2D DNA Self-Assembly for Satisfiability

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Abstract. DNA self-assembly has been proposed as a way to cope with huge combinatorial NP-HARD problems such as satisfiability. However, the algorithmic designs proposed so far either involve many biosteps or are highly dependent on the particular instance to be solved. This paper presents an algorithmic design for solving satisfiability problems using two-dimensional DNA self-assembly (tiling). The main driving factor in this work was the design and encoding of the algorithm in a general way that separates the algorithm from the data and minimizes the dependency on particular instances. In effect, a large amount of work and preparation can be done in advance as a batch process. In practice, it is likely that the total time for computation will be decreased significantly and laboratory procedures will be simplified.

1. The Satisfiability (SAT) Problem

The Boolean Satisfiability (SAT) problem is the most well known representative of the NP-HARD and NP-COMPLETE classes of problems. These problems require non-deterministic polynomial computation time for optimal solutions. As a result, they are theoretically 'intractable' and, as the instance size grows, they very quickly become impossible to solve on conventional digital computers.

An instance of the SAT problem consists of a number of Boolean variables \( x_1, x_2, \ldots, x_m \) and a number of clauses \( C_1, C_2, \ldots, C_n \). Each clause is a disjunction of distinct literals, whereby a literal is a single variable \( x_i \) or its negation \( \overline{x}_i \). A solution to the problem is a satisfying assignment, that is an assignment of binary truth values to the variables \( x_i \) such that the conjunction of all clauses is satisfied. Boolean formulas represented in this particular format are said to be in Conjunctive Normal Form (CNF); a conjunction of disjunctions.

An example with 5 variables and 8 clauses is given below (+ indicates disjunction; clauses are enclosed by a set of parentheses):

\[
(x_1 + x_2 + x_3)(x_2 + x_4)(x_1 + x_5)(x_1 + x_2 + x_3 + x_4 + x_5)(x_3)(x_2 + x_5)(x_5)(x_1)
\]

In this case, the assignment \((x_1, x_2, x_3, x_4, x_5) = (1, 1, 0, 1, 0)\) is satisfying and, therefore, a solution to the problem.

In general, the number of literals in a single clause is limited only by the total number of variables. However, even if it is required that each clause has exactly \( k \) literals \((k \geq 3)\), the problem (known as \( k\text{-CNF-SAT} \)) is still intractable. Moreover, it is proven that any SAT instance can be turned into an equivalent \( k\text{-CNF-SAT} \) instance [GJ79]. Among the most notable is the \( 3\text{-CNF-SAT} \) variation, because of the small-sized clauses and the low bound on the number of possible clauses. Notice that given \( m \) variables, there can be at most

\[
\binom{m}{k} \times 2^k
\]
possible clauses in a $k$CNF-SAT formula (choose $k$ variables and for each one either leave intact or negate). In particular, for $k = 3$, there are

$$\binom{m}{3} \times 2^3 = \frac{8}{6} \times m(m - 1)(m - 2) = O(m^3)$$

possible clauses in a 3CNF-SAT formula.

The number of satisfying assignments for a given formula can be anywhere between 0 and $2^m$. If it is 0, the formula is said to be unsatisfied, whereas if it is $2^m$, the formula is a tautology. In all other cases, the formula is simply satisfiable.

2. DNA Computation and Satisfiability

The basic idea is to exploit the massive parallelism possible in DNA operations in order to emulate a non-deterministic device that solves the SAT problem in polynomial time. Such emulation can be achieved by exponential-order parallelism.

Consider a particular assignment to the boolean variables in a CNF-SAT formula. On a conventional computer it is fairly easy to check whether this particular assignment is a solution to the problem, i.e. an assignment that satisfies all the clauses in the formula. In fact, this can be done in time linear in the size of the formula. It is the huge (exponential) number of different possible assignments and thus the huge sequence of checks that makes the problem difficult on a conventional computer. If there was a way to perform this check on a DNA-based computer, all checks could be done in parallel by an exponential number of computers, completing the whole computation in polynomial time.

