

Stochastic Strategies for a Swarm Robotic Assembly System

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Abstract—We present a decentralized, scalable approach to assembling a group of heterogeneous parts into different products using a swarm of robots. This activity forms the basis for a reconfigurable manufacturing system. While the assembly plans are predetermined, the exact sequence of assembly of parts and the allocation of subassembly tasks to robots are determined by the interactions between robots in a decentralized fashion in real time. Our approach is based on developing a continuous abstraction of the system derived from models of chemical reactions and formulating the strategy as a problem of selecting rates of assembly and disassembly. These rates are mapped onto probabilities that determine stochastic control policies for individual robots, which then produce the desired aggregate behavior. This top-down approach to determining robot controllers also allows us to optimize the rates at the abstract level to achieve fast convergence to the specified target numbers of products. Because the method incorporates programs for assembly and disassembly, changes in demand can lead to reconfiguration in a seamless fashion. We illustrate the methodology using a physics-based simulator with examples involving 15 robots and two types of final products.

I. INTRODUCTION

We develop an approach to designing a reconfigurable manufacturing system in which a swarm of homogeneous robots assembles static parts into different types of products. The system must respond quickly to produce desired amounts of products, which can change at discrete points in time, from any initial set of parts. Since it is difficult, if not impossible, to efficiently control a large robot population through centralized algorithms, we employ a decentralized strategy in which robots operate autonomously and use local communication. The strategy can be readily implemented on resource-constrained robots, and it is scalable in the number of robots and parts and robust to changes in robot population.

The robots move randomly within a closed arena, encountering parts and other robots at probability rates that are determined by the physical parameters of the system and by sensor and environmental noise. We model this behavior using a realistic 3D physics simulation, which we refer to as a *micro-continuous model* since it encompasses the continuous dynamics of individual robots. The system can be approximated as well-mixed, and the robot-part and

robot-robot interactions are analogous to chemical reactions between molecules. Hence, like a set of chemical reactions, our system can be represented by the Stochastic Master Equation [1]. Various methods exist to numerically simulate the evolution of a system governed by this model [2], [3]. We call this the *complete macro-discrete model* because it describes a continuous-time Markov process whose states are the discrete numbers of system components. When there are large numbers of parts, the system can be abstracted to an ordinary differential equation (ODE) model, which we call a *macro-continuous model* since its state variables are continuous amounts of the parts.

Our work is similar in objective to studies on programmable self-assembly for modular robots [4]–[7]. The system in this work is a set of homogeneous triangular robots that move around randomly on an air table and assemble into components according to a plan that is constructed using *graph grammars*, which are rules that define interactions between robots. Graph grammars can be constructed automatically to produce a given predefined assembly [7]. The probabilities of a newly formed component to detach into different combinations of parts can be optimized to maximize the number of a desired assembly at equilibrium [5]. However, this optimization requires the enumeration of all system states reachable from the initial state.

Our approach to assembly system design allows for improved scalability and flexibility. We consider a scenario in which a heterogeneous set of parts is assembled into two types of final products. To provide theoretical guarantees on performance, we employ the “top-down” design methodology presented in [8]–[10] for reallocating a swarm of robots among a set of sites/tasks in a desired distribution. This methodology was applied to a linear model; here we extend it to a nonlinear (specifically, multi-affine) model with robot interactions. We construct this *complete macro-continuous model* of the system using the Chemical Reaction Network (CRN) framework [11], [12], which has been studied extensively for theoretical insight into biochemical systems. We compute reaction rate constants in the model from physical properties of the robots and environment and check that the model accurately predicts the results of the micro-continuous model. Then we simplify this abstraction to a *reduced macro-continuous model* with the same rates and use the model to optimize these rates for fast convergence to a target distribution of products, using an approach similar to [13], [14]. The optimization problem is independent of the number of parts and scales only with the number of rates. We simulate the reduced model with optimized rates for different target distributions, map the rates onto probabilities

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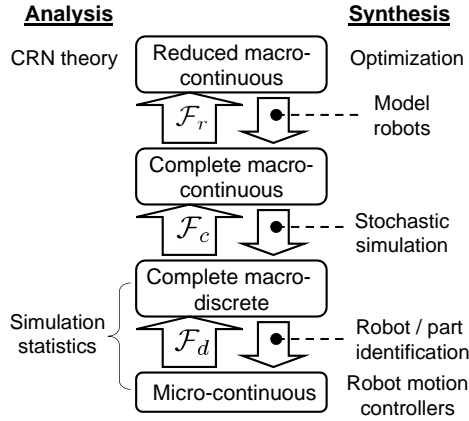


Fig. 1. Levels of abstraction of the assembly system with analysis and synthesis methodologies. The high-dimensional micro-continuous model is mapped to lower-dimensional representations, the macro-discrete and macro-continuous models, through the abstractions \mathcal{F}_d and \mathcal{F}_c using the theoretical justification in [2]. Under certain assumptions (see Section III-C), the complete macro-continuous model can be mapped to a lower-dimensional model via the abstraction \mathcal{F}_r .

of assembly and disassembly which are used as control policies in the micro-continuous model, and observe this system's convergence behavior.

