the right to delete itself. Head monomers from the right are shifted to their new rod. (g) When everything has moved into place, synchronization occurs along the single green rod, and the LSB (top) bit of the purple label is deleted.

organise the correct order of mergings, we use the purple labels, specifically their binary sequences, which are shown in Figure 11(f).

Merging algorithm. The following procedure is iterated until there is exactly one rod left. For each i , the i th rod checks its label, if the least significant bit (LSB) of the purple label is 1 (in Figure 11(f) the LSB is on top), then rod i attempts to merge with rod $i - 1$ to its left, by “moving” its head to the left distance $\lfloor \log_2 n \rfloor + 1$. This pairwise merging process is described in the caption of Figure 12. Rod i , and its label (but not its head) get deleted in the process. After merging of the pair of rods is complete, rod $i - 1$ deletes its LSB, thus shortening its purple region by 1. Rod $i - 1$ now has two heads, and signals to the light-grey line that it is done. After all pairs have merged, we have $\lceil n/2 \rceil$ rods, with 2 heads each. At this point the light-grey line synchronizes and the process iterates. After $O(\log n)$ rounds of pairwise merging we are left with one rod, which has no label and is carrying all n heads.

When a pair of rods are merging, the right rod needs to move distance $\lfloor \log_2 n \rfloor + 1$ to the left. In the worst case there are $\lceil n/2 \rceil$ collisions for a pair of rods, however, these are all resolved in parallel as described in Figure 11(c). So we have $\leq \lceil n/2 \rceil$ heads, that each need to independently walk distance $\lfloor \log n \rfloor + 1$ to the left which naïvely takes $O(\log^2 n)$ expected time, and by applying Lemma 2 takes $O(\log n)$ expected time.

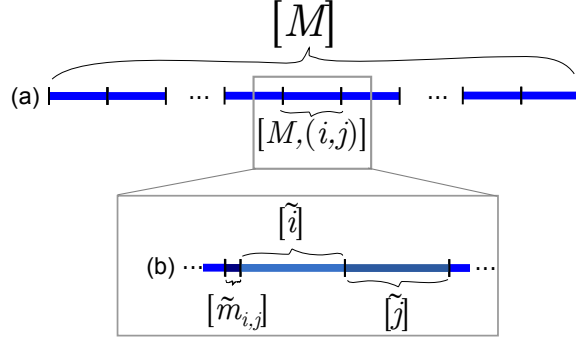


Figure 13: (a) Encoding of a Boolean matrix M as a line of monomers $[M]$. (b) Zoom-in of the encoding of a single matrix entry $M_{ij} \in \{0, 1\}$ as a line segment $[M, (i, j)]$ that contains a single monomer $[\tilde{m}_{i,j}]$ that encodes the bit $M_{i,j}$ and line segments of binary monomers, $[\tilde{i}]$ and $[\tilde{j}]$, that encode i and j .

Final steps. After merging is complete, the heads are on a single rod, sorted vertically upwards in increasing order of head value. The heads rearrange themselves on the rod to that they are separated by vertical distance exactly $\lfloor \log_2 n \rfloor + 1$, and then rotate down into a line configuration, giving the sorted list as shown in Figure 10(d).

6.3 Sorting details: time, space and states analysis

The expected time to complete the various stages of sorting was given above, and is dominated by growing and synchronizing the rods, which is $O(\log^3 n)$. For the space analysis, note that the length of the light-grey line is $O(n \log n)$ (giving the horizontal space bound). The rods are of height $O(n)$, and the purple labels are of height $O(\log n)$, giving a vertical space bound of $O(n)$. Hence we get a space bound of $O(n \log n) \times O(n)$. All counters and line growth algorithms use number of states that is constant, which can be seen by a careful analysis of each part of the construction. \square

7 Fast Boolean matrix multiplication

Let M and N be $n \times n$ Boolean matrices. Let $M_{i,j}$ denote the element at row i and column j of M , and let MN denote their matrix product. The following two definitions are illustrated in Figure 13 and describe our encoding of a square matrix as an addressed line of monomers.⁴

Definition 11 (Matrix element encoding). *An element $M_{i,j}$ of $n \times n$ Boolean matrix M is encoded in nubot monomers as a line of $O(\log n)$ monomers $[M, (i, j)] = [\tilde{m}_{i,j}][\tilde{i}][\tilde{j}]$ where $[\tilde{m}_{i,j}]$ is a nubot monomer that encodes the bit $M_{i,j}$, and $[\tilde{i}]$ and $[\tilde{j}]$ are lines of binary monomers of length $O(\log n)$ that encode the numerical values i and j , respectively (the segments $[\tilde{i}]$ and $[\tilde{j}]$ are each terminated by delimiter monomers).*

⁴Our choice of a 1D, rather than 2D, encoding simplifies our constructions. It would also be possible use a more direct 2D square encoding, which, it turns out, can be unfolded to and from our line encoding in expected time $O(\log n)$. We omit the details.

Definition 12 (Monomer encoded Boolean matrix). *An $n \times n$ Boolean matrix M is encoded in nubot monomers as a line of $O(n^2 \log n)$ monomers $[M] = [M, (1, 1)][M, (1, 2)] \dots [M, (n, n-1)][M, (n, n)]$ of all $[M, (i, j)]$ for $1 \leq i, j \leq n$, ordered from left to right, first by i , then by j .*

The main result of this section, Theorem 14, is a fast parallel algorithm for Boolean matrix multiplication.

Problem 13 (Monomer encoded Boolean matrix multiplication problem). *Input: Monomer encoded Boolean matrices $[A]$ and $[B]$, that represent $n \times n$ Boolean matrices A, B . Output: Monomer encoding of the Boolean matrix $[C] = [AB]$.*

Theorem 14. *The monomer encoded Boolean matrix multiplication problem can be solved in $O(\log^3 n)$ expected time, $O(n^4 \log n) \times O(n^2 \log n)$ space and with $O(1)$ monomer states.*

7.1 Parallel function evaluation in 2D

Before proving Theorem 14 we give a useful lemma that formalises a notion of nubots efficiently computing many (n^2 here) functions in parallel, where each function acts on two length k inputs. Figure 14 illustrates the proof.