This work proposes a way to perform the checking procedure above on molecular substrate using 2D DNA self-assembly. By creating billions of billions of copies of the participating DNA structures (tiles, in our case), we expect that the procedure will run in parallel on all possible assignments. The assignments will be created dynamically as part of the assembly. In effect, that will make the computation time linear in the size of the formula, while pushing the exponential dimension of the problem into the large number of DNA assemblies, and thus into the space (volume) occupied by the DNA molecules. If there is a satisfying assignment, we expect that at least one of these parallel checks will discover it.

The rest of this section reviews the main proposals for biomolecular solutions to the SAT problem, briefly describes the general DNA structures and methods that are used here and delineates our work.

2.1. Related Work. Lipton was the first to propose a DNA model for satisfiability [Lip95]. His proposal follows Adleman’s elimination method [Adl94], whereby the whole combinatorial space of solutions is created and subsequently the “good” ones are extracted by a series of separation steps. Lipton’s method was refined in [BDLS99]. Later, Hagiya et al. [HAK+99] presented an approach to evaluate and learn $\mu$-formulas (a particular form of Boolean formulas whereby each variable occurs at most once) using a technique that has become known as Whiplash PCR. This method was improved by Winfree [Win98b] to account for general CNF formulas. Finally, a proposal for CNF-SAT using hairpin microcrossover DNA tiles and linear assembly can be found in [WER99].

In Lipton’s method the sequence of separate/extract and combine/union operations performed on the DNA molecules is determined by the problem instance. As a result the number of biosteps required is linear in the size of the formula. In the other two approaches, the number of required biosteps is constant. It is generally desirable to perform as less biosteps as possible due to the error rate and inaccuracies associated with such operations. “Single pot” DNA computations seem to be more promising in this sense.

The initial DNA solution in Lipton’s method is not dependent on the problem instance. A feature of the other approaches and a potential practical problem is that construction of the participating DNA structures cannot really begin until a particular SAT instance is given at hand. This “instance-specific” design implies long total computation time (encoding/computation/decoding). In addition, all required man/machine resources need to be employed again and again as new instances are provided. This may also increase the likelihood of inadvertent encoding errors, which if occur will render the whole computation useless.
It is highly desirable for practical reasons to keep the number of biosteps small and the participating DNA structures (strands, tiles, etc.) as simple as possible. This will also allow experimentalists to do more experimental computations without being much concerned about costly operations and/or (re)synthesis of complex DNA structures. Obviously, there is a trade-off between these two goals.

A balanced solution to these problems, would possibly be an algorithmic design that requires minimal encoding for a given instance (some straightforward description of the input), whereas the main algorithm is coded in preconstructed “instance-independent” DNA molecules that can be created and even tested off-line in a batch fashion.

The DNA self-assembly technique employed here follows concepts and proposals from a model known as DNA tiling computation originally proposed by Winfree [Win98a] for $O(1)$ DNA computations. Basic components for DNA tiling have been prototyped and tested by Seeman and colleagues. In particular, double crossover molecules have been shown to be rigid and able to form planar lattices containing hundreds of thousands of tiles [WLWS98]. Winfree has shown how to solve the Hamiltonian Path problem using 2-dimensional DNA tiling [Win98a]. Further, LaBean, Winfree and Reif [LWR99] have been experimenting with parallel XOR and addition operations using DNA tiling.

2.2. 2D DNA Self-Assembly for Satisfiability. The main goal of the algorithmic design presented here is to utilize the computational power of DNA tiling to solve the SAT problem, while minimizing the dependency of the process on particular problem instances. In other words, it is an attempt to separate the algorithm (instance-independent) from the data (instance-dependent). This can be accomplished by coding the general algorithm as a library of preconstructed non-specific DNA tiles which, when combined with an appropriate encoding for a given instance (input), would perform the desired computation. A separation step afterwards could separate the successful computations from where a satisfying assignment could be drawn. Put another way, we would like an encoding that specifies the general algorithm for the problem and not one that specifies a specialized algorithm for solving a particular instance of the problem.

