Computation of decision problems within messages in DNA-tile-based molecular nanonetworks

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Abstract

Akyildiz et al. envisioned the use of nanonetworks as a new paradigm for computation and communication on a very small scale. We present a new approach to implement nanonetworks with molecular communication using tile-based self-assembly systems on the basis of DNA. In this model, the medium of communication is filled with DNA-based molecules. Furthermore, some nanobots are capable of creating or releasing said molecules. Once present, they can be detected by other nanobots and interpreted as messages. Some DNA-based molecule systems are capable of universal computation. We show that it is possible to construct systems, in which the evaluation to TRUE of an arbitrary decision problem is a precondition for the assembly of a message molecule. We relocate computations from nanobots into message molecules, thereby revolutionizing the paradigm for computation in nanonetworks. This approach can be interpreted as computation inside the communication channel. We further present message molecules that only assemble if a marker has been detected at least k times, as a proof of concept.

1. DNA-based nanonetworks

DNA has been proposed as a material for construction as early as 1982 by Seeman [1]. Among others, this early work inspired and accelerated research that targets computational processes at the nanoscale. In recent years, nanonetworks have been introduced as a methodology to compensate for the resource constraints to be expected at increasingly small scales [2]. In nanonetworks, nanoscale devices collaborate to solve tasks that exceed the capabilities of single devices. To achieve this, the devices have to communicate with each other. For example, terahertz communication and molecular communication have both been examined as supportive technologies.

Over the past decade, numerous scenarios have been proposed in which the application of nanonetworks might be beneficial. Material sciences, medical, pharmaceuticals as well as computer science and chemistry are among the most prominent contributors. Medical applications appear to be most established. Since molecular communication (e.g., hormones) is native to living organisms, it is of special interest in medical scenarios.

In [3], a huge variety of applications for molecular communications have been surveyed. The detection and treatment of diseases are of general interest. Both can be further itemized into categories like drug delivery, immune system support, nanosurgery or the constant monitoring of health parameters.

DNA-origami is an established, replicable technique which generates nanoscale objects (e.g., message molecules) with the desired physical properties [7]. The process utilizes a long strand of DNA that is pinned together at predetermined base pairings by a short DNA-strand. While it is possible to create complex structures using DNA-origami, the procedure is clearly limited in its capabilities. The longer the strands get, the higher the error-rate. Nevertheless, it has been shown that DNA-origami can create nanostructures that are capable of detecting certain markers and releasing a payload on detection [8].

We further investigate the tile-based self-assembly method that is also based on complex DNA-molecules. For this, we chose molecules that greatly differ from the conventional DNA double-helix that represents the genetic material of all organisms. An example is the DX-molecule that combines two parts of a double helix into a single molecule with open strands to both sides [9].
Those open strands can encode conditions, which have to be met for other DNA-molecules to form a stable binding.

If correctly arranged, DNA can be shaped into complex objects with multiple open ends in many directions. The molecules, often called tiles (see Fig. 1), can then be introduced into a common medium where they start to self-assemble into even more sophisticated objects. The process is mostly used for computational purposes or to create structures at the nanoscale [10].

In literature, it is often suggested that nanobots communicate through specific molecules [11]. We instantiate these message molecules through tile-assembly. A message molecule can either be a single tile or a more complex assembly of tiles. The process of tile-based self-assembly does not require nanobots to produce a complex molecule—a seed-tile is enough to initiate a self-assembly process to generate complex message molecules. Nanobots thus do not need to produce big molecules, simplifying nanobot design.

A message molecule may require a specific tile in order to finalize self-assembly. With all other tiles already present in the communication medium, a nanobot needs to release only a single tile as trigger to start or continue self-assembly. Tiles and assemblies of tiles (in this case message molecules) are subject to Brownian motion like any other molecule and may utilize it as a way of propagation. Every message molecule might be detected by other bots, which may amplify the message by producing/releasing additional tiles until a concentration threshold is met. Another option is the immediate release of therapeutics or fluorescence markers that indicate a positive diagnosis.

In the later presented scenario, the bots communicate among each other via self-assembly binding reactions of message molecules that might harbor binary information, as shown in Section 3.2. Communication to the macroscale may be realized by reading molecule concentrations, as with regular DNA- computing [12], or by reading information encoded on the message molecule with fluorescence markers.

Nanobots initiate communication with the production of only one, small type of tile. The required productions cost necessary might be greatly reduced. Other techniques require the production or release of complex and bigger molecules.