Lemma 15 (Parallel function evaluation in 2D). *Let \mathcal{F} be any function that maps a pair of length k adjacent parallel horizontal monomer lines $[X], [Y]$ to a length k horizontal monomer line $[Z]$, that is $\mathcal{F}([X], [Y]) = [Z]$, and moreover \mathcal{F} is nubot computable in $O(k)$ expected time, $O(k) \times O(1)$ space, and $O(1)$ states. Let $[A] = [A, 1][A, 2] \dots [A, n]$ and $[B] = [B, 1][B, 2] \dots [B, n]$ be monomer lines, each composed of n consecutive length k monomer lines (called “line segments”). Then, given $[A]$ and $[B]$ as input, the line $[C] = [C, 1][C, 2] \dots [C, n^2]$ consisting of all $[C, i + (j - 1)n] = \mathcal{F}([A, i], [B, j])$ for $1 \leq i, j \leq n$ is computable using nubots in $O(k + \log^2 n)$ expected time, $O(n^2 k) \times O(nk)$ space, and $O(1)$ states.*

Proof of Lemma 15. Figure 14 gives an overview of the construction. From an initial configuration with $[A]$ and $[B]$ adjacent as in Figure 14(a), $[B]$ rotates down to vertical (Figure 14(b)). $[B]$ is copied from the grey line which rotates down to horizontal as shown in Figure 14(c). In Figure 14(d.1), we duplicate each line segment $[A, i]$, for $1 \leq i \leq n$, n times, down to the grey vertical line, which acts as a barrier to stop the duplication. A one monomer horizontal gap is inserted between adjacent columns of green columns (of line segments), which triggers a vertical synchronization, shown as a vertical red line in Figure 14(d.2) of each completed green column. Next, monomer-to-monomer messages are passed, horizontally from right to left, within each green line segment to signify that monomers should change from being “vertically connected” to being “horizontally connected”. After this, the vertical red synchronization lines carry out another synchronization and then delete themselves in a way that keeps all green monomers horizontally connected. In Figure 14(d.3), each purple $[B, j]$ inserts a 1-monomer vertical gap between it and its neighbour $[B, j + 1]$. After all gaps insert, the purple vertical line synchronizes, and then n horizontal synchronizations happen which tell excess duplicates of $[A, i]$ to delete themselves to give the configuration in Figure 14(d.4).

Next, a duplication and deletion process occurs with $[B, j]$ line segments as shown in Figure 14(e) (similar to what we did before, but now horizontally rather than vertically). The $[B, j]$ ’s duplicate until they hit the vertical grey barrier on the right, at which point the system synchronizes. After

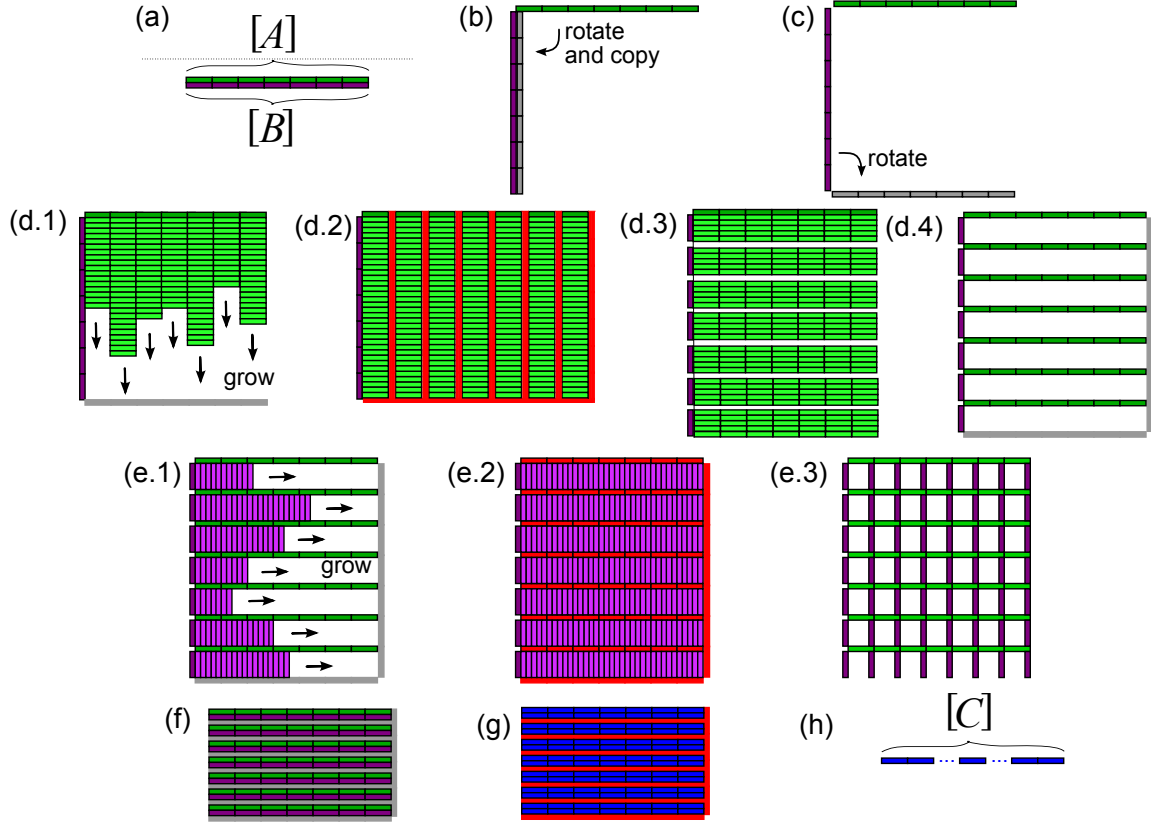


Figure 14: Parallel function evaluation in 2D, used in the proof of Lemma 15. (a) Initial configuration with line $[A]$ in green and line $[B]$ in purple, each has n line segments. We wish to compute \mathcal{F} on all n^2 pairs of line segments in $[A]$ and $[B]$. (b) $[B]$ rotates down to vertical and duplicates. (c) The duplicate of $[B]$ rotates down to horizontal creating a grey border. (d.1) Each segment of $[A]$ duplicates, and the resulting pair of segments duplicate, and so on iteratively. (d.2) The copied line segments of $[A]$ reach the bottom grey border line. A vertical gap is inserted between each *column* of green line segments, then synchronization occurs (red). (d.3) The vertical synchronization causes the system to change connectivity (to be a comb with horizontal teeth), allowing for segments of $[B]$ to insert 1-monomer vertical gaps between themselves. (d.4) Duplicates of $[A]$, not adjacent to a gap delete themselves; monomers rearrange and horizontal synchronization rows are regrown. (e.1) Segments of $[B]$ duplicate, iteratively. (e.2) When duplication finishes, synchronizations occur along the copied segments of $[A]$. (e.3) Duplicates of segments of $[B]$ not adjacent to the left/right ends of duplicates of segments of $[A]$ delete themselves. (f) Purple duplicated line segments of $[B]$ rotate up to align parallel with those of $[A]$, the structure shrinks vertically, and a new vertical synchronization row (grey) is formed on the right. (g) \mathcal{F} is evaluated in parallel on all line segments $[A, i]$ and $[B, j]$, to give the set of all line segments $\mathcal{F}([A, i], [B, j])$ for all $1 \leq i, j \leq n$ represented in blue. (h) The rectangle rearranges into the line $[C]$ of length $O(n^2 \log n)$, as in Figure 15.