Our design is described at the algorithmic level. We abstract each DNA tile as a square with labels at the corners (see figure 1 below). Each label indicates a particular kind of sticky end. Two sticky ends that can complement in the Watson-Crick sense and ligate are represented by identical labels. Each tile can have from one to four labels. Non-labeled corners indicate the absence of sticky ends. Experimental work [WYS98] has shown that, in principle, there exist parameter conditions (temperature, pH, ionic strength, etc.) under which tile binding to a slot defined by two sticky ends can be dominated by tiles with two matching sticky ends rather than single match tiles. Although these conditions are somewhat difficult to achieve in practice, this result shows that solution parameters can be tuned such that undesired and/or corrupted assemblies are avoided. It is taken for granted here that a tile would not occupy a slot unless both labels match.

Figure 1. Abstracted Tile Representation.

The actual implementation details are not discussed here since they fall outside of the scope of this manuscript. However, we believe that we make no arbitrary hypotheses. In fact, our work is based on the assumptions and achievements that come with DNA tiling computation in general.

2.3. The Non-Deterministic Algorithm. As was mentioned above, our design attempts to simulate a non-deterministic algorithm for the SAT problem. Non-determinism implies that at some step(s) the algorithm makes a non-deterministic choice (as if some oracle could tell the algorithm what the right choice is). In our case, this is simulated by an exponential number of DNA assemblies, expecting to cover all possible choices. The algorithm is given below. Notice that step 3 is the nondeterministic step.
NON-DETERMINISTIC SATISFIABILITY\((x_1, x_2, \ldots, x_m, C_1, C_2, \ldots, C_n)\)

1. Set all clauses to be unmarked
2. for \((i = 1, \ldots, m)\) do
3. Assign a value (0 or 1) to variable \(x_i\)
4. Mark all clauses \(C_j, j = 1, \ldots, n\) which are satisfied by this assignment
5. if (all clauses are marked) then return SUCCESS and output \(\{x_i\}\)
6. else return FAILURE

We present two slightly different designs for encoding this algorithm as a DNA tiling. The difference lies in what the tiles code. In design A tiles encode clauses; in design B tiles encode literals. After a detailed exposition of both designs a comparison is made.

3. Design A: Encoding Clauses

In this case we assume that the formula is given in 3-CNF form\(^1\). Since the number of possible clauses is countable in this case (see section 1), we can order all clauses and number them in some systematic way. With this mapping each clause \(C\) is represented by its corresponding number, say \(j\). For simplicity, in what follows we number a clause \(C\) with its number in the ordering as \(C_j\). Given a variable \(x_i\) and a clause \(C_j\), we can easily construct a function \(F(i, v, j)\) that determines whether the clause \(C_j\) is satisfied when variable \(x_i\) takes the binary value \(v\). The function \(F\) will be actually “precoded” in the structure of the DNA tiles.

We can represent the desired computation in a table format, that facilitates the transition to the explanation of the DNA assembly. Consider the 3-CNF formula

\[(\overline{x}_1 + \overline{x}_2 + x_3)(x_1 + \overline{x}_2 + \overline{x}_3)(\overline{x}_1 + x_2 + x_3)\]

and the table given below. There are 3 variables (represented in the first column of the table) and 3 clauses (represented in the last row) with numbers 1, 4, and 3 according to some ordering. The second column represents a possible assignment. Cells marked with “**” are helper cells.