Fig. 1 illustrates the relationships among the four levels of system abstraction discussed and the methods we use for designing the system and analyzing its behavior.

II. PROBLEM STATEMENT

A. Assembly task

There are four different types of parts, numbered 1 through 4, which can be combined to form larger parts according to the assembly plans in Fig. 2. Parts bond together through bi-directional connections at sites along their perimeters. The last step in each plan is the production of a final assembly, $F1$ or $F2$. The assembly task is executed by a group of robots in an arena that is sufficiently large to ignore the dynamics of small-scale interactions. Initially, robots and many copies of parts of type 1 through 4 are randomly scattered throughout the arena. There are exactly as many of these parts as are needed to create a specified number of final assemblies, and the number of robots equals the total number of scattered parts. Each robot knows the assembly plans a-priori and has the ability to recognize part types, pick up a part, combine it with one that is being carried by another robot, and disassemble a part it is carrying.

Our objective is to define robot controllers for moving around the arena and for picking up, assembling, and disassembling parts so that the robots produce target numbers of final assemblies as quickly as possible.

B. Micro-continuous model

We implement the assembly task in the robot simulator Webots [15], which uses the Open Dynamics Engine to accurately simulate physics. We use the robot platform Khepera III, which has infra-red distance sensors for collision

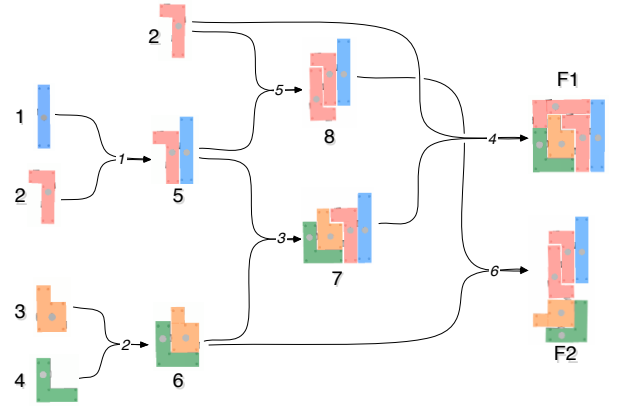


Fig. 2. Assembly plans for final assemblies $F1$ and $F2$.

avoidance. Each robot is outfitted with a protruding bar with a rotational servo at the tip. A magnet on the servo bonds to a magnet on the top face of a part, and the servo is used to rotate the bonded part into the correct orientation for assembly. Parts bond to each other via magnets on their side faces. Magnets can be turned off to deactivate a bond. Robots and parts are equipped with a radio emitter and receiver for local communication and for computing relative bearing, which is used to align robot and part magnets and to rotate a part for assembly. The task takes place inside the walled hexagonal arena shown in Fig. 3.

We use a control policy for the robots that is inspired by chemical processes: random movement patterns with probabilistic assemblies upon encounter, as well as random disassemblies. Our models assume that the system is well-mixed; to achieve this property, robots move according to a random walk, and we verify that the space is uniformly covered. Robots and parts switch between action states based on information they receive via local sensing and communication. When a robot encounters a part on the ground, it approaches and bonds to it and starts searching for a robot that is carrying a compatible part, according to the assembly plans. When one is found, the two robots align their parts and approach each other to join the parts. One robot carries off the newly assembled part while the other resumes searching for a part on the ground. A robot can disassemble a part it is carrying by dropping one of the component parts on the ground. To control the outcome of part populations, we can directly modify the probabilities of starting an assembly and performing a disassembly.