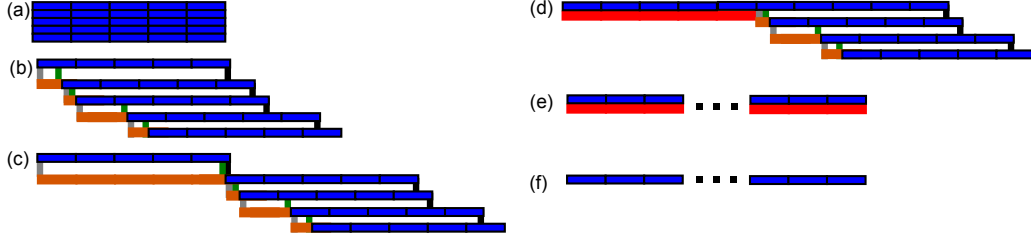


Figure 15: Unfolding a rectangle of n^2 monomer line segments, each of length $O(\log n)$ into a line of length $O(n^2 \log n)$. (a) Initial configuration. (b) A new “insertion line” (shown in orange) grows from the left of each row i . The left end of the orange insertion line at row i is attached to the left end of row $i - 1$ (above) by a monomer shown in grey. As the insertion line for row i grows, it “pushes” row i to the right, relative to row $i - 1$. A monomer (black) attached to the right end of row $i - 1$ and a monomer (green) attached to the right end of the orange insertion line i below are used as “hooks” so that the insertion line is stopped from growing beyond length $O(n \log n)$. (c) The green monomer of row 2 and the black monomer of row 1 become “hooked”. (d) Row 2 moves up to be horizontally aligned with row 1. The grey, green, and black monomers delete. When the orange insertion line in the second row is placed adjacent to row 1, it becomes a red synchronization row. All rows continue this process independently and in parallel. When all are done the insertion line becomes a synchronization row. (e) Ready to synchronize. (f) Final configuration.

this occurs, excess $[B, j]$ segments are deleted (using direct monomer-to-monomer message transfer as before). When this process is complete, we are at Figure 14(e.3).

Next, the duplicates of each $[B, j]$ rotate up to horizontal as shown, and the leftmost copy of $[B]$ deletes itself in a way that vertically “shrinks” the assembly to get Figure 14(f). During this process we make n grey synchronization rows, also shown in Figure 14(f). From Figure 14(f) to Figure 14(g), $\mathcal{F}([A, i], [B, j])$ is computed on each of these n^2 line segments (independently and in parallel), and by the lemma hypotheses this can be done in the allotted space. The horizontal red lines synchronize, and then the vertical red line synchronizes. After this occurs, we can delete the grey synchronization rows and unfold the result into a line, which is of length $O(n^2 \log n)$, to get the final configuration in Figure 14(h), using the technique shown in Figure 15.

Space, state and time analysis. By stepping through the construction (and Figure 14), it is straightforward to check that the entire construction is contained within space $O(n^2 k) \times O(nk)$, and uses $O(1)$ states.

For the time analysis, we first observe that rotation, and copying, of a length ℓ line can each be done in $O(\log \ell)$ expected time via a straightforward analysis [51]. Steps (b), and (c) of Figure 14 involve rotations and copying of lines of length $O(n \log n)$: this completes in expected time $O(\log n)$. The duplication processes of green and purple segments in Figures 14(d) and (e) take $O(\log^2 n)$ expected time. Each application of \mathcal{F} takes expected time $O(k)$, and we apply it independently in parallel n^2 times, hence via Lemma 2, all complete in merely $O(k)$ expected time. There are a number of other places where $\leq n^2$ independent processes, each with expected time $O(\log n)$, take place (deletions in Figures 14(d.4) and (e.3), and rotations in (f)), and by Lemma 2, they take expected time $O(\log n)$. In each of Figures 14(d.2), (e.2) and (g) there are n lines, each of length

$O(n \log n)$ that need to be synchronized. For example, in Figure 14(d.2), synchronization for each red vertical single line takes expected time $O(\log n)$, and since we must wait until all n vertical lines are synchronized (independently), and only then synchronize the horizontal line, this takes expected time $O(\log^2 n)$. Finally, the rearrangement in Figures 14(g) to (h) (given in detail in Figure 15) takes expected time $O(\log^2 n)$: each insertion line must grow $O(n \log n)$ monomers before a level is moved up. There are n of them that work independently, so Lemma 2 gives an expected time to finish of $O(\log^2 n)$. Besides computing \mathcal{F} , the slowest parts of the construction run in expected time $O(\log^2 n)$, and there are at most a constant number of these parts, so the entire construction finishes in expected time $O(k + \log^2 n)$. This concludes the proof of Lemma 15. \square

7.2 Proof of Theorem 14: fast Boolean matrix multiplication

Proof of Theorem 14. The multiplication $C = AB$ of two $n \times n$ Boolean matrices is defined as $C_{i,j} = \bigvee_{k=1}^n (A_{i,k} \wedge B_{k,j})$. To calculate $A_{i,k} \wedge B_{k,j}$, for each i, j, k , we begin by defining the function \mathcal{F}_{AND} which acts on two encoded matrix elements $[A, (i, k_1)]$ and $[B, (k_2, j)]$ as follows.

$$\mathcal{F}_{\text{AND}}([A, (i, k_1)], [B, (k_2, j)]) = \begin{cases} [\emptyset][\tilde{i}][\tilde{j}][\tilde{k}_1][\tilde{k}_2] & k_1 \neq k_2 \\ [\tilde{c}_{i,j,k_1}][\tilde{i}][\tilde{j}][\tilde{k}_1][\tilde{k}_2] & \text{if } k_1 = k_2 \end{cases}$$

where \emptyset is a special monomer denoting “no useful data here”, $[\tilde{c}_{i,j,k_1}]$ is the monomer encoding for the (useful) bit $A_{i,k_1} \wedge B_{k_1,j}$ when $k_1 = k_2$, and as usual $[\tilde{i}]$, $[\tilde{j}]$, $[\tilde{k}_1]$, $[\tilde{k}_2]$ denote the binary monomer line segments encoding of $i, j, k_1, k_2 \leq n$.

We now apply Lemma 15 to $[A]$ and $[B]$ setting \mathcal{F} to \mathcal{F}_{AND} . This gives a line of monomers with n^4 segments, each of length $O(\log n)$, and n^3 of which encode useful data. The remainder are the $n^4 - n^3$ segments ℓ for which $k_1 \neq k_2$. The entire line synchronizes and begins the process of deleting the useless line segments ℓ as follows. Each ℓ encodes an $O(\log n)$ bit number p (as the concatenation of the bit strings for i, j, k_1, k_2). The digits of p are used to organise the deletion of the segments. If the LSB of p in segment ℓ is 1, then ℓ deletes itself. Deletion of a segment works as follows: the rightmost monomer r of ℓ walks on top of its ℓ segment, walking left, sequentially deleting monomers until all of the ℓ segment is deleted. The monomer r is now adjacent to a new segment ℓ' (to the left of the former ℓ) which causes ℓ' to “delete” its LSB (sets its monomer state to \emptyset). The entire line grows a shift synchronization row and the process iterates. To stop the iteration: if an r monomer completes its walk left and meets a non- ℓ segment (i.e. it meets a useful segment) it initiates growth of a lift synchronization row, when all lift synchronization rows form, a lift synchronization occurs signalling that the entire deletion process has finished. This gives a line of monomer segments of the form $[\tilde{c}_{i,j,k_1}][\tilde{i}][\tilde{j}][\tilde{k}_1][\tilde{k}_2]$ with $k_1 = k_2$. Next each of the redundant $[\tilde{k}_2]$ segments deletes itself, then the line synchronizes. This gives a line of n^3 segments $[C, (i, j, k)]$ for $1 \leq i, j, k \leq n$.

