Competing associations in six-species predator-prey models

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Abstract. We study a set of six-species ecological models where each species has two predators and two prey. On a square lattice the time evolution is governed by iterated invasions between the neighboring predator-prey pairs chosen at random and by a site exchange with a probability $X_s$ between the neutral pairs. These models involve the possibility of spontaneous formation of different defensive alliances whose members protect each other from the external invaders. The Monte Carlo simulations show a surprisingly rich variety of the stable spatial distributions of species and subsequent phase transitions when tuning the control parameter $X_s$. These very simple models are able to demonstrate that the competition between these associations influences their composition. Sometimes the dominant association is developed via a domain growth. In other cases larger and larger invasion processes precede the prevalence of one of the stable associations. Under some conditions the survival of all the species can be maintained by the cyclic dominance occurring between these associations.

1. Introduction

Ecological systems exhibit a large number of associations of certain species which coexist on the same territory. These associations can be considered as evolutionary stable states (or Nash equilibria) whose stability and competitiveness depend on the environmental conditions [1, 2, 3, 4, 5, 6]. Generally, the systematic analysis of the competition between these spatial states is made difficult by the complexity of the phenomena and by the large number of parameters in the suitable models.

In order to have a deeper insight into these phenomena we consider now a family of toy models exemplifying some basic features and the richness of possible behaviors. In these predator-prey models the species, located on the sites of a lattice, invades one of the neighboring sites if it is occupied by its prey. The investigation of predator-prey systems on lattices has a long history reviewed recently in the papers [5, 7, 8]. In these systems the invasion processes can yield extinction of species and/or maintain some steady states in the spatial distribution if the corresponding food web (a directed graph) contains loops. In such a situation a self-organizing spatio-temporal pattern can be formed by the species dominating cyclically each other. Besides it, in the present model the neutral (neighboring) species are allowed to exchange their sites. This latter
process helps the formation of many well-mixed distributions of neutral species which
can also protect each other against the rest of species. As a result these models possess
many final states toward which such a system can evolve. In the spatial systems these
states can coexist by forming domains if the system is started from a random initial state
for sufficiently large sizes. The competition between these multi-species associations
along the boundaries can affect their internal structure and the evolutionary process
selects one of these states. Thus, the investigation of these simplified models can help
us to clarify some mechanisms related to Darwinian selection \[1\] and the emergence of
structural complexity \[9\] characterizing pre-biotic evolution too \[10\].

Several basic features are already well known for the simplest versions of the
above models. As mentioned above, if the spatial predator-prey model on the square
lattice contains three species with cyclic dominance (this model is also called as spatial
rock-scissors-paper game) then the cyclic invasions maintain a self-organizing pattern
with equal species concentrations \[11, 12, 13\]. Both the composition \[14, 15\] and the
geometrical features \[16, 17\] of this spatial pattern are affected by the independent
variation of invasion rates meanwhile this structure provides a stability against some
external invaders \[18, 19\]. Similar self-organizing pattern is found for those cyclic
 predator-prey (or voter) models where the number \(N_s\) of species is limited, namely
\(3 < N_s \leq 14\) \[13\]. For even \(N_s\), however, these systems are very sensitive to the
independent variation of invasion rates \[20\]. More complex behavior was observed for
those systems where the food web consists of several cycles that can be distinguished
from a topological point of view \[19\]. In these multi-cycle ecological models the final
(stationary) state is prevailed by a defensive alliance being the most stable cyclic
subsystem. These defensive alliances are composed of those species that form a cycle
in the food web thereby their spatial distribution is equivalent to those of one-cycle
systems mentioned above. The privileged role of these associations is related to their
most effective protection, provided by the topological features of the food web, against
the external species. In their enhanced protection the role of spatial structure is crucial
because under mean-field conditions these associations are not able to eliminate the
external invaders.

Very recently the investigation of a four-species cyclic predator-prey model has
indicated the spontaneous formation of another type of defensive alliances \[21\]. In this
latter model the lattice sites can be empty and the individuals are allowed to jump to
one of the neighboring empty sites. As a result the odd (even) label species can form
a well-mixed state ensuring the possibility of prompt counter-attack against the even
(odd) label species and by this means these associations can defend their territories.
In other words, the neutral species can also form defensive alliances in the presence of
some local mixing.