Another major benefit of tile-based message molecules is that computations are shifted from nanobots themselves into the assembly process of message molecules. This modifies the computation paradigm in nanonetworks, where it is generally assumed that nanobots execute computations. The space restrictions in the communication channel are far less severe. To the best of our knowledge, to this day, no major research explores the possibility of computations taking place in the communication channel.

Any decision problem [13] can be computed by a tile-assembly system as shown by [6]. However, it remains unclear if nanobots may perform computations like classical computers, or fulfill the space requirements imposed by complex computations [14]. Thus, it appears beneficial to utilize a well tested process to encode conditional behavior at the nanoscale, thereby answering the question of how computations may look like in nanonetworks.

The remainder of this work is structured as follows: Section 2 explains the necessary basics for tile-based self-assembly systems. Section 3 elaborates the use of tile-based self-assembly systems for the purpose of molecular communication and explains the relevant basic components. Section 4 compares the presented framework to other channel models and evaluates the approach. Section 5 concludes the work, and Section 5.1 presents ideas on future work and some open questions.

2. Preliminaries for tile-based self-assembly

This section provides definitions and examples of the mathematical constructs known as tiles and assemblies, as well as all relevant properties [6]. The definitions are based on [15]. For an in-depth introduction to tile-based self-assembly see [12]. Assemblies, made up out of tiles, serve as messenger molecules in this communication scheme, and can additionally compute arbitrary problems through self-assembly.

Definition 1. An n-dimensional tile \( t_n \) is an object in the \( \mathbb{Z}^n \) with unit length and angles of 90°. DNA-tiles cannot be rotated.

A side of a tile \( t_n \) is defined by a vector \( u_i \in \mathbb{Z}^n \). \( u_i \) has exactly one non-zero element with value 1. Sides are enumerated by:

\[
\text{side} : t_n \mapsto U_1 \times U_2 \cdots \times U_n
\]

\( U_i \) is the set of unique unit vectors \( u_i \) defining the sides of \( t_n \). The vector \(-u_i\) denotes the side opposite of \( u_i \) in the same dimension.

In the following, we often omit the dimension of tiles when this information is irrelevant in the presented context. We focus on two- and three-dimensional tiles. A 2D-tile \( t_2 \) is a square in the \( \mathbb{Z}^2 \) and a 3D-tile \( t_3 \) is a cube in the \( \mathbb{Z}^3 \). Fig. 2 shows an example of a 2D-tile.

Each side of a tile can have one or multiple glues:

Definition 2. A glue \( g \in G \), where \( G \) is the set of utilized glues, is defined by a label \( \Sigma \in \Sigma^* \), where \( \Sigma \) is an alphabet and \( s \in \mathbb{N} \) a glue-strength, as well as the functions:

\[
\text{label} : G \mapsto \Sigma^* \\
\text{strength} : G \mapsto \mathbb{N}
\]

The glue label is often referred to as color. Glues are instantiated by open DNA strands that can encode both conditional bindings and varying strengths.

In the DNA-computing community, tiles are implemented with DNA [12]. An example structure can be seen in Fig. 1. The open strands can be used to create a variety of different glues.

Each tile can occupy exactly one position in the discrete space \( \mathbb{Z}^n \), given by the vector \( v_i \in \mathbb{Z}^n \). Such a position is called a site.

Definition 3. Two tiles \( t \) and \( t' \) that occupy sites defined by the vectors \( v_i \) and \( v_i' \in \mathbb{Z}^n \) are neighbors iff \( |v_i - v_i'| = 1 \) and the resulting vector \( e = v_i - v_i' \) has exactly one non-zero element.

A tile can have zero or exactly one neighbor on any of its sides. There is no interaction between the tiles \( t \) and \( t' \) unless they are neighbors. The rules for this interaction are specified by their glues.

Definition 4. An n-dimensional tiletype \( T_n \) is a template for a tile \( t_n \). It is defined by a label \( \Sigma \in \Sigma^* \) and a set of glues \( g_i \in G \), one on each side \( u_i \in U_i \) of \( t_n \), and the functions:

\[
\text{glue} : T_n \times U_i \mapsto G \\
\text{strength} : G \mapsto \mathbb{N}
\]

All tiles that have identical glues and the same markers are of the same tiletype.

All not explicitly mentioned or shown glues have the label \( \Sigma = \text{void} \) and strength \( s = 0 \).

A tile \( t \) that has a glue of strength zero or no glues on side \( u_i \) cannot interact with its neighbor \( t' \) at the side \( u_i \).
Two tiles can **bind** together if they are neighbors and have glues of matching color ("matching" with DNA-tiles means that both tiles have complementary open strands) as well as a strength of at least the temperature $\tau$.