III. MACRO-CONTINUOUS MODELS

A. Definitions

Interactions between parts and robots in the assembly system are modeled in the form of a Chemical Reaction Network (CRN). A set of reactions can be represented as a directed graph, $\mathcal{G} = (\mathcal{V}, \mathcal{E})$. The set of vertices, \mathcal{V} , signifies the *complexes*, which are the combinations of parts and/or robots that appear before and after reaction arrows. The set of directed edges, \mathcal{E} , represents the reaction pathways between

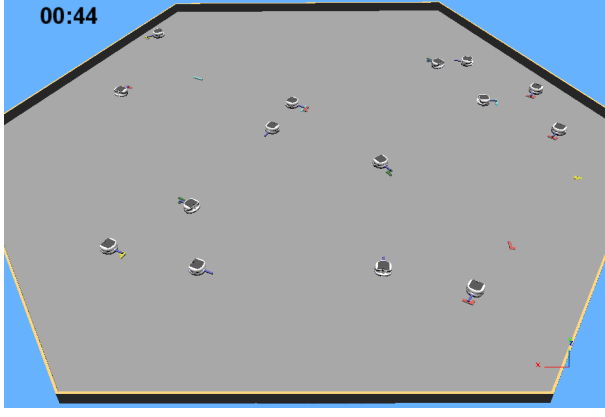


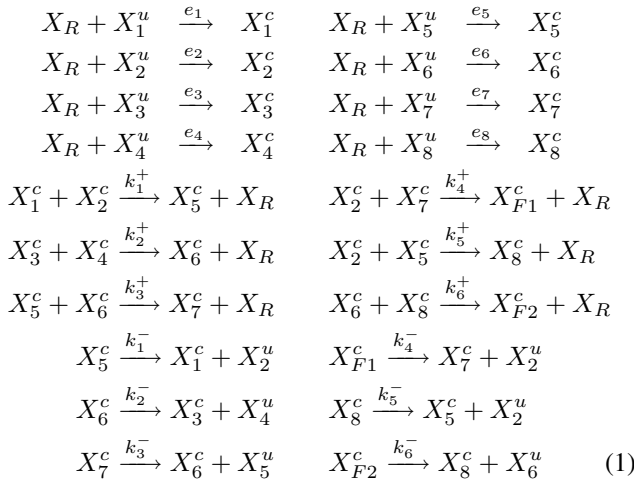
Fig. 3. Snapshot of the arena in the realistic physical simulation. Robots carry parts at the end of a protruding bar.

the complexes. Each pathway is denoted by an ordered pair $(i, j) \in \mathcal{V} \times \mathcal{V}$, which means that that complex i transforms into complex j , and is associated with a positive *reaction rate constant*.

Each part of type i in Fig. 2 is symbolized by X_i , and a robot is symbolized by X_R . X_i may be further classified as X_i^u , an unclaimed part on the ground, or as X_i^c , a claimed part i and the robot that is carrying it. Let M be the number of these variables, or *species*, in a model of the system. Then $\mathbf{x}(t) \in \mathbb{R}^M$ is the vector of the species populations, which are represented as continuous functions of time t .

B. Complete macro-continuous model

We define a CRN that represents each possible action in the micro-continuous model of the assembly system:



In this CRN, e_i is the rate at which a robot encounters a part of type i , k_i^+ is the rate of assembly process j , and k_i^- is the rate of disassembly process j . We theoretically estimate these rates as functions of the following probabilities:

$$e_i = p^e, \quad k_i^+ = p^e \cdot p_i^a \cdot p_i^+, \quad k_i^- = p_i^- . \quad (2)$$

p^e is the probability that a robot encounters a part or another robot. Using the assumption that robots and parts are distributed uniformly throughout the arena, we calculate p^e

from the geometrical approach that is used to compute probabilities of molecular collisions [16], [17]: $p^e \approx vTw/A$, where v is the average robot speed, T is a timestep, A is the area of the arena, and w is twice a robot's communication radius, since this is the range within which a robot detects a part or robot and initiates an assembly process.

p_i^a is the probability of two robots successfully completing assembly process i ; it depends on the part geometries.

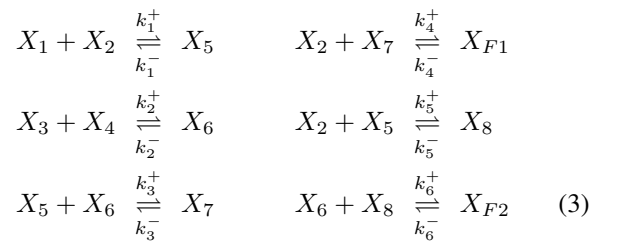
p_i^+ is the probability of two robots starting assembly process i , and p_i^- is the probability per unit time of a robot performing disassembly process i . These are the *tunable parameters* of the system.

We compute p_i^a and the parameters for p^e using measurements from the micro-continuous model (Webots simulations): $A = 23.4 \text{ m}^2$ (hexagon of radius 3 m), $w = 1.2 \text{ m}$, $v_R = 0.128 \text{ m/s}$ from an average over 50 runs, and $\mathbf{p}^a = [0.9777 \ 0.9074 \ 0.9636 \ 0.9737 \ 0.8330 \ 1.0]$ (entries follow the numbering of the associated reactions) from averages over 100 runs. We set $T = 1 \text{ s}$.

