Self-Organizing Formation Algorithm for Active Elements

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In this paper, we propose a novel method of self-organizing formation. It is assumed that elements are not connected to each other, and they can move in continuous space. The objective is to arrange elements in certain spatial pattern like a crystal, and to make the outline of the group in desired shape. For this purpose, we proposed a method by using virtual springs among the elements. In this algorithm, an element generates virtual springs between neighbor element based on information how many other elements exist in neighborhood with a certain radius. Although the elements interact locally only by virtual springs, and they don't have global information at all, they form a shape much larger than the sensory radius. By simulation study, we confirmed convergence to a target shape from a random state in very high probability. This kind of algorithm gives a new principle of self-organizing formation, and its simplicity will be useful for design of self-assembling nano machines in future.

1 Introduction

There are various phenomena in which many identical elements make the whole form by self-organization. For example, in the growth process of a crystal, many atoms or molecules form regular lattice structure according to the principle of free energy minimization. Although atoms are very small and simple, when they aggregate, they make a symmetrical crystal in a macroscopic scale. Form of a living organism is also a result of self-organization process. Beginning from fertilization, it makes a peculiar form by dividing cells repeatedly and arranging them in a certain order. Primitive elements of this process are biological cells containing equivalent genetic information. The cells determine the form, through exchanging chemical substances. Although the behavior of a cell is farther complicated than the atom in a crystal, there is a common process in which global order of the system emerges from the cooperation of homogeneous elements influencing locally.

However, in the world of the artificial things, such kind of self-organization seldom appears. It is because an artifact is usually designed along a certain purpose, and its form and structure are optimized to realize desired purposes. However, in the artificial system of the next generation, the self-organization processes will be applied more purposefully. For example, for a machine in a nano scale, it is not realistic to pick a part and assemble them one by one. Like a protein molecule, many elements must self-aggregate to form a functional component (self-assembly). Self-organization is also important to realize self-repairing artifacts. For instance, several modular robot systems capable of self-repair have been considered recently. They consist of many homogeneous mechanical modules, and it can restore itself by replacing a broken module by a new module (self-repair). In such a system, it does not know beforehand which module will break. Therefore, the self-organization algorithm in which the group of elements cooperates and forms a target shape is needed.

Formation processes of homogeneous elements can be classified roughly into the following three classes.

- A. Elements are not connected to each other, and they can move in continuous space.
- B. Elements are connected to each other, and they are constrained to certain discrete relative positions.
- C. Elements are connected to each other, but continuous change in relative positions is allowed.

Class A is a system like a flock of birds or a school of fish, or a group of mobile robots as an artificial system. Class B is a system like a crystal or a snowflake, and its artificial example is a modular robot. Class C can be found only in natural systems such as protein aggregation or multi-cellular organisms.

Algorithms of self-organizing formation have been proposed for each class. For class A, Yamashita proposed an algorithm to align a group of homogeneous mobile robots in a circle¹. An alignment algorithm based on nonlinear reaction-diffusion is also proposed². Methods to control not a group's form but the dynamic behavior of a group are also studied for class A^{3,4}. For class B, a self-assembly algorithm for a modular robot is proposed based on local connection style among modules⁵, and a self-repair algorithm using hierarchical description of the robot shape is proposed for the same modular robot^{6,7,8}. About class C, although many models of developing organisms are proposed, it is thought that an artificial hardware corresponding to this class does not exist yet.

In this paper, we assume that elements are not connected to each other, and they can move in continuous space (class

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A). However, our target is like class B. Namely, the objective is to arrange elements in certain spatial pattern like a crystal. and to make the outline of the group in desired shape. For this purpose, as the simplest model which can consider the size of an element, we proposed a method by using virtual springs among the elements. In this algorithm, an element generates virtual springs between neighbor elements based on how many other elements exist in the neighborhood with a certain sensory radius. Although the elements interact locally by virtual springs, and they don't have global information at all, they form a shape much larger than the radius. By simulation study, we confirmed convergence to a target shape from a random state occurs in very high probability. This kind of algorithm gives a new principle of self-organizing formation, and its simplicity will be useful for design of self-assembling nano-machines in future.