The temperature is modeled after the physical temperature. As the physical temperature increases, molecules move faster, thus increasing the chance of molecular bonds to break. Molecules with stable and stronger bonds are more stable at higher temperatures than those with fewer or weaker bonds.

**Definition 5.** The **temperature** $\tau$ of a tile system describes the minimum glue strength required to form a stable binding.

Due to the nature of DNA, some bindings will always be incorrect [5]. However, the number of incorrect bindings can be reduced by designing special sets of tiletypes that account for error behavior. [12] shows two approaches to mitigate errors for two-dimensional assemblies.

The **binding strength** $s$ between the tiles $t$ and $t'$ is equal to the sum of the matching glues on the involved sides. The strength of a glue is denoted by the number of black bars on a side of the tile. In Fig. 2, all glues are of strength 1.

**Definition 6.** Two tiles $t, t'$ of type $T$ and $T'$ **correctly bind** at temperature $\tau$ if:

1. $t$ and $t'$ are neighbors
2. $\exists u_i \in U : \text{label}(\text{glue}(t, u_i)) = \text{label}(\text{glue}(t', -u_i))$
   \[ \land \text{strength}(\text{glue}(t, u_i)) \geq \tau \]
   \[ \land \text{strength}(\text{glue}(t', -u_i)) \geq \tau \]

Bindings that do not fulfill the conditions (1) and (2) are considered **incorrect** or **errors**.

**Definition 7.** An $n$-dimensional **tile assembly**, or just assembly, is a partial function $\alpha : \mathbb{Z}^n \mapsto T$, where $T$ is a set of $n$-dimensional tiletypes $T_n$. An assembly $\alpha$ is $\tau$-stable if no tile $t_\tau \in T$ can be removed without breaking bonds of at least strength $\tau$, where $\tau \in \mathbb{N}$.

**Definition 8.** The **border** of an assembly $\alpha$ is a subset of tiles of $\alpha$. It includes all tiles that have an unoccupied site, that is, where the tile has no neighbor.

Only border tiles can interact with tiles added to the assembly at a future time.

**Definition 9.** The **growth front** of a $n$-dimensional assembly is a subset of all sites in the $\mathbb{Z}^n$. A site belongs to the growth front iff it is not occupied by a tile and a neighboring site is occupied by a border tile with a minimum glue-strength of $\tau$ towards the empty site.

The sites of the growth front may change when a tile is added to or removed from the assembly. For the sake of simplicity, we assume that a single random tile is added to the assembly $\alpha$ at every discrete time step.

The initial assembly $\alpha_0$ at time step 0 is the seed-tile or seed assembly $\sigma$. Starting from $\sigma$, one tile is non-deterministically added to $\alpha$ at a growth front site at each time step. Due to the non-deterministic nature, it can be complicated to design tile sets that assemble into a unique terminal assembly.

**Definition 10.** A **Tile Assembly Model (TAM)** is a three-tuple $T_{\tau} = (T, \sigma, \tau)$, where $T$ is a finite set of tiletypes, $\sigma$ a seed assembly and $\tau \in \mathbb{N}$ the temperature of the TAM.

In the remainder of the text $\tau$ is assumed to be 2 and the stability of a TAM is omitted.

**Definition 11.** $A[T]$ denotes the set of all terminal assemblies that can be produced by a TAM $T$ in a finite number of time steps. An assembly $\alpha \in A[T]$ is **terminal** if no tile can be $\tau$-stably added to it.
Fig. 3. (a) An example tile set of a 2D-TAM as a visual representation. (b) The tile assembly sequence of a TAM with a tile set as depicted in (a). The seed tile is S and temperature $\alpha = 2$. $\alpha_0, \ldots, \alpha_3$ depict the three time steps the TAM requires to reach a terminal assembly.

**Definition 12.** Let $\alpha_i$ be an assembly produced by a TAM $\mathcal{T}$. An assembly sequence of TAM $\mathcal{T}$ is the sequence $A = (\alpha_0, \alpha_1, \ldots)$, where $\alpha_{i+1}$ is obtained from $\alpha_i$ by adding one tile to $\alpha_i$. If the sequence $A$ is finite, the last element of the sequence $A$ is the result of $\mathcal{T}$.

Fig. 3(a) shows an example tile set of a 2D-TAM, where the seed is the tile labeled S. The assembly sequence of this TAM with temperature $\tau = 2$ is depicted in Fig. 3(b). $\alpha_0$ is the initial assembly which consists of only the seed tile. $\alpha_2$ and $\alpha_3$ show the addition of one tile each to the previous assembly.