<table>
<thead>
<tr>
<th>*</th>
<th>T</th>
<th>T</th>
<th>T</th>
<th>T</th>
<th>SS</th>
</tr>
</thead>
<tbody>
<tr>
<td>x₃</td>
<td>1</td>
<td>OK</td>
<td>OK</td>
<td>OK</td>
<td>*</td>
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<tr>
<td>x₂</td>
<td>0</td>
<td>OK</td>
<td>OK</td>
<td>C₃</td>
<td>*</td>
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<tr>
<td>x₁</td>
<td>1</td>
<td>C₁</td>
<td>OK</td>
<td>C₃</td>
<td>*</td>
</tr>
<tr>
<td>*</td>
<td>*</td>
<td>C₁</td>
<td>C₄</td>
<td>C₃</td>
<td>*</td>
</tr>
</tbody>
</table>

Following the algorithm above, the table is filled in a bottom-up manner, one row at a time. A cell corresponding to variable \(x_i\) and clause \(C_j\) will be marked as “OK” if and only if the clause \(C_j\) is satisfied when variable \(x_i\) takes the binary value \(v_i\) indicated in the second column or the cell below is already labeled “OK”. Otherwise, it is marked with the clause name “\(C_j\)”. Therefore, as we move up, “\(C_j\)” is propagated up as long as the clause is not satisfied. Once it is satisfied, it turns to an “OK” label, which propagates to the top independently of the assignments of the remaining variables. In effect, the entries of the table reflect the function \(F\).

It remains to check whether all clauses are satisfied. This is done in the first row of the table. Initially, we assume that the conjunction of all clauses is satisfied (label “T”=TRUE in the second column)\(^2\). The label “T” will propagate to the right as long as there are “OK”s to the right in the row below. The upper right cell is filled with “SS” (=SUCCESS) if and only if the cell to the left is “T” and the cell below is “**” (end of formula). Therefore, the formula is satisfied with this assignment if and only if the symbol “SS” appears in the table. Notice that if it was not satisfied, “T” would not had propagated to the end and “SS” would never appear, in which case we could propagate a label “F” (=FALSE) or leave the table incomplete.

\(^1\)Generalization to \(k\)CNF-SAT can be easily done, albeit increasing the number of required DNA tiles.

\(^2\)By definition, an empty conjunction is satisfied [DSW94].
The idea illustrated with the table above can be carried out almost directly by assembly of DNA tiles. The input is coded as a concatenation of tiles representing the first column and the last row of the table. This input structure is reproduced in billions and is mixed with a DNA solution that already contains tiles from a fixed library to be described shortly. The appropriate tiles will self-assemble on this input layer. Values are assigned to the variables in a random manner. Each assembly is testing one possible assignment to the variables. The input DNA structure and the resulting tiling computation of a satisfying assignment (in fact, the one in the table above) is shown in Figure 2.

![Figure 2. Input tiles and a successful assembly computation for design A.](image)

For illustration purposes, the whole computation is unfolded step-by-step in Figure 3. Initially, there is only one slot where a tile can bind. This tile will be an assignment tile; the label “V” (=Value) indicates that a value is expected for the particular variable. Once a value is assigned to $x_1$, there are two open slots. The slot to the left “asks” for an assignment to variable $x_2$. The slot to the right “wants” to check whether the assignment of $1$ to $x_1$ (shown as $x_1+$) satisfies clause $C_1$. This slot will be filled by the appropriate tile that contains the “answer”. Unfortunately, $x_1+$ does not satisfy $C_1$ and thus the label $C_1$ is propagated up, to be checked against the remaining assignments. At the same time, $x_1+$ is propagated at the other side to check for the remaining clauses. At the third step, there are three slots open. One for assigning a value to $x_3$, one for checking $C_1$ against $x_2-$, and one for checking $C_4$ against $x_1+$. Notice that both clauses are satisfied in this case, and thus “OK” is propagated up and left. This continues until assignment of values has been completed at which time the final check begins as well (indicated by label “C” for Check). If all clauses are satisfied the assembly will continue until the “T” (=True) label meets the “E” (=End) label and the success marker will be placed on the top. If there is some unsatisfied clause, the “T” label cannot propagate and the assembly will remain incomplete and thus without the success marker. At the very end, a separation procedure that isolates the assemblies containing the success marker will provide one or more solutions to the input instance.