In the present paper we consider a set of models allowing the appearance of both
types of above mentioned defensive alliances. In these six-species models the food web
contains many cycles and the local mixing is provided by a site exchange between
the neutral pairs. For all the models the probability \(X_s\) of site exchange will be the
only tunable parameter. Using Monte Carlo (MC) simulations it will be shown that, despite their simplicity, these models are able to account different stationary states and subsequent phase transitions when varying the value of $X_s$. The stationary states are composed of different species whose number varies from 2 to 6. Evidently, the time scales of the formation of these states are dependent on the microscopic mechanism (invasion or mixing). Sometimes equivalent states are formed via a nucleation mechanism and they can destroy each other (cyclically) when the boundaries of these growing domains meet.

Apparently the combination of our previous models \[19, 21\] means a simple extension of the six-species models. This extension, however, exemplifies that a small variation of a sufficiently complex ecological model can yield surprisingly large increase in the types of behaviors and phase transitions whose clarification raises many other questions. At the same time, many aspects of these features seem to be valid for the sufficiently complex ecological systems (or evolutionary games) as discussed in the Conclusions.

2. Spatial predator-prey models

In all the present models the site $i$ of a square lattice is occupied by an individual belonging to one of the six species and the corresponding distribution is given by a set of site variables ($s_i = 0, \ldots, 5$). The predator-prey relation between the species is defined by a food web. On the corresponding directed graph the nodes (labelled from 0 to 5) refer to the species and the directed edges point to the prey from its predator. Our analysis is focused on those systems where each species has two predators, two prey, and a neutral one (not linked). In this case each possible food web (satisfying the above conditions) is isomorphic to one of the four graphs shown in Fig. 1 and the corresponding model is named after their labels A, B, C, and D [19].

The evolution of the species distribution is governed by iterating nearest neighbor invasions and site exchanges. More precisely, starting from a random initial state we repeat the following steps: 1) two nearest-neighbor sites are chosen at random; 2) if these sites are occupied by a predator-prey pair then the prey is killed and an offspring of the predator occupies the site of prey; 3) for neutral pairs the species exchange their sites with a probability $X_s$. Evidently, nothing happens if the individuals belong to the same species. For example, in the model A the randomly chosen pair (1, 2) transforms into (1, 1) (invasion) and the pair (1, 4) transform to (4, 1) (site exchange) with a probability $X_s$, otherwise the neutral pair remains unchanged. Notice that the invasion rates between the predator-prey pairs are unique and the models have only one parameter ($X_s$) characterizing the strength of mixing between the neutral species.

These models are investigated by a series of MC simulations performed on a square lattice with sites $N = L \times L$ under periodic boundary conditions at different values of $X_s$. The simulations are started from a random initial state. Within a time unit (MCS) each pair has a chance once on the average to perform an invasion (between predator
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Figure 1. Food webs defining the four classes of predator-prey relations where each species has two predators and two prey. The decorated lines connect the members of the defensive alliances providing mutual protection against the external invaders via cyclic invasions.

and prey) or a site exchange with a probability $X_s$ (between neutral species). During the simulations we have recorded the current values of species concentrations $[\rho_\alpha(t); \alpha = 0, \ldots, 5]$ as well as the probability of finding predator-prey $[P_{pp}(t)]$ and neutral pairs $P_n(t)$ on two neighboring sites. The stationary states are characterized by the averages values of these quantities (denoted as $\rho_\alpha$, $P_{pp}$, and $P_n$) which are determined by averaging over a long sampling time varied from $10^4$ to $10^6$ MCS after a suitable thermalization time. Evidently, $\sum_\alpha \rho_\alpha = 1$ and the quantity $1 - P_{pp} - P_n$ gives the probability of finding the same species on two neighboring sites. At the same time we have determined the fluctuation of these quantities defined as $\chi_\alpha = N(\langle \rho^2_\alpha \rangle - \langle \rho_\alpha \rangle^2)$ where $\langle \ldots \rangle$ refers to the averaging over the sampling time. To avoid undesired size effects the linear size is varied from $L = 500$ to $2500$. The larger sizes are used in the close vicinity of transition points where the fluctuations diverge.

Now we discuss the general features of the possible stationary states. For all these models there exist six homogeneous states denoted as $\Phi^{(h)}(i)$ where $s_i = \alpha$ ($\alpha = 0, \ldots, 5$) for each site $i$. These states remain unchanged and the evolution of the species distribution is stopped whenever one of these (absorbing) states is reached. At the same time, the homogeneous state is unstable against the invasion of the suitable predators.