A tile-assembly system (TAS) is an extension of a TAM. It governs the rules at which and at what pace tiles are added to or removed from a growing assembly. There are several prominent tile-assembly systems. The most popular ones date back to Eric Winfree [6] and are the abstract Tile-Assembly Model (aTAM) and the kinetic Tile-Assembly Model (kTAM) [12]. All tileset verifications in this work use the realistic kTAM model. It reflects the nature of DNA by introducing a probability that tiles may break free from the assembly.

**2.1. Errors in self-assembly systems**

There are three types of errors commonly observed in tile-based self-assembly systems like the kTAM [12].

**Definition 13.** A growth-error occurs when a tile is attached to the assembly in such a way that at least one of its glues mismatches an adjacent glue.

**Definition 14.** A facet-error occurs when a tile is attached to the assembly without any mismatched glues but with insufficient binding strength to satisfy the temperature requirement.

Both error types can become locked in, when a new tile is attached at a neighboring site, providing correct bonds to the erroneous tile. Therefore, the erroneous tile’s total binding strength may satisfy the temperature requirement and thus be unlikely to detach from the assembly. Fig. 4 depicts the process. Growth-errors and facet-errors can be significantly reduced by employing block-replacement strategies like snaked-proofreading or $k \times k$-proofreading [16,17].

The third kind of error differs from the other two. It occurs, when the assembly process starts with tiles other than a seed tile.

**Definition 15.** A nucleation-error occurs, when a tile-assembly system starts an assembly process with no seed tile present or with any other tile than the seed.

**3. Tile-based communication systems**

This section introduces the most important components of tile-based molecular communication systems, namely senders, channels and receivers as well as messages and a mechanism for debugging. Furthermore, it details a construction scheme for a message molecule that computes a logical 4-bit AND.
every glue in a TAM to globally suppress assembly. Blue tiles are conditional-tiles that actually calculate binary decision problems. Note, that the marker in the middle of the conditional tile does not represent its binary value—the presence of a tile represents a logical TRUE. The white tile is the seed-tile that starts the assembly and the red tiles enable growth in the y-dimension.

Fig. 5(b) shows the resulting message molecule. This assembly fulfills a multitude of required properties. It consists of a seed-tile to the right labeled ‘S’. The assembly advances by adding a border tile to the top and the bottom of the seed-tile. The tiles at the border of the assembly have as few glues to the outside as possible, to prevent erroneous bindings. The four tiles to the left of the seed-assembly are conditional-tiles. In this case, they encode the computation of a 4-bit logical AND.

The message molecule requires the presence of all tiles — especially the four different conditional-tiles — to fully assemble. Releasing conditional-tiles only under special conditions (e.g. in the presence of a disease-related biomarker) allows for the encoding of binary information into conditional tiles. By adding additional distinct conditional tiles, one can easily compute a k-bit AND. This approach can be utilized to find a distributed consensus concerning certain events among nanodevices. By employing this strategy, the number of false positives can be greatly reduced.

3.3. Assembly errors in message molecules

Message molecules are prone to the three types of errors presented in Section 2.1. Facet-errors and growth-errors can both be reduced significantly by employing block-replacement schemes like snaked-proofreading or \( k \times k \)-proofreading [16,17]. Fig. 6 shows an example. \( k \times k \)-proofreading simply increases the scale of the assembly, whereby rendering growth-error \( 1/k \) times as likely. In addition, snaked-proofreading also prevents facet-errors in one direction completely, by omitting the internal glue between the two lower tiles of the snaked-block. By replacing each tiletype by a block of tiletypes, more errors are required for a whole original tile to be erroneous. The in-message calculation remains equivalent.

Nucleation-errors are a different matter. In theory, the leftmost two columns of the message molecule in Fig. 5 can form without the presence of a seed-tile, thereby forming a ligand without a successful computation. Two facet errors combined with a nucleation-error are sufficient to produce this behavior. The probability for a single facet-error — namely two tiles attaching next to each other with insufficient strength at once, thereby locking each other in place — is very low. Two of them happening at the same nucleation-error is even more unlikely. Due to the high concentration of tiles, the described scenario will eventually occur, however unlikely.

This can be circumvented by the presented snaked block-replacement strategy. However, it is necessary to adapt the outer glue structure to ensure that only two outer glues are present.

Another possibility would be a redesign of both receptor and message molecule to also enforce the presence of the seed-tile. Fig. 7 shows a possible adjustment. An appropriate receptor can easily be designed. This way, the presence of both the seed-tile and the ligand are enforced and therefore the effects of nucleation-errors are mitigated.