3.1. Complexity of Design A. The complexity of the design is considered in terms of computation time, computation space and number of distinct tiles required. It is obvious from the examples given that the computation time $T(A)$ is equal to the depth (diagonal) of the assembly. In fact, it is

$$T(A) = (m + 1) + (n + 2) - 1 = m + n + 2 = \Theta(m + n) = O(m^3)$$
for $m$ variables and $n$ clauses, $\Theta(m + n)$ is linear in the size of the formula. $O(m^3)$ is an upper bound polynomial to the number of variables. We have used the fact that $n = O(m^3)$ for 3CNF-SAT (see section 1). The space $S(A)$ taken for each assembly is the area of the assembly.

$$S(A) = (m + 2) \times (n + 3) = \Theta(m \times n) = O(m^4)$$

which is upper-bounded polynomially to the number of variables. Finally, the library of fixed tiles need contain the following tiles:

- **Variables.** There must be $m$ tiles coding $m$ variables, where $m$ is the maximum number of variables that can appear in a formula.
- **Clauses.** There must be \( n = \frac{4}{3} m (m-1)(m-2) = \Theta(m^3) \) tiles coding all possible clauses.

- **Input Boundaries.** There are 4 such tiles to mark the end of variable list, the end of clause list, the beginning of value assignment and the beginning of computation in the input assembly.

- **Assignment.** Each variable can take one of two values, so there is a total of \( 2m \) tiles for assigning values. A + sign indicates an assignment of 1, whereas a – sign indicates an assignment of 0.

- **Computation Boundaries.** For each variable assignment there must be an ending tile, thus a total of \( 2m \) such tiles.

- **Computation.** For each variable assignment and for each clause there must be a tile that indicates whether the clause is satisfied or not. There are \( 2mn \) such tiles. Further, there must be tiles to propagate the “OK”s of the satisfied clauses to the end of the assembly. There are \( 2m \) of those tiles.

- **Final Check.** Finally, there must be some tiles to check if all clauses are satisfied and mark the result. There are 3 such tiles.

Summing up all the numbers, we have the total number of tiles for design A.

\[
N(A) = 7m + (2m + 1)n + 7 = \Theta(m \times n) = \Theta(m^4)
\]

**3.2. An Example.** In order to reinforce the method, Figure 4 shows another example of successful computation for a formula with 8 variables and the following 10 clauses.

\[
\begin{align*}
C_1 &= (\overline{x}_2 + x_3 + x_7) \\
C_2 &= (x_3 + \overline{x}_5 + \overline{x}_8) \\
C_3 &= (\overline{x}_1 + x_2 + x_4) \\
C_4 &= (\overline{x}_3 + x_5 + x_6) \\
C_5 &= (x_1 + \overline{x}_4 + x_5) \\
C_6 &= (x_5 + x_7 + x_8) \\
C_7 &= (\overline{x}_2 + x_4 + x_6) \\
C_8 &= (x_3 + \overline{x}_4 + x_5) \\
C_9 &= (x_1 + x_4 + \overline{x}_6) \\
C_{10} &= (x_2 + \overline{x}_6 + x_8)
\end{align*}
\]
4. Design B: Encoding Literals

This design attempts to overcome the large number of tiles required by design A. In particular, it operates at a lower level, encoding literals that appear in each clause rather than the clause itself. The basic idea is the same. Consider the same 3-CNF formula

\((\overline{x}_1 + \overline{x}_2 + x_3)(x_1 + \overline{x}_2 + \overline{x}_3)(\overline{x}_1 + x_2 + x_3)\)

and the table given below. There are 3 variables (represented in the first column of the table) and 3 clauses (represented in the last row) with all literals listed explicitly. The “s” (=SEPARATOR) is used to separate clauses. The second column represents a possible assignment as before. Cells marked with “*” are helper cells.