In the thermodynamic limit, which includes the condition that populations approach infinity, the physical system represented by (1) can be abstracted to an ODE model [2]. This is illustrated in the next section. We numerically integrate this macro-continuous model with the rates we calculated and also use the StochKit toolbox [18] to efficiently perform a stochastic simulation of the macro-discrete model. We compare the results to those for the micro-continuous model in Fig. 4, using $p_i^+ = 1$, $p_i^- = 0 \ \forall i$. The results for all models are very similar, although discrepancies arise from two factors. First, certain events that happen in the physical simulation are not modeled: parts sticking together or being wedged against a wall, and deviations from the assumption of uniform spatial distribution. Second, the ODE approximation is valid only for large numbers of parts, and the system modeled had only 15 robots and 15 parts. Overall, the macro-continuous model accurately predicts the evolution of part populations, and hence we can use it to design the rates to direct the system's behavior, provided that the system has very large numbers of parts.

C. Reduced macro-continuous model

We simplify the complete model by abstracting away robots and retaining only interactions between parts, assuming that the time to find a part is small and there are more robots than parts:



The rates are also defined by equation (2).

We define a vector $\mathbf{y}(\mathbf{x}) \in \mathbb{R}^{12}$ in which entry y_i is the

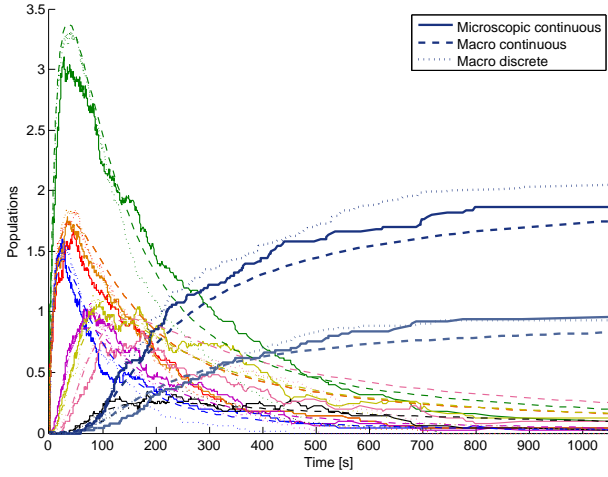


Fig. 4. Part populations in the complete macro-continuous, macro-discrete, and micro-continuous models for 3 final assemblies and 15 robots. Macro-discrete and micro-continuous results are averaged over 100 runs.

part or product of parts in complex i :

$$\mathbf{y}(\mathbf{x}) = \begin{bmatrix} x_1 x_2 & x_5 & x_3 x_4 & x_6 & x_2 x_7 & x_{F1} \\ x_5 x_6 & x_7 & x_2 x_5 & x_8 & x_6 x_8 & x_{F2} \end{bmatrix}^T. \quad (4)$$

We also define a matrix $\mathbf{M} \in \mathbb{R}^{10 \times 12}$ in which each entry M_{ji} , $j = 1, \dots, 10$, of column \mathbf{m}_i is the coefficient of part type j in complex i (0 if absent). We relabel the rate associated with reaction $(i, j) \in \mathcal{E}$ as k_{ij} and define a matrix $\mathbf{K} \in \mathbb{R}^{12 \times 12}$ with entries

$$K_{ij} = \begin{cases} k_{ji} & \text{if } i \neq j, (j, i) \in \mathcal{E}, \\ 0 & \text{if } i \neq j, (j, i) \notin \mathcal{E}, \\ -\sum_{(i,l) \in \mathcal{E}} k_{il} & \text{if } i = j. \end{cases} \quad (5)$$

Then our ODE abstraction of the system can be written in the following form [19]:

$$\dot{\mathbf{x}} = \mathbf{MKy}(\mathbf{x}). \quad (6)$$

One set of linearly independent conservation constraints on the part quantities is:

$$\begin{aligned} x_3 - x_4 &= N_1 \\ x_1 + x_5 + x_7 + x_8 + x_{F1} + x_{F2} &= N_2 \\ x_2 + x_5 + x_7 + 2(x_8 + x_{F1} + x_{F2}) &= N_3 \\ x_3 + x_6 + x_7 + x_{F1} + x_{F2} &= N_4 \end{aligned} \quad (7)$$

where N_i , $i = 1, \dots, 4$, are computed from the initial part quantities.

Theorem 1: System (6) subject to (7) has a unique, stable equilibrium $\bar{\mathbf{x}} > \mathbf{0}$.