3.4. Message receivers

Fig. 8 shows two examples of possible receptors for message molecules. Both receptors and ligands are based on DNA. The binding of a ligand at a receptor is a DNA binding reaction that is subject to the temperature \( \tau \).

The gray squares represent the arbitrary ligand/receptor-part of the message molecules. The black squares represent individual glues of strength 1. In order for a message molecule to form a stable binding with a nanobot, at least a binding strength equal to the system temperature \( \tau \) is necessary.

If designed accordingly, nanobots can detect the last three tiles that were added to a message only at once. This is necessary in order to prohibit the binding of single tiles with the receptors and thus reducing the probability of binding errors at receptors.

Once a message molecule binds at a receptor, this might trigger any kind of reaction. This could, e.g., be an own release of seed-tiles and thereby help spread messages through the medium. A single nanobot is not necessarily limited to one type of receptor.

If the goal is just communication, single tiles are enough to transmit binary information. Assemblies of multiple tiles are only required if the message should perform additional computation.
3.5. Example for in-message computation

We envision the application of DNA-tile based nanonetworks in the blood stream of living organisms. Since there are numerous unknown environmental parameters in the human body – in addition to the technical challenges – we illustrate the usefulness of the presented approach in a more constrained, but easily achievable, realistic scenario.

To quickly and reliably detect a disease, a DNA-tile based nanonetwork can be used in a controlled environment like a petri dish. The components for the nanonetwork and a blood or tissue sample can be mixed in a petri dish, where it is easier to control the environmental parameters that are required for self-assembly systems.

If a disease has been detected, conditional tiles are released and distributed via Brownian motion. In a petri dish, the tile concentration can be chosen as high as necessary to reliably assemble message molecules. Additionally, we extrapolate from wet lab experiments that successfully assembled a Sierpinski triangle with a tile-based approach.

Once the molecules are fully assembled, they behave like any other molecule that acts as message carrier in a molecular communication system and can transmit the information to a receiving nanobot, that in turn releases a chemical that starts a fluorescence reaction. This scenario illustrates how nanonetworks could be used to help diagnose diseases with a high level of confidence.

We design a tile-based nanonetwork consisting of a number of nanobots and sensors that dispatch specific tiles once a marker is detected. To ensure that few “false positive” messages occur, our tile-assembly based nanonetwork guarantees that at least 4 individual nanobots detected a marker for a whole message molecule, with ligands, to assemble. We utilize the set of tiletypes from Fig. 5 for this scenario.

We include the computation of a 4-bit logical AND into our message molecules, thus proposing a method to immediately release therapeutics or fluorescence markers once a disease is reliably detected. The 4-bit AND ensures a distributed consensus about an event with high probability.

Generally speaking, tile-assembly systems are Turing complete at temperature 2 or higher [6,13], thus similar systems can compute arbitrary decision problems and then add appropriate ligands that can be detected by other nanobots to identify a positive computational result.

Fig. 9 illustrates the vision: 1. Nanosensors detect the disease-related biomarkers (orange squares) and 2. release a number of conditional tiles 1–4 in return. 3. All other tiles constantly circle the channel (blood stream/petri dish) and start to further assemble once the conditional tiles are present. When all four tiles are gathered, the ligand-tiles can stably attach to the message molecule. 4. With the ligand function now present, other nanobots can bind the message molecule at their corresponding receptors, thus communicating the information that a disease has been reliably detected. 5. Nanobots release an antigen to help combat the detected disease.

The nanobots only detect a message molecule once it is fully assembled, and a message molecule only fully assembles once all required tiles are present. The set of tiletypes is designed in a way that it always assembles into exactly one terminal assembly while erroneous bindings are rare. All presented sets of tiletypes were tested and verified in a kTAM simulator [19,20].

3.6. Scenario modularization

The presented medical scenario serves to clarify the vision of nanonetworks. In reality, the experiments will start with much simpler scenarios in vitro. To test such a scenario in wet lab experiments, past experiences have shown that it can be beneficial to decompose a vision into smaller parts. Those components are then tested on their own, thus validating parts of the approach at lower cost.

Such a wet lab experiment could be realized by using a piece of human tissue in a petri dish and then testing the nanonetwork components separately. As a last step, all functioning parts have to be re-assembled into the scenario. A non-medical testing approach for the whole scenario, after parts have been tested, could be to use nanonetworks to monitor and manage agriculture or fish tanks.