2 Self-organizing Formation

2.1 Problem formulation

The objective of formulation problem is to arrange elements in predetermined order without depending on their initial position and direction (Figure 1). We assume properties of the elements as follows:

- All the elements have the same character.
- When a distance between two elements is less than a sensory radius R, they interact each other by a virtual spring between them.
- The number of elements which exist in R is countable.

2.2 Algorithm of self-organizing formation

We have developed an algorithm that utilizes the number of connection between elements under the problem setup above.

2.2.1 Connection number and virtual spring

In the algorithm, virtual springs shall be generated to all the elements that exist in R. We call the number of elements in the circle with a radius of R, number of connection. For example, elements a, b, c, d, e in Figure 2 have connections 2, 4, 2, 3, 3, respectively.

Virtual springs used in the algorithm are assumed to have a large spring constant k. By setting natural length of virtual springs l slightly smaller than R (i.e.90% of R), a triangular lattice is made among the elements (Figure 2). This is regarded as "nominal" setting. We determined the value of k and l by trial and error.

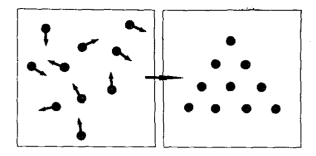


Figure 1: Self-organizing formation

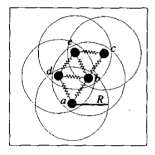


Figure 2: The number of connection and triangular lattice

2.2.2 Tuning virtual springs

In the formation problem, we need to control not only the internal structure pattern, but also the outline of the global structure. In order to do this, we changed characteristics of the virtual springs. Namely, spring constant and natural length of a spring is defined by the numbers of connection of the elements at the both ends of the springs.

For combinations of the number of connection exist in the target form, spring constant and natural length is set to their nominal values. In addition to them, we can separate elements with a certain combination of the number of connection by setting the natural length to a value bigger than R. Combination is canceled by separation, because springs vanish when the element pushed out of R. The product of the spring constant and the spring length can tune the strength of ejection.

We also put a probability P of existence to each virtual spring. By using the probability, we can control frequency of the separation.

For example, in the situation of Figure 3(left), assume that we want to separate a virtual spring a-c or b-c, because the combination of the number of connection 2-5 is not included in the target form. To do this, we can set the virtual spring 2-5 with large spring constant, natural length and probability of existence compared with nominal springs. As a consequent,

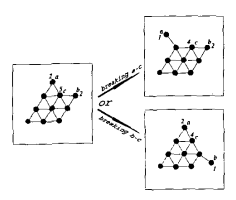


Figure 3: Breaking unnecessary bonds

spring a-c or b-c will be replaced by a large spring, and it pushes the elements, and then vanishes because it goes out of range. By introducing suitable combination of springs, we can generate desired shapes.

3 Simulation

3.1 Simulation Model

Elements move on a 2-Dimensional space. Subscript i and j are identification numbers of the elements.

From Figure 4, equation of motion for the element i is

$$m\ddot{\xi_{i}} + c\sum_{j=1}^{n_{i}} (\dot{\xi_{i}} - \dot{\xi_{j}}) + k_{n_{i} - n_{j}} \sum_{j=1}^{n_{i}} (|\xi_{i} - \xi_{j}| - l_{n_{i} - n_{j}}) u_{ji} + d\dot{\xi_{i}} + \epsilon \xi_{i} = 0$$

where, position of the element ξ_i is defined as a vector, and m is mass of the element, and c, $k_{n_i-n_j}$, $l_{n_i-n_j}$, d denote damper coefficient, spring coefficient, natural length of spring, and friction coefficient, respectively. n_i and n_j are the number of connection of elements i and j. u_{ji} is the unit vector which goes to element j from element i. We assume the additional force $\epsilon \xi_i$ to aggregate the elements to the origin. This is the force proportional to the distance between the element and the origin of the plane. This is added to increase the efficiency of convergence, but it is weak enough so that the formation is not affected.