Proof: Each equilibrium of the system, $\{\bar{\mathbf{x}} \mid \mathbf{MKy}(\bar{\mathbf{x}}) = \mathbf{0}\}$, can be classified as either a *positive* equilibrium $\bar{\mathbf{x}} > \mathbf{0}$ or a *boundary* equilibrium in which $\bar{x}_i = 0$ for some i , which can be found by solving $\mathbf{y}(\bar{\mathbf{x}}) = \mathbf{0}$ [19]. From definition (4) of $\mathbf{y}(\mathbf{x})$, it can be concluded that in each boundary equilibrium, all $x_i = 0$ except for one of the four combinations of variables (x_1, x_3) , (x_1, x_4) , (x_2, x_3) , (x_2, x_4) . Since we only consider systems that can produce x_{F1} and x_{F2} , it is not possible

for the system to reach any of these equilibria; each one lacks two part types needed for the final assemblies.

The *deficiency* δ of a reaction network is the number of complexes minus the number of linkage classes, each of which is a set of complexes connected by reactions, minus the network rank, which is the rank of the matrix with rows $\mathbf{m}_i - \mathbf{m}_j$, $(i, j) \in \mathcal{E}$ [20]. Network (3) has 12 complexes, 6 linkage classes, and rank 6; hence, $\delta = 0$. Also, the network is *weakly reversible* because whenever there is a directed arrow pathway from complex i to complex j , there is one from j to i . Because the network has deficiency 0, is weakly reversible, and does not admit any boundary equilibria, it has a unique, globally asymptotically stable positive equilibrium according to Theorem 4.1 of [21]. ■

IV. RATE OPTIMIZATION

We consider the problem of designing the system described by model (6) subject to (7) to produce desired quantities of parts as quickly as possible. The objective will be posed as the design of *optimal rates* k_i^+ , k_i^- , $i = 1, \dots, 6$, which define an *optimal rate matrix* \mathbf{K}^* according to (5), that minimize the convergence time of the system to a vector of target part quantities, \mathbf{x}^d . Note that although only the amounts of the final assemblies $F1$ and $F2$ may need to be specified in practice, our optimization problem requires that target quantities of *all* parts be defined.

We first specify $x_1^d, x_2^d, x_3^d, x_5^d, x_8^d$ and a parameter

$$\alpha \equiv x_{F1}^d / (x_{F1}^d + x_{F2}^d). \quad (8)$$

Then we compute the dependent variables x_4^d, x_6^d, x_7^d , and $x_{F1}^d + x_{F2}^d$ from the conservation equations (7) and definition (8) and check that they are positive to ensure a valid \mathbf{x}^d . In this way, we can keep $x_{F1}^d + x_{F2}^d$ and the target non-final part quantities constant while adjusting the ratio between x_{F1} and x_{F2} using α . Theorem 1 shows that we can achieve \mathbf{x}^d from any initial distribution \mathbf{x}^0 by specifying that $\bar{\mathbf{x}} = \mathbf{x}^d$ through the following constraint on \mathbf{K} ,

$$\mathbf{MKy}(\mathbf{x}^d) = \mathbf{0}. \quad (9)$$

Now we consider the aspect of minimizing the convergence time to \mathbf{x}^d . We quantify this time in terms of the system *relaxation times* τ_i , $i = 1, \dots, 6$, the times in which different modes (dynamically independent variables) of the the system converge to a stable equilibrium after perturbation [22], [23]. Various measures of the average relaxation time of a CRN have been defined, but they are applicable only under certain conditions, such as a linear reaction sequence [24]. For instance, one such measure was minimized in the optimization of rates for the linear chain in [25].

To estimate the relaxation times, we first reformulate the system in terms of new variables. Define v_i , $i = 1, \dots, 6$, as the difference between the forward and reverse fluxes associated with reaction i in system (3). For example, $v_1 = k_1 x_1 x_2 - k_2 x_5$. Let $\mathbf{v}(\mathbf{x}) = [v_1 \dots v_6]^T$ and let $\mathbf{S} \in \mathbb{R}^{6 \times 10}$

denote the stoichiometric matrix of the system, which is defined such that model (6) can be written as [26]:

$$\dot{\mathbf{x}} = \mathbf{S}\mathbf{v}(\mathbf{x}). \quad (10)$$

The dynamical properties of a CRN are often analyzed by linearizing the ODE model of the system about an equilibrium and studying the properties of the associated Jacobian matrix $\mathbf{J} = \mathbf{S}\mathbf{G}$, where the entries of \mathbf{G} are $G_{ij} = dv_i/dx_j$ [23]. Denoting the eigenvalues of \mathbf{J} by λ_i , a common measure of relaxation time is $\tau_i = 1/|\text{Re}(\lambda_i)|$. Since the λ_i are negative at a stable equilibrium, one way to yield fast convergence is to choose rates that minimize the largest λ_i . However, in our system it is very difficult to find analytical expressions for the λ_i . We use an alternative estimate of relaxation time that is also derived by linearizing the system around its equilibrium \mathbf{x}^d [26],

$$\tau_j = \left(\sum_{i=1}^{10} (-s_{ij}) \frac{dv_j}{dx_i} \right)_{\mathbf{x}=\mathbf{x}^d}^{-1}. \quad (11)$$