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<tbody>
<tr>
<td>*</td>
<td>F</td>
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<td>T</td>
<td>T</td>
<td>F</td>
<td>T</td>
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<tr>
<td>x_3</td>
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<td>\overline{x}_1</td>
<td>OK</td>
<td>OK</td>
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<td>OK</td>
<td>OK</td>
<td>\overline{x}_3</td>
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<td>OK</td>
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<td>x_2</td>
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<td>\overline{x}_1</td>
<td>OK</td>
<td>x_3</td>
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<td>x_1</td>
<td>1</td>
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</tr>
</tbody>
</table>

Again, following the algorithm given in section 2.3, the table is filled in a bottom-up manner, one row at a time. However, now a cell corresponding to variable \(x_i\) and literal \(y_j\) will be marked with “OK” if and only if literal \(y_j\) is TRUE when variable \(x_i\) takes the binary value \(v_i\) (indicated in the second column) or the cell below is already labeled “OK”. Otherwise, it is marked with the literal name “\(y_j\)” to propagate the computation to the next row. At the end of the day, the second row of the table will contain “OK”’s and/or some “\(y_j\)”’s depending on what satisfies what.

The checking step is a little more involved compared to the previous one. We need to check whether each individual clause is satisfied and further whether the whole formula is satisfied. Since each clause is a disjunction, we initially assume that it is not satisfied\(^3\). This is denoted by the “F” (=FALSE) label in

\(^3\text{By definition, an empty disjunction is not satisfied [DSW94].}\)
the second column. As long as the particular clause has not been satisfied (i.e., there are \( y_j \)'s to the right and bottom of “F”), the label “F” propagates to the right unchanged. Once, an “OK” label is encountered to the right and bottom of an “F”, it turns to a “T” (=TRUE) label, indicating that the clause has been satisfied. “T” propagates until the separator symbol “s” is met to the bottom-right. If “T” meets the separator, that means that the current clause is satisfied and we can continue with the next one by initializing to “F” again. However, if “F” meets the separator, that means that this assignment failed to satisfy the formula and computation is halted. Finally, if the “F” label hits the “ss” marker at the end of the table, it is implied that the whole formula is satisfied and therefore the success symbol “SS” marks the upper-right corner of the table. As previously, an assignment is satisfying if and only if the symbol “SS” appears at the upper-right corner of the table.

Having the concept of the table in mind, it is easy to make the transition to the DNA assembly. The basic idea is again the same as previously, but the tiles and the coding are somewhat different. Figure 5 below shows the input tile assembly and the successful computation of the table above.

![Figure 5. Input tiles and a successful assembly computation for design B.](image)

Notice that by encoding literals we are not restricted to 3CNF-SAT (or to any \( k \)-CNF-SAT) anymore. We can encode any SAT formula given in CNF format in a straightforward manner.

4.1. Complexity of Design B. Again, the complexity of the algorithm is considered in terms of computation time, computation space and number of distinct tiles required. The computation time \( T(B) \)
is

\[
T(B) = (m + 1) + (l + n + 2) - 1 = m + l + n + 2
\]
\[
= \Theta(m + l + n) = O(m + mn + n) = O(mn)
\]
The last equality follows from the fact that in the worst case all clauses contain all the variables, that is \(l = mn\), and therefore \(mn\) is an upper bound for \(l\). If we consider 3CNF-SAT only, then \(l = 3n\) and therefore

\[
T(B) = (m + 1) + (4n + 2) = \Theta(m + n) = O(m^3)
\]

which is of the same order as in the previous design. The space \(S(B)\) taken for each assembly is

\[
S(B) = (m + 2) \times (l + n + 3) = \Theta(m(l + n)) = O(m^2n)
\]

For 3CNF-SAT, \(l = 3n\) and so \(S(B) = \Theta(mn) = O(m^4)\) as before. The tile library is somewhat different in this case.

- **Variables.** There are \(m\) tiles coding the \(m\) variables, where \(m\) is the maximum number of variables that can appear in a formula.

- **Literals.** There are \(2m\) tiles coding all possible literals. They are used for coding the clauses of the formula.

- **Input Boundaries.** There are 5 such tiles: four are the same as in design A, plus the separator.

- **Assignment.** The same as before. There is a total of \(2m\) such tiles.

- **Computation Boundaries.** As before, for each variable assignment there must be an ending tile, thus a total of \(2m\) such tiles.