To calculate the elements $C_{i,j} = \bigvee_{k=1}^n C_{i,j,k}$, for $1 \leq i, j \leq n$, we begin by sorting the line segments first by i , then by j , then by k . From this sorted line of elements, for each i, j we calculate $[C, (i, j)]$, the encoding of the matrix element $C_{i,j}$, in a two-step process as follows. First, for all i, j, k where $k \neq 1$, the $[\tilde{i}]$ and $[\tilde{j}]$ line segments delete themselves from each of the $[C, (i, j, k)]$ line segments, and the entire line synchronizes when done. The n segments $[\tilde{k}]$, for all $k \leq n$, then rotate perpendicular to their original orientation, and translate (horizontally “shrink”) so that the n monomers of the form $[\tilde{c}_{i,j,k}]$ lie horizontally adjacent to each other. At this point we have a structure consisting of vertical columns of $[\tilde{c}_{i,j,k}]$ and $[\tilde{k}]$, ordered horizontally by i , then by j then by k , for $1 \leq i, j, k \leq n$ (and also with those $[\tilde{k}]$ that encode 1, which still have their horizontal $[\tilde{i}]$

and $[j]$ segments). The set of monomer bitstrings $\{[\tilde{k}] \mid k \leq n\}$ are used to organise iterated pairwise ORing. For each $[\tilde{k}]$ with LSB 1 delete the LSB from the $[\tilde{k}]$ segment and OR its $[\tilde{c}_{i,j,k}]$ bit with its neighbour $[\tilde{c}_{i,j,k'}]$ to the left, with the result bit being stored in the $[\tilde{c}_{i,j,k'}]$ monomer to the left. Then a shift synchronization occurs. This is iterated $O(\log n)$ times until all n bits have been ORed and finished with a lift synchronization. We are left with a line of segments of the form $[\tilde{C}, (i, j)] = [\tilde{c}_{i,j}][\tilde{i}][\tilde{j}]$, for $1 \leq i, j \leq n$, ordered first by i and then by j . This is exactly the monomer representation of the matrix $C = AB$ that we desire.

Time, space and state analysis. The time of Boolean matrix multiplication is dominated by Lemma 15 and the sorting algorithm. Since the expected time of \mathcal{F}_{AND} is $O(\log n)$, then $k = O(\log n)$ in the hypothesis of Lemma 15, giving $O(\log^2 n)$ as the expected time of the application of Lemma 15. There are $O(n^3)$ monomer segments (each of length $O(\log n)$) to be sorted when calculating the ORs, so the expected time for sorting is $O(\log^3 n)$. Hence, the entire matrix multiplication takes expected time $O(\log^3 n)$.

The most space-consuming aspects of Boolean matrix multiplication are Lemma 15 and the sorting algorithm. Lemma 15 takes space $O(m^2 k) \times O(mk)$ for two lines each containing m line segments each of which is of length k . For matrix multiplication, we are starting from two encoded $n \times n$ matrices, each of which has n^2 elements. Setting $m = n^2$ and $k = O(\log n)$, the total space for Boolean matrix multiplication is thus $O(n^4 \log n) \times O(n^2 \log n)$, and since the sorting algorithm takes less space than that, we are done.

A careful analysis of the algorithm shows that the number of monomer states is $O(1)$. \square

8 Boolean circuit simulation

Our main result, Theorem 1 is a restatement of the following theorem. Definition 3 defines what it means to decide languages with the nubot model.

Theorem 16. *Let $L \in \text{NC}^j$ be a language decided by a logspace-uniform Boolean circuit family \mathcal{C} of circuits that have depth $O(\log^j n)$, for some $j \geq 1$, size $O(n^k)$ and input length $|x| = n$, and let $[\tilde{x}]$ denote the representation of $x \in \{0, 1\}^*$ as a line of binary monomers. Then there is a set of nubot rules \mathcal{N}_L such that, for all $x \in \{0, 1\}^*$, starting from an initial configuration containing only $[\tilde{x}]$, \mathcal{N}_L decides whether $x \in L$ and uses space $n^{O(1)} \times n^{O(1)}$, monomer states $O(1)$, and expected time $O(\log^{j+3} n)$.*

The proof is contained in Section 8.2, where we give a nubot algorithm that given x , quickly generates a Boolean circuit, and then simulates that circuit on input x . Before that, in Section 8.1, we present a nubot algorithm that simulates function-computing logspace Turing machines in polylogarithmic expected time. This fast Turing machine simulation will be used in the circuit generation part of Section 8.2.

8.1 Fast parallel simulation of space bounded Turing machines

Here, we give a polylogarithmic expected time simulation of deterministic logspace Turing machines that compute functions with domain and range $\{0, 1\}^*$.

Lemma 17. *Let \mathcal{M} be a deterministic Turing machine that on input $x \in \{0, 1\}^*$, of length $|x| = n$, generates an output $y \in \{0, 1\}^*$, in $O(\log n)$ workspace and time t . There is a set of nubot rules $\mathcal{N}_{\mathcal{M}}$*

such that for all $x \in \{0, 1\}^*$, starting with the initial configuration containing only the line $[\tilde{x}]$ (that represents x), \mathcal{M} computes $[\tilde{y}]$ (the representation of y) using $O(1)$ states, $n^{O(1)} \times n^{O(1)}$ space, and $O(\log^4 n)$ expected time.

Before giving the proof of Lemma 17 we state some assumptions about \mathcal{M} :

1. \mathcal{M} follows the standard conventions for logspace Turing Machines: there are 3 tapes: a read-only input tape, a $O(\log n)$ space bounded work tape, and a write-only output tape whose head moves in one direction only.
2. \mathcal{M} uses the alphabet $\{0, 1\}$ on all 3 tapes (the input is delimited with the symbol $\#$).
3. A configuration consists of the input tape head position and read symbol, worktape contents and head position, worktape read symbol, and machine state, and (unusually⁵) the output tape head position and write symbol.
4. \mathcal{M} always ends its computation in a halting, accept state.

There are n possible positions for the head on the input tape and $O(\log n)$ head positions on the work tape. We note that each configuration of \mathcal{M} can be written as a string over $\{0, 1\}^*$ of length $O(\log n)$. This follows from the fact that in a given configuration $O(\log n)$ bits describe the position of the input and output tape heads, $O(\log \log n)$ bits describe the position of the worktape head, $O(\log n)$ bits describe the contents of the work tape, and $O(1)$ symbols describe the read and write symbols on the various tapes, and the machine state. Thus, on length- n input, \mathcal{M} visits at most $2^{O(\log n)} = n^{O(1)}$ configurations before halting, or looping forever, in other words $t = n^{O(1)}$. We next define the *configuration matrix* of a space-bounded Turing machine.