Each reaction j in system (3) is of the form $X_k + X_l \xrightleftharpoons[k_j^-]{k_j^+} X_m$. Thus, $v_j = k_j^+ x_k x_l - k_j^- x_m$, and the entries of column j in \mathbf{S} are all 0 except for $s_{kj} = s_{lj} = -1$ and $s_{mj} = 1$. Then according to equation (11), the relaxation time for each reaction is

$$\tau_j = (k_j^+ (x_k^d + x_l^d) + k_j^-)^{-1}. \quad (12)$$

Define $\mathbf{k} \in \mathbb{R}^{12}$ as the vector of all rates k_i^+, k_i^- . Using equation (12), we define two possible objective functions $f: \mathbb{R}^{12} \rightarrow \mathbb{R}$, the average τ_j^{-1} and the minimum τ_j^{-1} , to maximize in order to produce fast convergence to \mathbf{x}^d :

$$f_{ave}(\mathbf{k}) = \frac{1}{6} \sum_{j=1}^6 \tau_j^{-1}, \quad (13)$$

$$f_{min}(\mathbf{k}) = \min\{\tau_1^{-1}, \dots, \tau_6^{-1}\}. \quad (14)$$

Finally, we write the rates k_i^+, k_i^- in terms of the tunable probabilities p_i^+, p_i^- using equation (2) and define these probabilities as the optimization variables. Let $\mathbf{p} \in \mathbb{R}^{12}$ be the vector of all p_i^+, p_i^- . Then the optimization problem can be posed as **Problem P** below. It will be referred to as **Problem P1** when $f = f_{ave}$ and as **Problem P2** when $f = f_{min}$.

$$\begin{aligned} [\mathbf{P}] \quad & \text{maximize} \quad f(\mathbf{k}(\mathbf{p})) \\ & \text{subject to} \quad \mathbf{M}\mathbf{K}(\mathbf{p})\mathbf{y}(\mathbf{x}^d) = \mathbf{0}, \quad \mathbf{0} \leq \mathbf{p} \leq \mathbf{1}. \end{aligned}$$

Problems P1 and P2 are both linear programs, which can be solved efficiently. To check that they do in fact minimize convergence time, we implemented a Monte Carlo method [27], which is more computationally expensive, to find the $\mathbf{k}(\mathbf{p})$ that directly minimizes this time. We measure the degree of convergence to \mathbf{x}^d by $\Delta(\mathbf{x}) = \|\mathbf{y}(\mathbf{x}) - \mathbf{y}(\mathbf{x}^d)\|_2$ and say that one system converges faster than another if it takes less time for $\Delta(\mathbf{x})$ to decrease to some small fraction, here defined as 0.1, of its initial value. At each iteration, $\mathbf{k}(\mathbf{p})$ is perturbed by a random vector and projected onto

the null space of linearly independent rows of a matrix \mathbf{N} defined such that $\mathbf{N}\mathbf{k} = \mathbf{M}\mathbf{K}\mathbf{y}(\mathbf{x}^d) = \mathbf{0}$. Once $\mathbf{k}(\mathbf{p})$ also satisfies $\mathbf{0} \leq \mathbf{p} \leq \mathbf{1}$, it is used to simulate the reduced macro-continuous ODE model to find $\Delta(\mathbf{x})$ after some time. Since the system is stable by Theorem 1, $\Delta(\mathbf{x})$ always decreases monotonically with time, so a Newton scheme can be used to compute the exact time $t_{0.1}$ when $\Delta(\mathbf{x}) = 0.1\Delta(\mathbf{x}^0)$.

V. RESULTS

A. Optimization of rates

We solved optimization problems P1 and P2 for $\alpha \in \{0.01, 0.02, \dots, 0.99\}$ using $\mathbf{x}^0 = [60 \ 120 \ 60 \ 60 \ 0]^T$ and $\mathbf{x}^d = [0.5 \ 2.5 \ 1 \ 1 \ 0.5 \ 1 \ 1 \ 1 \ 57\alpha \ 57(1-\alpha)]^T$. As Table I shows, the computed rates are constant for each α except for those corresponding to assembly and disassembly processes 4 and 6. This means that the system is flexible enough to yield any α when only the rates of assembling and breaking apart the final assemblies are modified. We also ran the Monte Carlo program for $\alpha = 0.4$. For the optimal set of rates, which were very similar to those for Problem P1, $t_{0.1} = 4.69$, as compared to 4.68 for Problem P1 and 8.49 for P2. This provides evidence that Problem P1 is minimizing the system convergence time.