3.2 Triangle formation

We simulated the formation process of a triangle. We changed a set of parameters such as, spring constant, natural length, and generating probability according to combination of connection number. For each set of parameters, formation process is simulated 1000 times, and the rate of success and an average step number are evaluated. Here, throughout the simulation we assumed m=10, r=5 (radius of an element

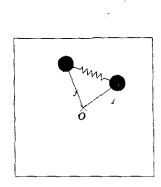


Figure 4: Movement of Element i

Table 1: Virtual springs for triangle formation

n_i	n_j	$k_{n_i-n_j}$	$l_{n_i-n_j}$	$P_{n_i-n_j}$
2	4			
4	2	!		}
4	4	200	90	1
4	6			
6	4			{
6	6			
$\overline{2}$	5	250	120	0.7
3	4	300	120	0.005
3	3	300	120	0.001
oth	iers	200	120	0.0001

body), d=2, c=2, $\epsilon=0.3$, R=100, and $|\xi_{init}|=2$ (initial velocity), where ξ_{init} has a random direction.

As a result, the best parameter set was selected for The number of elements n = 15 (see Table 1) and it was applied to various size of triangles (n = 6, 10, 21).

3.2.1 Parameter tuning for triangle formation

Combinations n_i - n_j contained in the target triangle are 2-4, 4-2, 4-4, 4-6, 6-4, and 6-6. Because these are structural members, we put nominal property $(k_{n_i-n_j} = 200, l_{n_i-n_j} = 90)$ for them. $P_{n_i-n_j}$ (generation probability of the springs) is set to 1.

These springs are not sufficient to control the global outline shape of the system. We need to add other kind of springs to reshape outline to the target shape. We introduced springs with a larger spring constant and larger natural length to cut a specific unnecessary connection (lower half of Table 1).

Frequency of breaking the connections that are not included in a triangle is also important.

If the spring with the combinations such as $n_i = 4, 5$,

Table 2: Simulation results by size of target

Average	4423	7447	7662	27508
Steps in worst case	31684	55774	42364	178253
Steps in best case	100	160	313	257
Success rate(%)	100	100	100	83.7
Size of target triangle	6	10	15	21

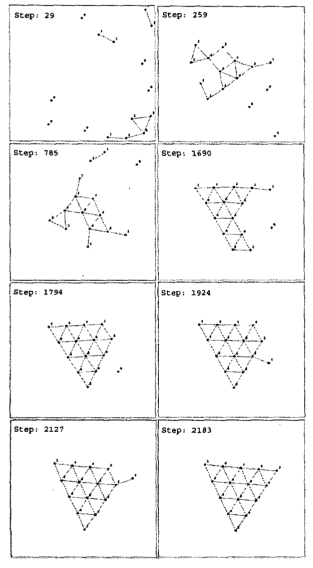


Figure 5: Simulated formation process of a triangle

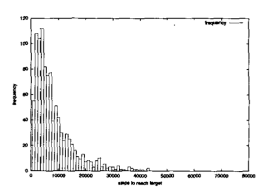


Figure 6: Histogram of steps to reach target (n = 15)

and 6, is generated for separation in high probability, it will never converge. On the contrary, if the probability is too low, then other formations easily appear. We found that moderate generation probability for 3-3 and 3-4 is efficient to form a triangle.

Deadlock shapes usually contain obtuse vertices (120°) , and the connection number of them is 3. We, therefore, cut connections including $n_i = 3$ in low probability. However, since $n_i = 3$ also appears during the desired formation process of a triangle, the probability must be small enough. Combination 2-5 has a particular function. It gives an element freedom of motion along the edge, instead of completely cutting it off. We set a very small probability to all the other springs not contained in the above list. This is effective to prevent deadlocks in general.

3.2.2 Simulation results

Simulation results are shown in Table 2 and Figure 5. The rate of convergence was evaluated by sizes of targets. We used the parameter set tuned for a triangle with 15 elements for all of them.

In Figure 6, the peak of the necessary step for convergence appeared in the left of the plot shows that it is basically a random walk process. It is a process without memory, because the number of activation for each rule is linearly increased against convergence time (Figure 7).