Definition 18 (Configuration matrix). *Let \mathcal{M} be a deterministic Turing machine with space bound $s(n)$. Consider the set of all $k = n^{O(1)}$ possible configurations on a length n input (we include all syntactically valid configurations for a worktape with $s(n)$ tape cells, even though on a given input many will be unreachable). We define M to be the $k \times k$ Boolean matrix where for $1 \leq i, j \leq k$, $m_{i,j} = 1$ if and only if there exists a one-step transition from configuration c_i to configuration c_j , and $m_{i,j} = 0$ otherwise.*

Proof of Lemma 17. A logarithmic space-bounded Turing machine \mathcal{M} can be simulated efficiently in parallel (in polynomial time, using polynomial processors/resources) in a variety of parallel models by iterated squaring of \mathcal{M} 's Boolean configuration matrix M [38]. Specifically, we can determine whether there exists a sequence of one-step transitions from any configuration c_i to any c_j by beginning with matrix M , computing $M := M^2 + M$, and iterating this procedure $O(\log n)$ times. A path between the two configurations exists only if entry $m'_{i,j} = 1$ in the resulting matrix M' . We call matrix M' the *path-complete matrix*. Since \mathcal{M} is deterministic and always accepts (\mathcal{M} is total: for any input 1^n , $n \in \mathbb{N}$ it outputs a circuit c_n), there is exactly one path, through \mathcal{M} 's configuration graph, that leads from the start configuration to the halt (accept) configuration. The technique of iterated squaring is sufficient for simulating Turing machines that decide languages, but here we want to simulate a function-computing machine. We do this by modifying the iterated squaring technique: our configurations contain (extra) information about what is written to the output tape,

⁵Our configurations include an output tape write symbol and an output tape head position which is not standard practice [38], but will be useful in our construction.

we appropriately extract this information during our simulation of \mathcal{M} by iterated squaring. The remainder of the proof describes how we do all of this in the nubot model.

We generate all possible configurations of Turing machine \mathcal{M} , in parallel. First, we build a counter that counts up to $2^{s(n)} = n^{O(1)}$, the upper bound on the number of distinct worktape contents of \mathcal{M} . Once completed, each row of the counter then generates its own counter, counting up to the number of different positions that the head can be on the input tape. This process is iterated for each of the (constant number of) attributes in a Turing machine configuration to give a final counter with $k = n^{O(1)}$ rows, one for each distinct configuration (see Definition 18 for k , and see [51] for details on efficiently growing a counter). The counter backbone synchronizes, giving a $k \times O(\log k)$ rectangle, whose bond structure forms a “comb”. The counter rearranges itself into a line $[C] = [C, 1][C, 2] \dots [C, k]$ where line segment $[C, i]$ encodes configuration c_i .

We next use our encoding $[C]$ of all possible configurations to generate an encoding of the configuration matrix M . First, $[C]$ is copied so that we have two parallel instances of $[C]$, side-by-side. Next, we apply Lemma 15, setting $\mathcal{F} = \mathcal{F}_{\mathcal{M}}$ where $\mathcal{F}_{\mathcal{M}}$ takes as input the pair of parallel line segments $[C, i]$ and $[C, j]$, and a copy of the input line segment⁶ $[\tilde{x}]$ that encodes x , and returns a segment $[M, (i, j)] = [m, i, j][C, i][C, j]$ where $[m, i, j]$ is a binary nubot monomer representing element $m_{i,j}$ in \mathcal{M} ’s configuration matrix⁷. In other words, given the encoding of two configurations c_i, c_j , the function $\mathcal{F}_{\mathcal{M}}$ determines if there is a one-step transition from c_i to c_j via Turing machine \mathcal{M} . $\mathcal{F}_{\mathcal{M}}$ works by straightforward message-passing and state changes from monomer to monomer along the pair of encoded configurations. To satisfy the hypotheses of Lemma 15, $\mathcal{F}_{\mathcal{M}}$ should work in time linear in the encoded configurations’ length ($|[C, i]|, |[C, j]|$) which is easily achieved. It is also the case that the space and states bound in the hypotheses of Lemma 15 are met by $\mathcal{F}_{\mathcal{M}}$. After applying Lemma 15 we get an encoding of the configuration matrix M as a single line $[M]$ of consecutive line segments $[M, (i, j)] = [m, i, j][C, i][C, j]$ for $1 \leq i, j \leq k$.

We make a copy of the encoded matrix $[M]$ (a line of monomers) and then use Theorem 14 to square $[M]$, giving $[M^2]$ (another line of monomers). After we have both lines $[M]$ and $[M^2]$, “adding” the lines together (to compute $M^2 + M$) is easy: matrix elements with the same (i, j) coordinates are adjacent when $[M]$ and $[M^2]$ are orientated parallel and next to each other, so the addition can be carried out “locally”. The iterated squaring (and addition) are carried out $1 + \log k = O(\log n)$ times. The result is an encoding of M ’s path-complete matrix M' .

Consider the path-complete configuration matrix M' , with start configuration c_{start} and halt configuration c_{halt} . We need to (i) determine which configurations are on the unique path, in the configuration graph, from c_{start} to c_{halt} , and (ii) follow this unique path keeping track of what was written to the output tape at each step. For any i , if configuration c_{start} leads to configuration c_i in ≥ 1 steps then $M'_{c_{\text{start}}, c_i} = 1$. Similarly, if configuration c_i leads to configuration c_{halt} in one or more steps then $M'_{c_i, c_{\text{halt}}} = 1$. Hence it is sufficient to extract row c_{start} and column c_{halt} from M' , and compare them, in order to find the entire path of configurations from c_{start} to c_{halt} .

We do this by first deleting all encoded matrix elements of $[M']$ that are not in row c_{start} or not in column c_{halt} . This results in two lines of monomers, that are then aligned parallel and side-by-side. Next each entry i is compared, if there is a 1 in both we keep the entry, otherwise the entry is deleted. We are left with the list of configurations on the path from c_{start} to c_{halt} . This line of

⁶Each configuration is of length polynomial in $|x| = O(|[\tilde{x}]|)$, hence including $[\tilde{x}]$ here does not change the asymptotics.

⁷The line segments in an encoded matrix usually encode the matrix element’s (i, j) coordinates, here we do things slightly differently: we are using encoded configurations, rather than natural numbers, as the matrix indices. This simplifies our constructions a little.

monomers synchronizes.

Next, the encoded matrix entries (bits) are deleted leaving the list of encoded configurations. Configurations that do not write anything to the output tape are deleted. The remaining configurations are sorted in increasing order of output-tape write location. Since the output tape head moves one way only, this gives the list of outputting configurations in the order they are executed by the Turing machine \mathcal{M} . Finally, all monomers that do not represent a symbol written by the output tape head are deleted, and the result is compressed into a line. We are left with a monomer line encoding y , the output tape contents.

State, space and time analysis of Turing machine simulation. The state complexity of $O(1)$ can be seen from stepping through the algorithm. In the proof, the configuration matrix dimension size is $k \times k$, where $k = n^{O(1)}$. From Theorem 14, matrix multiplication for two $k \times k$ matrices takes space $O(k^4 \log k) \times O(k^2 \log k)$ using nubots, and since $k = n^{O(1)}$ this gives the space bound in the lemma statement. The space complexity of our Turing machine simulation is dominated by this.

For the time analysis, first note that generating the configurations consists of running a counter that takes expected time $O(\log^2 n)$, see [51] for details. After the configurations are generated, each iteration of matrix multiplication, addition, and deletions is bounded by time $O(\log^3 n)$. Since we do $O(\log n)$ matrix multiplications and additions the expected time is $O(\log^4 n)$ using nubots. The expected time of the other rearrangements and computations during the construction is dominated by that of matrix multiplication. This completes the proof of Lemma 17. \square

8.2 Generating and simulating a Boolean circuit: proof of Theorem 16

We define the nubot monomer encoding of gates and Boolean circuits. Boolean circuits were defined in Section 2.2.