We integrated the reduced macro-continuous model for $\alpha = 0.4$ and $\alpha = 0.8$ using the optimized rates from Problems P1 and P2 and a set of non-optimal rates that were chosen to satisfy constraint (9) and $\mathbf{0} \leq \mathbf{p} \leq \mathbf{1}$ but were not optimized for some objective. The evolution of the model for each set of rates is shown in Fig. 5, with time in log-scale. The optimized systems first quickly converge to an “intrinsic equilibrium” at $t \approx 10^2$ that is independent of α and then redistribute much more slowly to the target ratios, which are attained for all set of rates. For both α , the optimized models converge faster to equilibrium than the non-optimal model. For $\alpha = 0.4$, the non-optimal model displays a large over- and under-shooting of the target ratios, whereas the optimized models converge quickly with little or none of this effect. Problem P2 produces a more efficient system in this respect, although both optimized models converge at comparable rates.

B. Mapping rates onto the micro-continuous model

For $\alpha = 0.4$, we mapped the rates optimized by Problem P1 and the non-optimal rates onto the micro-continuous model to see whether the physical system would behave similarly to the reduced continuous model. We did this in the following way. Let R be a uniformly distributed random number between 0 and 1 and let Δt be the simulation timestep. A robot carrying a part that can be disassembled according to process i computes R at each timestep and disassembles the part if $R < p_i^- \Delta t$. A robot about to begin assembly process i computes R and executes the assembly if $R < p_i^+ \Delta t$. Fig. 6 shows the time evolution of the micro-continuous model averaged over 30 runs for both sets of rates, using 15 robots and 15 parts (3 final assemblies). The non-optimal model converges to the target ratios but initially over- and under-shoots x_{F1}^d and x_{F2}^d , respectively,

TABLE I

VALUES OF OPTIMIZED RATES FOR VARYING α . Continuous rates evolve continuously with respect to α .

Reaction i	1	2	3	4	5	6
P1 Optimized p_i^+	1.0					
P1 Optimized p_i^-	0.01885	0.00754	0.00377	continuous	0.00942	continuous
P2 Optimized p_i^+	0.36	0.666	1.0	continuous	0.4705	continuous
P2 Optimized p_i^-	0.006855	0.005027	0.00377	continuous	0.00443	continuous

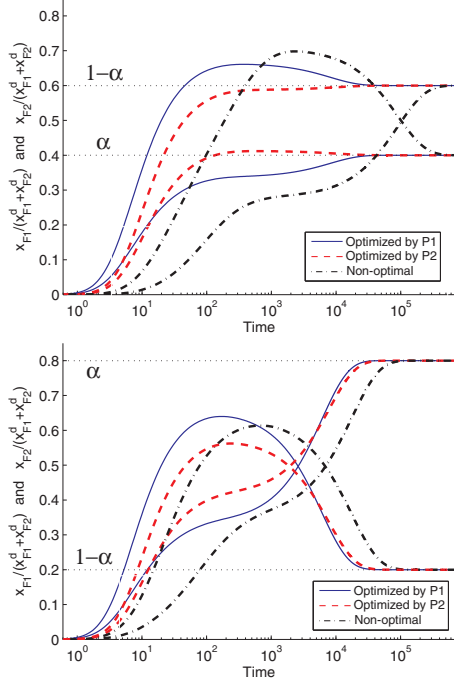


Fig. 5. Evolution of final assembly ratios in the reduced macro-continuous model for $\alpha = 0.4$ (top) and $\alpha = 0.8$ (bottom), using rates optimized by Problems P1 and P2 and non-optimal rates.

as in Fig. 5. The ratios in the optimized model rise quickly to the target ratios but then deviate from them toward 0.5.

The discrepancy between the optimized model results and the target ratios can be explained by inaccurate modeling of some low-level effects. In the micro-continuous model, when a robot disassembles a part, a nearby robot is likely to pick up the fallen component and quickly reassemble the original part with the other robot, which generally hasn't traveled far. This increases the rate of assembly of the part and violates the well-mixed assumption. In addition, robots that pick up dropped parts spend a significant amount of time carrying them before reassembly, a factor that was abstracted away in the reduced model. Adding more robots tends to alleviate this effect. Random failures at disassembly, incorrect internal state definitions by the robots and parts, and collision errors by the physics engine sometimes occur in the simulation but are not modeled. Finally, the continuous abstraction may not accurately capture the system behavior for such a low number of parts. However, it becomes more computationally expensive to simulate the system as the number of parts and robots increase.