- **Computation.** For each pair of literals \(x_i, y_j\) there must be a single tile. If the literals do not refer to the same variable, or they refer to the same variable but are complementary, the result is simply propagation of the labels along the diagonals of the tile. However, if they match, \(x_i\) is propagated and the literal \(y_j\) is turned to an “OK”. This requires

\[
\binom{2m}{2} = m(2m - 1)
\]

such tiles. Further, for each variable and each possible assignment there must be tiles to propagate the “OK” labels up. There are \(2m\) of those tiles.

- **Final Check.** Finally, the tiles that check for satisfiability at the end are a little more involved than in the previous case. First, “F” has to propagate over literals \(y_j\) and turn into an “T” if “OK” is encountered. “T” propagates over \(y_j\)’s and “OK”’s. Finally, two tiles are needed for the extrema and one for resetting “T” to “F”. In total, \(4m + 5\) tiles.
Summing up all the numbers, we have a total number of tiles:
\[ N(B) = 2m^2 + 12m + 10 = \Theta(m^2) \]

which is two orders of magnitude less than the total number of tiles in design A.

5. Comparison and Discussion

The two designs implement the same algorithm in a slightly different way. It is our belief that design B is better compared to design A for two main reasons: (1) the input formula can be any CNF formula, and (2) the total number of required tile types is only \( \Theta(m^2) \), where \( m \) is the maximum number of variables. In contrast, design A assumes that the formula is given in 3CNF and requires \( \Theta(m^4) \) tile types. On the other hand, design A results in smaller computation time and space but asymptotically it seems that the difference disappears (see the analysis above).

There is a final detail that is crucial for the success of both algorithms. The concentrations of assignment tiles corresponding to variable \( x_i \), i.e. tiles for \( x_i+ \) and \( x_i- \), have to be equal so that there is equal chance of assigning either value. If this is not the case, there might be assignments that will never be explored because of this “discrimination” in assigning values.

A limitation of the algorithm, which is common for most DNA computations, comes from the fact that the exponential dimension of the problem has been pushed into the physical space (volume) occupied by the DNA molecules. This will eventually become a restrictive factor. The input size and thus the DNA volume cannot grow forever. This implies an upper bound to the size of instances that can be solved in practice. Obviously, the practicality of a DNA algorithm for satisfiability is heavily dependent on whether this upper bound is well above the upper bound for instances that can be solved on a conventional computer.

6. Future Work

It is an open question whether the design(s) will work well in practice or not, a fact that can be verified only experimentally. Encouragement comes from the recent investigations of several DNA tile structures [WLWS98]. In particular, TAO35 (see figure 1) is a general DNA tile that is currently being investigated for use in self-assembly computations [LWR99]. Moreover, it was recently demonstrated by LaBean, Winfree and Reif that input layers like the ones we use can be constructed relatively easily using a long “scaffold strand” of DNA which traverses all input tiles [LWR99]. The scaffold strand acts as a nucleation front for assembly of input layer tiles. The completed input layer then acts as the foundation for growth of the overall tile assembly, as described above.

Another line of research will focus on ways to enhance the algorithm will well-known heuristics for satisfiability, such as the unit propagation rule (if there is a clause with a single literal, force the corresponding variable to take the value that makes the clause true) and the purification rule (if a variable appears in the formula in exclusively negated or non-negated form, assign to this variable the value that makes all instantiations true). Actually, both of these rules can be taken into account in the current designs simply by altering the concentrations of the assignment tiles to the corresponding variables so that only the desired value is given as an option. However, this way it becomes a manual step that is perform only at the beginning. Alternatively, preprocessing of the formula could eliminate such variables. Our goal is to incorporate them in the algorithm since the need for unit propagation and/or purification might reappear during computation.

7. Conclusion

We presented an algorithmic design for solving the satisfiability problem using 2-dimensional DNA self-assembly. It seems that the peculiarity of DNA-based computation requires a reconsideration of our thinking about algorithms and computation in general. New designs or redesigns of traditional algorithms
aiming to match the requirements and capabilities of DNA computation models seem to be an important step toward useful DNA-based computers.

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