Definition 19 (Nubot monomer encoding of a gate). *The encoding \tilde{g} of a Boolean circuit gate g is as follows: a single gate monomer encodes the gate type (AND, OR, or NOT), directly above the gate monomer are k line segments of monomers called result segments where k is the gate's out-degree. Each result line segment encodes a destination gate number using $O(\log n)$ binary monomers, where n is the circuit size. There is an empty region of height $O(\log n)$ below the gate monomer called the input region (Figure 16(b): dotted blue regions).*

A gate is simulated as follows. The input region of \tilde{g} is an empty region to which a line of monomers, that encode the inputs to g , can attach (Figure 1(d): dotted blue regions). Upon attachment of the input lines, the gate monomer computes g 's Boolean function. Let g be a gate with out-degree 2, and which outputs to the gates g_1 and g_2 . The result region of nubot gate \tilde{g} consists of two lines of binary monomers that encode the wires that lead to g_1 and g_2 (Figure 1(d): solid blue regions). The simulation of wires is covered in the proof below.

Definition 20 (Nubot monomer encoding of a Boolean circuit). *A Boolean circuit c is encoded as a nubot configuration consisting of the encoded gates (Definition 19) written in layers, one for each layer in c (see Figure 1). Within a layer, the encoded gates are horizontally spaced apart by the circuit size.*

Proof of Theorem 16. The proof has two parts, circuit generation and circuit simulation.

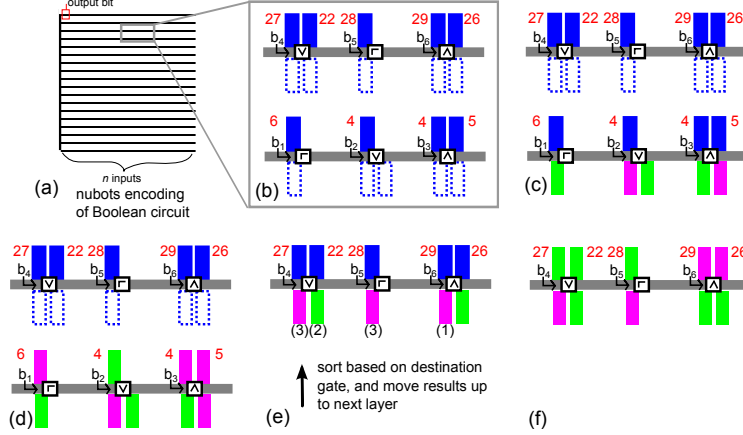


Figure 16: Nubots configuration that simulates a Boolean circuit. Dotted blue regions denote gate input regions. Blue rectangles denote gate result regions. The gate address encoded by a line segment is in red. Gates are monomers. Pink encodes 0, green encodes 1. (a) Initial configuration that encodes the entire circuit and its input. (b) Zoom-in shown the initial configuration of 6 gates. (c) Input line segments arrive at the gate. (d) The gate monomers calculate their respective gates’ result, and communicate the result bit to the result segments (shown as colour change). (e) The input monomers and grey line are deleted. The result monomers sort themselves using the sorting algorithm (not shown) and are moved upwards to be the input to the next layer. The gate from which each input monomer originates is shown in parenthesis (these values came from gates 3, 2, and 1—not shown). (f) The gate monomers of the next layer calculate their result values.

8.2.1 Circuit generation

Let $c_n \in \mathcal{C}$ be a Boolean circuit with n input gates that we wish to simulate. From the theorem statement \mathcal{C} is uniform by logspace Turing machine \mathcal{M} . To generate the encoding of c_n as nubot monomers, first, \mathcal{M} on input 1^n is simulated via Lemma 17 to give a line of monomers $[\hat{c}_n]$ that encodes $c_n = \mathcal{M}(1^n)$. Next, this “linear” encoding of c_n geometrically unfolds into a two-dimensional “ladder” format, with one encoded circuit layer per rung, as shown in Figure 1(c) and defined in Definition 20. We use the folding technique from [51] that takes expected time $O(\log^2 \ell)$ to fold a length ℓ line into a square (here we modify the technique to fold a line into a comb, then on the teeth of the comb the gate result monomers fold out from each of the teeth to give the structure in Figure 1). Since $||[\hat{c}_n]||$ is polynomial in n , the rearrangement happens in expected time $O(\log^2 n)$.

8.2.2 Circuit simulation

We have a nubot configuration that encodes a circuit as shown in Figure 16(a). We evaluate the encoded Boolean circuit layer by layer, from input layer (bottom) to output layer (top), with each layer being evaluated in parallel. Evaluating a layer is a 3-step process shown in Figures 16(c), (d) and (e). First assume we have the configuration shown in Figure 16(c): along the bottom grey line there are gate monomers (that encode AND, OR, or NOT), and below the bottom grey line there are pink and green line segments of monomers that encode gate input bits 0 and 1 respectively, and

above are blue line segments that encode the wires out of these gates (i.e. destination addresses to the next layer above). Evaluation of a gate is straightforward: since the gates have fan-in ≤ 2 , the gate simply reads the 1 or 2 pink/green line segments below by reading its lower neighbours' states. The gate monomer computes the encoded gate's Boolean function and passes the resulting bit to the result line segments above. Note that gates may have fan-out (or out-degree) as large as the circuit size, i.e. polynomial in input length, so for this we assume adjacent gates on a layer are spaced at least as far apart horizontally as the circuit size. Then a gate communicates its result to all of them via shift or lift synchronization (in expected time logarithmic of circuit size). After all gate monomers in a layer have completed this process, they synchronize. By now we have reached Figure 16(d).

Boolean circuits may be non-planar and so have crossing wires when drawn in 2D, hence naïvely moving bit-encoding monomers in the plane to the next layer above may cause unintended collisions. We resolve this problem using our nubot sorting algorithm from Section 6. After a layer has synchronized, the (blue) line segments in the gate result regions of that entire layer are organised into a horizontal line, to serve as input to our sorting procedure. These gate result regions are then sorted by increasing wire number. The gates on the next layer above are assumed to be encoded in increasing (gate index) order. After sorting, the gate result regions are aligned with the gate above them (using counters) and pushed vertically upwards to the relevant gate. This is done in such a way that when it is finished there are no monomers below the new layer (any excess monomers are deleted); this deletion leaves enough space below for the sorting algorithm on the next iteration.

After the monomer that encodes the circuit's unique output gate computes its result bit, it destroys itself, leaving a single monomer encoding the output bit. No rules are applicable and so the system has halted with its answer.

Time, state, and space analysis of circuit simulation. There are $O(1)$ gate types, and all numbers are written using binary monomers. All other parts of the construction from previous sections use $O(1)$ states. By stepping through the simulation with this in mind it is straightforward to obtain a state complexity of $O(1)$.