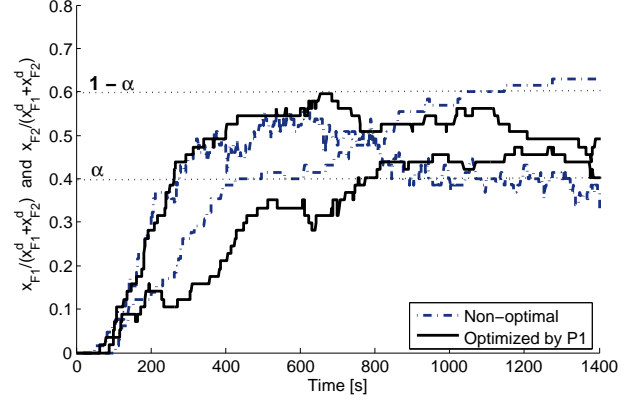


Fig. 6. Evolution of final assembly ratios in the micro-continuous model for $\alpha = 0.4$, using rates optimized by Problem P1 and non-optimal rates.

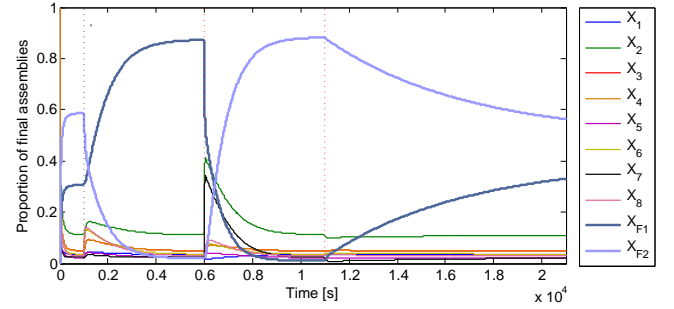


Fig. 7. Online adaptation of the reduced macro-continuous model to changes in target equilibrium, which occur at the vertical dashed lines.

C. Reconfigurable manufacturing

We apply our methodology to a “green manufacturing” task, in which finished products are recycled into new products in ratios that depend on the current demand. We perform a simulation of the reduced macro-continuous model that corresponds to a situation in which robots switch between sets of optimized rates at discrete points in time to produce different target numbers of parts. The sequence of target α is 0.4, 0.99, 0.01, and 0.5, and \mathbf{x}^0 is the same as in Section V-A. The system evolution is shown in Fig. 7 using rates from Problem P1. The system quickly adapts to new target ratios, although some take longer to attain than others (e.g. $\alpha = 0.5$). The results demonstrate that high-level control of the system is possible in real-time.

VI. CONCLUSIONS AND FUTURE WORK

We developed and presented a method to systematically derive decentralized control policies for a swarm of robots

performing assembly tasks to manufacture different products in response to varying demand. We represent the system using a multi-level modeling methodology. The micro-continuous model is derived from the dynamics of the robots and the assembly task and is implemented on a physics-based simulator. The collective behavior of the system, including the physical interactions between robots, is abstracted to a macro-continuous ODE model. The model incorporates parameters that govern the stochastic control policies running on individual robots for performing the assembly task. By tuning the parameters of the ODE, we can also tune the performance of the assembly system. This optimization relies on global stability properties of a specific class of chemical reaction networks that are modeled by the macro-continuous model. We implement the optimization as a linear program with constraints on target amounts of parts at equilibrium, using two possible objective functions that are based on an estimate of the system convergence time. We simulate the macro-continuous model and observe that it achieves the target final product amounts faster with optimized rates than with non-optimal rates, and that it can quickly respond to changes in the target equilibrium. Finally, we map the rates onto probabilities of assembly and disassembly in the micro-continuous model. We find that the resulting system can produce the target product amounts, although discrepancies arise due to violation of the well-mixed property, low part numbers, and failure to capture certain physical effects in the macro-continuous model.

Our future work will focus on tuning the macro-continuous model to match the physical system more closely, addressing in particular the lack of spatial homogeneity due to a small number of robots and parts. In addition, we also want to investigate the synthesis of the discrete assembly plans and incorporate feedback into the process. This direction draws inspiration from bio-molecular pathways in which intermediate subassemblies or molecules can promote or inhibit chemical reactions. We would like to be able to optimize the discrete assembly plan by constructing feedback loops to improve the yield rate.

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