The mixed states of neutral species are denoted as $\Phi^{(n)}$ if $s_i = 0$ or 3. Evidently there exist two additional neutral pairs for all the four models. In such a state there are no invasions at all. Consequently, the ratio of populations is fixed during the evolution meanwhile the system tends towards an uncorrelated spatial distribution for $X_s > 0$. 
For all the four models there exist several three-species associations whose members invade cyclically each other. For example, in model A one of the corresponding states is denoted as $\Phi_{024}^{(c)}$ where the subscripts refer to the three species being present with the same average concentrations, $\rho_0 = \rho_2 = \rho_4 = 1/3$. The corresponding spatial distribution is maintained by the cyclic invasions yielding a short average life time for the individuals (the survival probability of individuals decreases exponentially with a relaxation time $\tau \simeq 1.8$ MCS) and the correlation length ($\xi \simeq 2.5$) is characterizing the typical domain size $[11, 13, 22]$. In general, the stability of the possible three-species associations against the external invaders is determined by the food web topology $[19]$. For the model A the food web contains only two equivalent three-species cycles which are called defensive alliances because their members protect cyclically each other against the external invaders. For example, within the state $\Phi_{024}^{(c)}$ the species 2 can only be invaded by the “external” species 1. During the cyclic invasion processes both the internal species 2 and the substituted invader 1 are killed by their common predator (0), in the meantime the third member (4) of the association feeds the species 2 and blocks the spreading of invaders. Thus, sooner or later the offspring of invaders will be eliminated from the system. One can easily check that other types (3 or 5) of invaders become extinct in the same way because of the cyclic symmetry in this food web. For the food web B the system has only one cyclic defensive alliance ($\Phi_{135}^{(c)}$) that will dominate the final stationary state although this food web has four additional three-species cycles with lower stability. Model C has eight equivalent three-species cycles, and neither of them can be considered as a defensive alliance in the sense mentioned above.

Model D represents another situation where one can find four three-species and three four-species cycles. In this case the four-species cycle $\Phi_{0234}^{(c)}$ proved to be the most stable spatial association that satisfies the conditions to be a cyclic defensive alliance $[19]$. For $X_s = 0$, the stability of the different states is already determined in a previous study when considering the effect of mutations on the population in similar six-species predator-prey models $[19]$. The present analysis will be focused on the effect of site exchange between the neutral pairs ($X_s > 0$) for the absence of mutation. For small sizes all the above mentioned states can be observed as a final state. For sufficiently large system sizes, however, the system tends towards the real stationary state independent of the initial state.

3. Four-species subsystems

In this section we study the behavior of several four-species subsystems whose understanding will help the interpretation of results obtained for the more complicated systems. More precisely, our analysis is focused on three systems (labelled as a, b, and c) whose food web are given in Fig. 2.

The food web of subsystem a is equivalent to a four-species cyclic predator-prey model where each species has only one predator and one prey. Such a situation can
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Figure 2. Food webs for the four-species subsystems.

occur in the models $A$, $B$, and $D$ if two species are missing, and its analysis becomes particularly important for model $D$ where $\Phi^{(c)}_{0234}$ is the stable state for $X_s = 0$. In addition, similar cyclic dominance has been reported very recently by Traulsen et al.\cite{23,24} considering a four-strategy prisoner’s dilemma game.

Some features of this system is already investigated previously by several authors for $X_s = 0$ \cite{13,20,21}. In this case the cyclic invasions maintain a self-organizing pattern with $\rho_0 = \rho_1 = \rho_2 = \rho_3 = 1/4$ and with a probability $P_n = 0.0518(5)$ to find neutral pairs on two neighboring sites. Here $P_n$ characterize the concentration of interfaces separating neutral domains (e.g., $\Phi_0^{(b)}$ and $\Phi_2^{(b)}$). Along these interfaces the evolution is blocked until one of the invading predators reaches the interface. For $X_s > 0$, however, along these neutral interfaces the site exchange increases the value of $P_n$ and supports the formation of the well-mixed phase of neutral species.

The MC simulations indicate clearly how the value of $P_n$ increases monotonously with $X_s$ until a threshold value $[X_s < X_{cr}^{(4a)} = 0.02662(2)]$. At the same time the probability $P_{pp}$ of predator-prey pairs is decreasing as shown in Fig. 3.

Figure 3. Probability of neutral (pluses) and predator-prey (open diamonds) pairs as a function of $X_s$ for the four-species cyclic food web (see a in Fig. 2).