The postulated scenario is rather generic. The phases are depicted in Fig. 9. The scenario can be generalized to the detection of an arbitrary marker. In essence, the scenario consists of the following parts:
1. Detection of a generic marker by a nanosensor. Li et al. proved in a wet lab experiment that therapeutics can be released by detecting a specific molecule \[8,21\].
2. Storing therapeutics, tiles or other payload in/on a nanobot. A variation of this problem has even been solved in vivo by Li et al. \[8,22\].
3. Assembling the 4-bit and message molecule under laboratory conditions. In general, this should be possible, since many experiments in the DNA-computing community yield a lot more complex structures \[23\].
4. Attaching a fully assembled message molecule to a pre-assembled receptor. In theory, this should also be possible. The DNA-computing community created the so-called 2HAM model that simulates the interaction between partial assemblies. The process seems to work under laboratory conditions \[24\]. A receptor may be assembled using a similar approach as in the message molecule. A receptor can therefore be an assembly.
5. Release of therapeutics after a message molecule has been detected at a nanodevice. Again, Li et al. solved variations of this problem \[8\].

These partial problems have mostly been solved using a DNA-based approach. To the best of our knowledge, reliable molecule release mechanisms remain an open problem in tile-based self-assembly systems. It would be necessary to locally decrease the stability of a nanobot or molecule to open it and release a payload. In theory, DNA-tiles are compatible with other DNA-based approaches and it should thus be possible to adapt already functioning technologies, like the mechanical opening mechanism presented by Li et al. \[8\].

4. Evaluation

Since no wet lab experiments have yet analyzed communication in tile-based nanonetworks, we mainly investigate theoretical channel models for molecular communication with a coherent biological basis. We evaluate our proposed scenario by both simulation and comparison with other molecular communication approaches—which is only partially possible. Further, we extrapolate from previously conducted wet lab experiments to further strengthen our reasoning.

4.1. Simulation and validation

We simulated the presented scenario using the software ISUTAS by Patitz \[20\]. We instantiated the tilesets for both the kTAM and the 2HAM model. The kTAM realistically simulates the assembly of exactly one message molecule. The model accurately predicts the assembly of tiles as validated in \[25\]. We used the 2HAM to validate the interaction between receptors and message molecules and analyzed the interactions between intermediate assembly products. Once the message molecules are fully assembled, we expect them to behave like regular molecules used for communication. We refer to established molecular communication models to further strengthen our line of argumentation.

Fig. 10 shows the tileset for both the kTAM and the 2HAM simulation. The blue tiles model the receptor. The remaining tiles assemble into a message molecule.

**Fig. 10.** The tileset for both the receptor (blue tile types) and the message molecule. The text in the center of a tile represents its label. The number of lines on the edge of a tile type represents the strength of a glue. The color of a glue is denoted by a label next to it. A dotted line means that a glue has strength 0. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

K TAM. We simulated the assembly of the message molecule with parameters \( G_{se} = 10.4 \) and \( G_{mc} = 17.0 \)—the same values as in \[25\]. \( G_{se} \) describes the free energy cost of breaking a single bond and \( G_{mc} \) the monomer concentration. The ratio of the binding cost to the strength of a single bond \( \tau = G_{mc} / G_{se} \) represents the temperature parameter of the kTAM.

A value of \( G_{se} = 10.4 \) led to more errors in the simulation result in \[25\]. We therefore simulated under problematic conditions. A \( G_{mc} = 17 \) value corresponds to 0.8 µM per tile and represents the monomer concentration. A \( G_{se} = 10.4 \) value corresponds to a temperature of 32.7 °C. Lower values of \( G_{se} \) represent a higher temperature, e.g., \( G_{se} = 8.5 \) corresponds to 41.8 °C.

The simulator assumes that every tile type occurs equally often in the medium and sufficient tiles for additional growth are always available—this includes tiles for the receptor, which slows down assembly. In reality, the individual tile concentrations can be manipulated to further enable control over the growth process. The simulation showed that a message molecule fully assembled on average after approximately 3000 attempted binding reactions. Ideally, only 28 binding reactions are necessary for the assembly. Fig. 11 shows the result of the simulation.

**Fig. 12.** A histogram of the results of 100 assembly simulations. The number of binding reactions (both forming and breaking bonds) necessary to fully assemble a message appears to be Log-normal distributed, with a slight bias towards higher numbers (\( \sigma \approx 0.5 \)). Only one fully assembled message occurred in under 1500 reactions. In reality, numerous binding reactions happen in parallel and the required number of binding reactions is an indicator of the time required to fully assemble a message. As long as \( 2G_{se} > G_{mc} > G_{se} \) holds, tile associations are more likely than tile dissociations and a fully assembled message molecule will eventually form \[26\].