Since those simulations were obtained by the parameters for 15 elements, convergence for 6 and 10 elements were worse. It will be improved if parameters are tuned according to that target size. Although for the target larger than 15, we need longer convergence time, the rate of a success did not get worse so much, since the target shape is the only stable state.

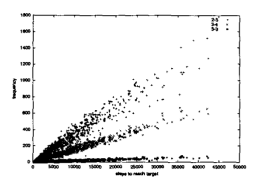


Figure 7: Frequency of separation by spring 2-5, 3-4, 3-3 (n = 15)

Table 3: The generating virtual springs for a ladder

n_i	n_j	$k_{n_i-n_j}$	$l_{n_i-n_j}$	$P_{n_i-n_j}$
$\overline{2}$	3			
3	2			
2	4			
4	2	200	90	1
3	4			
4	3			
4	4			
2	5	250	120	0.7
oth	ers	250	120	0.001

3.3 Formation of other shapes

We applied the same algorithm to other shapes such as a linear ladder and a hexagon.

For a ladder, we assume n=15, m=10, r=5, d=0, $c=2, \epsilon=0, R=100$, and for a hexagon n=19, m=10, $r=5, d=2, c=2, \epsilon=0.3, R=100$.

Combinations in the ladder are 2-3, 3-2, 2-4, 4-2, 3-4, 4-3, 4-4, and combinations in the hexagon are 3-6, 6-3, 4-6, 6-4, 6-6. $k_{n_i-n_j}$, $l_{n_i-n_j}$ and $P_{n_i-n_j}$ were set to the same value as 3.2. We designed other springs as shown in Tables 3 and 4.

By preliminary simulation, we succeeded in forming these shapes as is shown in Figures 8 and 9. However, the tuning of the parameters was difficult, and the convergence was much slower than the case of triangle. In order to improve the efficiency, we need to add some springs to remove deadlock states.

Table 4: The generating virtual springs for a hexagon

n_i	n_j	$k_{n_i-n_j}$	$l_{n_i-n_j}$	$P_{n_i-n_j}$
3	6			
6	3	,		
4	6	200	90	1
6	4			
6	6			
2	4	250	120	0.7
2	5	250	120	0.7
3	3	300	120	0.0005
oth	ers	300	120	0.0001

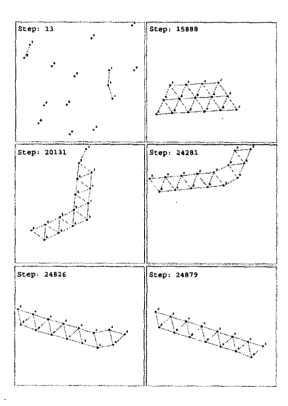


Figure 8: Formation process of a ladder

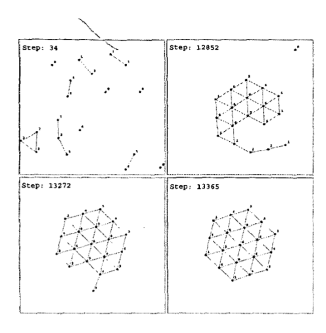


Figure 9: Formation process of a hexagon

4 Conclusion

In this paper, under the constraint that the elements locally interact with neighbors within radius R, we proposed an algorithm to form a shape much larger than R by using virtual springs, where, the properties of the spring were designed based on the number of connection of the element.

. In the proposed algorithm, it is necessary to tune parameters for the specific shape and the specific number of elements. In order to obtain the optimal parameter automatically, we can use genetic algorithms for instance. However, since it is a random process that does not have memory, it is difficult to drastically improve convergence speed. To extend this algorithm we need to introduce some additional properties such as nonlinear characteristics of the virtual springs and internal state of the element. For instance, by introducing the state, it becomes a system that has memory. By using the memory, we can not only improve convergence but assemble hierarchical structures, and more complicated shapes will be formed by such an extension.

As a future work, we need to prove that various formations are possible by this algorithm that does not exist in natural crystals. We also plan to redesign the algorithm in simpler form aiming at hardware embodiment by a cluster of small mobile robots. The simplicity of the formation algorithm may also lead us to applications such as nano/molecular machines.

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