We are simulating a circuit of size $O(n^k)$ and depth $O(\log^j n)$. Each layer of the circuit is encoded as a monomer layer of height $O(\log n)$, giving a total height of $O(\log^{j+1} n)$ for the encoded circuit. The width of an encoded layer is $O(n^{2k})$ which comes from the circuit size being $O(n^k)$, and from each gate being horizontally separated by a further $O(n^k)$ to handle fan-out (note a horizontal separation of a mere $O(\log n)$ monomers is sufficient for the sorting algorithm). This gives $O(n^{2k}) \times O(\log^{j+1} n)$ space to lay out the circuit. However, sorting n numbers, each written as length $O(\log n)$ bit strings, takes space $O(n \log n) \times O(n)$. Thus the total space complexity for the circuit simulation is $O(n^{2k}) \times O(n)$. Lemma 17 tells us that circuit generation takes space $n^{O(1)} \times n^{O(1)}$ (the hidden constants are coming from the logspace bounded Turing machine), which dominates the total space for both circuit generation and circuit simulation.

The asymptotically slowest part of simulating a circuit layer is the sorting algorithm, which takes expected time $O(\log^3 n)$ per layer. There are $O(\log^j n)$ layers, thus the total expected time for the simulation is $O(\log^{j+3} n)$. The circuit generation takes time $O(\log^4 n)$ from Lemma 17, but since we assumed that $j \geq 1$ in the statement of Theorem 16, this leaves the total expected time for both circuit generation and circuit simulation at $O(\log^{j+3} n)$. This completes the proof of Theorem 16. \square

Acknowledgments.

We thank Erik Winfree for valuable discussion and suggestions on our results, Paul Rothmund for stimulating conversations on molecular muscle, Niall Murphy for informative discussions on circuit complexity theory, and Dhiraj Holden and Dave Doty for useful discussions. Damien thanks Beverley Henley for introducing him to developmental biology many moons ago.

References

- [1] G. Aggarwal, Q. Cheng, M. H. Goldwasser, M.-Y. Kao, P. M. de Espanes, and R. T. Schweller. Complexities for generalized models of self-assembly. *SIAM Journal on Computing*, 34:1493–1515, 2005.
- [2] E. Allender and M. Koucký. Amplifying lower bounds by means of self-reducibility. *JACM*, 57(3):14:1–36, 2010.
- [3] G. Aloupis, S. Collette, M. Damian, E. Demaine, R. Flatland, S. Langerman, J. O’Rourke, V. Pinciu, S. Ramaswami, V. Sacristán, and S. Wuhler. Efficient constant-velocity reconfiguration of crystalline robots. *Robotica*, 29(1):59–71, 2011.
- [4] G. Aloupis, S. Collette, E. D. Demaine, S. Langerman, V. Sacristán, and S. Wuhler. Reconfiguration of cube-style modular robots using $O(\log n)$ parallel moves. In *ISAAC: Proc. of the 19th Annual International Symposium on Algorithms and Computation*, pages 342–353, 2008.
- [5] F. Becker, E. Remila, and I. Rapaport. Self-assembling classes of shapes, fast and with minimal number of tiles. In *Proceedings of the 26th Conference on Foundations of Software Technology and Theoretical Computer Science (FSTTCS 2006)*, volume 4337 of *LNCS*, pages 45–56. Springer, Dec. 2006.
- [6] Z. Butler, R. Fitch, and D. Rus. Distributed control for unit-compressible robots: goal-recognition, locomotion, and splitting. *IEEE/ASME Transactions on Mechatronics*, 7:418–430, 2002.
- [7] S. Cannon, E. D. Demaine, M. L. Demaine, S. Eisenstat, M. J. Patitz, R. T. Schweller, S. M. Summers, and A. Winslow. Two hands are better than one (up to constant factors): Self-assembly In the 2HAM vs. aTAM. In *STACS: 30th International Symposium on Theoretical Aspects of Computer Science*, pages 172–184, 2013.
- [8] H. Chandran, N. Gopalkrishnan, and J. Reif. Tile complexity of approximate squares. *Algorithmica*, 66(1):1–17, 2012.
- [9] A. Condon. A theory of strict P-completeness. *Computational Complexity*, 4(3):220–241, 1994.
- [10] N. Dabby and H.-L. Chen. Active self-assembly of simple units using an insertion primitive. In *SODA: Proceedings of the Twenty-fourth Annual ACM-SIAM Symposium on Discrete Algorithms*, pages 1526–1536, Jan. 2012.
- [11] E. Demaine, M. Demaine, S. Fekete, M. Ishaque, E. Rafalin, R. Schweller, and D. Souvaine. Staged self-assembly: nanomanufacture of arbitrary shapes with $O(1)$ glues. *Natural Computing*, 7(3):347–370, 2008.
- [12] E. D. Demaine, M. L. Demaine, S. P. Fekete, M. J. Patitz, R. T. Schweller, A. Winslow, and D. Woods. One tile to rule them all: simulating any Turing machine, tile assembly system, or tiling system with a single puzzle piece. In *ICALP: Proceedings of the 41st International Colloquium on Automata, Languages, and Programming*, volume 8572 of *LNCS*, pages 368–379. Springer, 2014. Arxiv preprint [arXiv:1212.4756](https://arxiv.org/abs/1212.4756) [cs.DS].
- [13] E. D. Demaine, S. Eisenstat, M. Ishaque, and A. Winslow. One-dimensional staged self-assembly. In *DNA: 17th Intl Conf on DNA Computing & Molecular Programming*, volume 6937 of *LNCS*, pages 100–114, Pasadena, CA, 2011.

- [14] E. D. Demaine, M. J. Patitz, T. A. Rogers, R. T. Schweller, S. M. Summers, and D. Woods. The two-handed tile assembly model is not intrinsically universal. In *ICALP: 40th International Colloquium on Automata, Languages and Programming. Proceedings, part 1*, volume 7965 of *LNCS*, pages 400–412, Riga, Latvia, July 2013. Arxiv preprint [arXiv:1306.6710](#) [cs.CG].
- [15] D. Doty. Randomized self-assembly for exact shapes. *SICOMP*, 39:3521, 2010.
- [16] D. Doty, J. H. Lutz, M. J. Patitz, R. T. Schweller, S. M. Summers, and D. Woods. The tile assembly model is intrinsically universal. In *FOCS: Proceedings of the 53rd Annual IEEE Symposium on Foundations of Computer Science*, pages 439–446, Oct. 2012.
- [17] D. Doty, J. H. Lutz, M. J. Patitz, S. M. Summers, and D. Woods. Intrinsic universality in self-assembly. In *STACS: Proceedings of the 27th International Symposium on Theoretical Aspects of Computer Science*, pages 275–286, 2009.
- [18] B. Fu, M. Patitz, R. Schweller, and B. Sheline. Self-assembly with geometric tiles. In *ICALP: The 39th International Colloquium on Automata, Languages and Programming*, volume 7391 of *LNCS*, pages 714–725. Springer, 2012.
- [19] D. Gillespie. A rigorous derivation of the chemical master equation. *Physica A: Statistical Mechanics and its Applications*, 188(1):404–425, 1992.
- [20] R. Greenlaw, H. J. Hoover, and W. L. Ruzzo. *Limits to parallel computation: P-completeness theory*. Oxford University Press, USA, 1995.
- [21] J. Hendricks, J. E. Padilla, M. J. Patitz, and T. A. Rogers. Signal transmission across tile assemblies: 3D static tiles simulate active self-assembly by 2D signal-passing tiles. In *DNA19: The 19th International Conference on DNA Computing and Molecular Programming*, volume 8141 of *LNCS*. Springer, 2013. Arxiv preprint: [arXiv:1306.5005](#) [cs.ET].
- [22] J. Hendricks and M. J. Patitz. On the equivalence of cellular automata and the tile assembly model. In *MCU: Proceedings of Machines, Computations and Universality*, volume 128, pages 167–189, University of Zürich, Switzerland. September 9-12, 2013. Electronic Proceedings in Theoretical Computer Science.
- [23] N. Jonoska and D. Karpenko. Active tile self-assembly, self-similar structures and recursion. 2012. Arxiv preprint [arXiv:1211.3085](#) [cs.ET].
- [24] N. Jonoska and G. L. McColm. Complexity classes for self-assembling flexible tiles. *Theoretical Computer Science*, 410(4):332–346, 2009.
- [25] M. Kao and R. Schweller. Reducing tile complexity for self-assembly through temperature programming. In *SODA: Proceedings of the Seventeenth Annual ACM-SIAM Symposium on Discrete Algorithms*, pages 571–580. ACM, 2006.
- [26] M. Kao and R. Schweller. Randomized self-assembly for approximate shapes. In *ICALP: Proceedings of the 35th International Colloquium on Automata, Languages and Programming*, volume 5125 of *LNCS*, pages 370–384. Springer, 2008.
- [27] A. Keenan, R. Schweller, M. Sherman, and X. Zhong. Fast arithmetic in algorithmic self-assembly. In *UCNC : The 13th International Conference on Unconventional Computation and Natural Computation*, July 2014. Arxiv preprint [arXiv:1303.2416](#) [cs.DS].
- [28] E. Klavins. Directed self-assembly using graph grammars. In *Foundations of Nanoscience: Self Assembled Architectures and Devices*, Snowbird, UT, 2004.
- [29] C. Malchik and A. Winslow. Tight bounds for active self-assembly using an insertion primitive. In *ESA: The 22nd European Symposium on Algorithms*, 2014. Accepted. Arxiv preprint [arXiv:1401.0359](#) [cs.FL].
- [30] A. C. Martin, M. Kaschube, and E. F. Wieschaus. Pulsed contractions of an actin–myosin network drive apical constriction. *Nature*, 457(7228):495–499, 2008.