No simulation run showed an erroneous interaction between the message molecule and the tiles that model the receptor. Even after more than 100,000 binding reactions, the fully assembled message showed no errors and remained stable.

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1.5. Conclusion

TAM and the 2HAM models were the only systems to simulate the interaction between parts of a message molecule. Therefore, we chose both models for the simulation.

The kTAM model represents the interaction between parts of a message molecule. The interaction parameters can be chosen in such a way that the required number of binding reactions is the same as in the 2HAM model. In our simulations, we used the same interaction parameters as in the kTAM model. The number of binding reactions was not the only parameter that influences the assembly speed. The temperature parameter \( \tau \) also influences the assembly speed. The higher the temperature parameter, the faster the assembly.

The 2HAM model was used to simulate the interaction between the receptors and the message molecule. The interaction parameters were chosen in such a way that the required number of binding reactions is the same as in the kTAM model. In our simulations, we used the same interaction parameters as in the 2HAM model. The number of binding reactions was not the only parameter that influences the assembly speed. The temperature parameter \( \tau \) also influences the assembly speed. The higher the temperature parameter, the faster the assembly.

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Fig. 11. The fully assembled message molecule from a kTAM simulation with ISUTAS at \( G_{se} = 10.4 \) and \( G_{mc} = 17.0 \) after 3000 binding reactions.

Fig. 12. Result of 100 message molecule assembly simulations. The histogram shows the number of finished molecules with their respective number of binding reactions.

Table 1
Size and number of possible partial assemblies and the final assembly that may occur simultaneously when many seeds are present at once.

<table>
<thead>
<tr>
<th>Size</th>
<th>Number</th>
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<tbody>
<tr>
<td>1</td>
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Fig. 13. The message molecule (right) binding at the receptor (left).

Sierpinski triangle experiment. [25] shows the result of a wet lab experiment with DNA-tiles. In reality, DNA-based tiles assemble like a hybrid of both kTAM and 2HAM. The assembled Sierpinski triangle occurs multiple times in the medium. Apart from that, the assembly process behaves the same as proposed in our scenario. A tile concentration of 0.2 \( \mu \)M for every tile has been used, which is 1/4 of the simulated concentration. For comparison, the sodium levels in one liter of human blood are 0.14 \( \mu \)M. The results show that structures with up to 100 tiles can self-assemble with low error rates.

Since the proposed message molecule consists of only 15 tiles, it appears likely that it self-assembles without errors. Extrapolating from the results of [25], we propose to use a tile concentration of 0.2 \( \mu \)M for every tile type.

Derived parameters. A comparison between the simulation and the physical experiments is difficult. At a tile concentration of 0.8 \( \mu \)M each tile, the simulator completed a message molecule after on average 3000 steps. The wet lab experiment was terminated after one hour at a tile concentration of 0.2 \( \mu \)M for each tile. The simulated message molecule contained 15 tiles, while the resulting assemblies of the wet lab experiment contained on average 100 tiles. By extrapolating from these results, we assume that it would take at most 10 min to fully assemble a message molecule of size 15. In general, bigger assemblies grow faster due to the bigger border.

For the number of required nanosensors and nanobots, a lower bound can be given. To ensure that for every seed-tile a message molecule may assemble, we have to guarantee that every conditional-tile occurs at least as often in the medium/the nanosensors. If placed in a DNA-box of size \( 42 \times 36 \times 36 \) nm\(^3\) [22], 472.5 conditional tiles of size \( 14.4 \times 4 \times 2 \) nm\(^3\) [25] can fit into the volume of a box. Equally distributed, \( 1.275 \times 10^{10} \) DNA boxes would be necessary to harbor the amount of tiles used in the experiment with the volume of 50 \( \mu \)l [25].

In theory, we can freely choose how to distribute the conditional tiles among the nanosensors. In [25], the amount of basic components were overestimated a little to ensure proper assembly of the tiles—therefore we propose to do the same.

In summary, we expect the assembly of message molecules and receptors to properly function in a controlled environment like a petri dish. Given a high enough concentration of message molecules – which can be chosen appropriately – we expect the
molecular messages to assemble. In the presented scenario, a high particle density is of little concern.

The binding reaction between receptor and message molecule works like other molecular communication approaches that are based on Brownian motion with or without drift. We refer to [27] for a detailed analysis and a case study on molecular communication in e.g., the blood stream.