- [31] P.-E. Meunier, M. J. Patitz, S. M. Summers, G. Theyssier, A. Winslow, and D. Woods. Intrinsic universality in tile self-assembly requires cooperation. In *SODA: ACM-SIAM Symposium on Discrete Algorithms (2014)*, pages 752–771, 2014. Arxiv preprint: [arXiv:1304.1679](https://arxiv.org/abs/1304.1679) [cs.CC], 2013.
- [32] S. Murata and H. Kurokawa. Self-reconfigurable robots. *Robotics & Automation Magazine, IEEE*, 14(1):71–78, 2007.
- [33] N. Murphy, T. J. Naughton, D. Woods, B. Henley, K. McDermott, E. Duffy, P. J. van der Burgt, and N. Woods. Implementations of a model of physical sorting. *International Journal of Unconventional Computing*, 4(1):3–12, 2008.
- [34] N. Murphy and D. Woods. AND and/or OR: Uniform polynomial-size circuits. In *MCU: Machines, Computations and Universality*, volume 128, pages 150–166. EPTCS, Sept. 2013. [arXiv:1212.3282](https://arxiv.org/abs/1212.3282) [cs.CC].
- [35] T. Neary and D. Woods. P-completeness of cellular automaton rule 110. In *ICALP: The 33rd International Colloquium on Automata, Languages and Programming*, volume 4051 of *LNCS*, pages 132–143. Springer, 2006.
- [36] J. Padilla, W. Liu, and N. Seeman. Hierarchical self assembly of patterns from the Robinson tilings: DNA tile design in an enhanced tile assembly model. *Natural Computing*, 11(2):323–338, 2011.
- [37] J. Padilla, M. Patitz, R. Pena, R. Schweller, N. Seeman, R. Sheline, S. Summers, and X. Zhong. Asynchronous signal passing for tile self-assembly: Fuel efficient computation and efficient assembly of shapes. In *Unconventional Computation and Natural Computation*, volume 7956 of *LNCS*, pages 174–185. Springer, 2013.
- [38] C. M. Papadimitriou. *Computational complexity*. Addison-Wesley, 1994.
- [39] M. J. Patitz. An introduction to tile-based self-assembly. In *Unconventional Computation and Natural Computation*. 7445: 34–62, *LNCS*, Springer, 2012.
- [40] P. Prusinkiewicz and A. Lindenmayer. *The algorithmic beauty of plants*. Springer-Verlag, 1990.
- [41] J. Reif and S. Slee. Optimal kinodynamic motion planning for 2D reconfiguration of self-reconfigurable robots. *Robot. Sci. Syst.*, 2007.
- [42] P. W. K. Rothmund and E. Winfree. The program-size complexity of self-assembled squares (extended abstract). In *STOC: Proceedings of the thirty-second annual ACM symposium on Theory of computing*, pages 459–468. ACM Press, 2000.
- [43] D. Rus and M. Vona. Crystalline robots: Self-reconfiguration with compressible unit modules. *Autonomous Robots*, 10(1):107–124, 2001.
- [44] D. Schultes. Rainbow sort: sorting at the speed of light. *Natural Computing*, 5(1):67–82, 2006.
- [45] D. Soloveichik, M. Cook, E. Winfree, and J. Bruck. Computation with finite stochastic chemical reaction networks. *Natural Computing*, 7(4):615–633, 2008.
- [46] S. Summers. Reducing tile complexity for the self-assembly of scaled shapes through temperature programming. *Algorithmica*, pages 1–20, 2012.
- [47] H. Vollmer. *Introduction to Circuit Complexity: A Uniform Approach*. Springer-Verlag, 1999.
- [48] E. Winfree. *Algorithmic Self-Assembly of DNA*. PhD thesis, California Institute of Technology, June 1998.
- [49] D. Woods. Upper bounds on the computational power of an optical model of computation. In *Algorithms and Computation*, pages 777–788. Springer, 2005.
- [50] D. Woods. Intrinsic universality and the computational power of self-assembly. In *MCU: Proceedings of Machines, Computations and Universality*, volume 128 of *Electronic Proceedings in Theoretical Computer Science*, pages 16–22, Univ. of Zürich, Switzerland. Sept. 9–12, 2013. Open Publishing Association. [dx.doi.org/10.4204/EPTCS.128.5](https://doi.org/10.4204/EPTCS.128.5).

- [51] D. Woods, H.-L. Chen, S. Goodfriend, N. Dabby, E. Winfree, and P. Yin. Active self-assembly of algorithmic shapes and patterns in polylogarithmic time. In *ITCS'13: Proceedings of the 4th conference on Innovations in Theoretical Computer Science*, pages 353–354. ACM, 2013. Full version: [arXiv:1301.2626](#) [cs.DS].
- [52] D. Woods and T. J. Naughton. Parallel and sequential optical computing. In *Optical supercomputing*, volume 5172 of *LNCS*, pages 70–86. Springer, 2008.
- [53] B. Yurke, A. J. Turberfield, A. P. Mills, Jr., F. C. Simmel, and J. L. Nuemann. A DNA-fuelled molecular machine made of DNA. *Nature*, 406:605–608, 2000.