4.2. Information exchange efficiency

Table 2 displays a list of molecular communication categories and their corresponding capabilities. We evaluate the approaches with regard to underlying principle, propagation method, biological invasiveness and bandwidth in bits per particle. Since message molecules constitute of tiles, they can be arbitrarily big. Each tile can encode at least 1 Bit of information.

Table 2 shows that most techniques employ diffusion as a method for propagation. Only bacteria-based approaches differ, as they are capable of chemotaxis. Particle-based approaches usually use endogenous molecules, which can interfere with the body’s own communication. Tiles and some bacteria probably will not conflict. We thus consider them to be less bio-invasive.

Tile-based molecular communication exhibits additional properties, which to the best of our knowledge are not possible with other approaches:

1. Message molecules can compute decision problems [6].
2. Message molecules can transport binary information in a constructive manner, e.g., by labeling parts of a tile with fluorescence markers.
3. Nanonetworks with tile-based message molecules can employ error correction for in-message computation [28].

Nevertheless, it is difficult to compare parts of our approach with other communication models. The hierarchical process of message molecule assembly is simply not represented. If only fully assembled message molecules are considered, they can be simulated in the same way as molecular communication approaches that are based on particle types.

5. Conclusion

DNA-based molecular communication in nanonetworks is a bio-compatible way of exchanging messages within nanonetworks ex vivo. When message molecules comprise of DNA-tiles, they can incorporate computation capabilities into the messages. This in turn eases the computational burden on single nanobots, which become easier to design and construct.

This work introduced a modeling framework for molecular communication with self-assembling message molecules built from DNA-tiles. By drawing tiles from the ambient medium, these messages assemble in specific patterns. The exact assembly result depends on the seed and conditional tiles released from nanobots. A message may not fully assemble unless a logical condition is met. In the example scenario, four different nanobots have to release a conditional tile each in order to assemble a message, thereby implementing a distributed consensus. More generally, a message can compute arbitrarily complex decision problems (for example, AND, OR, XOR, MAJORITY, THRESHOLD...) and only fully assemble if these evaluate to true.

The self-assembly property of DNA-tiles yields a constructive and reproducible implementation. The resulting message molecules possess a huge variety of structure and binding properties, while maintaining the same size. Careful design of the comprising tiles and terminal messages can reduce assembly errors, for example by adding tiles without glue to the outside of a message.

While on average approximately 3000 binding reactions are necessary to fully assemble a message molecule, this fact is of little concern in experiments with DNA. The monomer concentration can be increased until satisfactory results are met. As long as $2G_{se} > G_{mc} > G_c$, holds, tile associations are more likely than tile disassociations and a fully assembled message molecule will eventually form, given enough time [26].

DNA-tile based message molecules are naturally bio-compatible as well as bio-degradable. If applied in a living organism, the lifetime of message molecules is naturally limited by filter organs that influence DNA. Furthermore, very few antigens against DNA or RNA exist. The construction of DNA-tiles and DNA-origami is already possible in laboratories. Both are well tested, and can probably be combined. As such, DNA-tiles are a complete scheme for computing and molecular communication, which is implementable today.

Assembled message molecules may be inspected by outside observers. For example, some tiles can receive fluorescent labels, whose combination within a molecule can be detected from outside. A controller may even further influence or stop the assembly process by adding inhibitor-tiles or controlling the ambient temperature.

In summary, we presented and evaluated a scenario in which a tile-based nanonetwork can be applied to diagnose diseases ex vivo with high confidence. This is possible, since the involved nanobots calculate a distributed consensus about the presence of markers by a 4-bit AND.

5.1. Future work and open questions

The example scenario for a 4-bit AND message breaks down into five major parts. Each part can be verified in wet lab experiments on their own. Experiments regarding construction of message molecules and their interaction with receptors have already been conducted. Other parts, especially reliable tile dispensation, will need further investigation.

In this work, we presented only simple logical functions using tile-based self-assembly systems. Future work should analyze more complex approaches that have been proposed to be utilized in medical scenarios. We want to investigate how to combine the basic tilesets for logical operators into more complex logical formulas, and the implications for a tile-based messaging scheme under realistic conditions.

In a last step, we want to test the whole proposed scenario in a wet lab experiment, preferably in a petri dish and later in vivo.

In addition to wet lab experiments, it would be beneficial to simulate the presented scenario as accurate as possible. The human blood stream is of special interest. Previous experiments have shown the general feasibility of the approach ex vivo. For this, a complete model of the communication channel with the most important environmental parameters is necessary. Those include flow effects in the channel, collision detection, binding reactions in a fluid medium and tile diffusion.
References


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