Above this threshold value ($X_s > X_{cr}^{(4a)}$) the system segregates into two types ($\Phi_{02}^{(a)}$ and $\Phi_{13}^{(a)}$) of growing domains which are mixtures of neutral pairs. During the domain
growing processes these phases are in contact throughout a wide boundary layer (of type \( \Phi_{0123}^{(c)} \)) which serves as a species reservoir and sets the compositions to be symmetric [21]. Finally, one of the two equivalent mixed states will prevail the whole finite system and then \( P_{pp} = 0 \) and \( P_n = 0.5 \).

Subsystem \( b \) represents the situation when the three-species cyclic defensive alliance \( (\Phi_{123}^{(c)}) \) is attacked by an external invader (of type 0). The MC simulations indicate clearly that after some relaxation process the species 0 dies out and the system remains in the state \( \Phi_{123}^{(c)} \) if \( X_s < X_{cr}^{(4b)} = 0.0527(1) \). Conversely, the system develops into the symmetric mixed phase \( \Phi_{02}^{(n)} \) (\( \rho_0 = \rho_2 = 0.5 \)) for \( X_s > X_{cr}^{(4b)} \). In the vicinity of the transition point the visualization of the species distribution illustrates the formation of domains of \( \Phi_{02}^{(n)} \) whose area increases (decreases) slowly above (below) the threshold value. The average velocity of the interfaces (separating the domains of \( \Phi_{02}^{(n)} \) and \( \Phi_{123}^{(c)} \)) seems to be proportional to \( X_s - X_{cr}^{(4b)} \) as it is observed for another model [21]. This feature can explain why the ordering process becomes so slow in the close vicinity of the transition point.

The four-species subsystem \( c \) possesses two equivalent, three-species cycles that have two common species (1 and 3). In this case the spatial distribution of species segregates into two types (\( \Phi_{123}^{(c)} \) and \( \Phi_{103}^{(c)} \)) of growing domains and finally the whole system is dominated by one of them with equal probabilities. Notice that the bulk of these phases are not influenced by the site exchange mechanism whose effect is limited to the boundaries separating the phases \( \Phi_{123}^{(c)} \) and \( \Phi_{103}^{(c)} \). Furthermore the species within the mixed two-species phase \( \Phi_{02}^{(n)} \) are not able to protect each other because species 1 can invade their territories without restraint. This is the reason why the domain growing process is not prevented for \( X_s > 0 \) and the site exchange causes only a slight variation in the velocity of growing process.

4. Model A

As mentioned above the six-species model \( A \) has two equivalent three-species cycles consisting of disjoint sets of species, namely \( \Phi_{024}^{(c)} \) and \( \Phi_{135}^{(c)} \). According to the simulations one of these phases (with equal probability) will overwhelm the whole finite system after a domain growing process if the site exchange probability does not exceeds a threshold value, i.e., \( X_s < X_{cr}^{(6A)} = 0.05592(1) \). On a sufficiently large scale this domain growing process is similar to those that one can observe for the kinetic Ising model below the critical temperature if the system is started from a random initial state (for a survey of domain growth see [25]).

Above the threshold value \( (X_s > X_{cr}^{(6A)}) \) the system develops into one of the three (equivalent) two-species states composed from neutral species. The topological features of the food web \( A \) implies that the phases \( \Phi_{03}^{(n)} \), \( \Phi_{14}^{(n)} \), and \( \Phi_{25}^{(n)} \) can also be considered as defensive alliances because their members protect each other against the external predators in the well-mixed state. During the transient time one can observe a domain growth with these three types of domains. Apparently this growth is similar to
those observed in the three-state Potts model below the critical temperatures \([26, 27]\). However, along the interfaces separating two domains (say \(\Phi_{03}^{(n)}\) and \(\Phi_{14}^{(n)}\)) there occur a four-species cycle \(\Phi_{0134}^{(c)}\) whose transversal extension is limited by invasions from both sides. Notice that the corresponding four species subsystem (see the food web \(a\) in Fig. 2) exhibit a transition from the four-species cycle \(\Phi_{0134}^{(c)}\) in the mentioned example) to one of the mentioned neutral pair states at \(X_s = X_{cr}^{(4a)}\). As \(X_{cr}^{(6A)} > X_{cr}^{(4a)}\) therefore the coexistence of the four species is limited to the narrow strips separating the well-mixed phases of the neutral pairs.

The three-color maps have special points (called three-edge vertices) where three domains meet. Instead of it, here there are patches with phases of either \(\Phi_{024}^{(c)}\) or \(\Phi_{135}^{(c)}\). This inevitable coexistence of many different phases (during the decomposition processes) may be the reason why \(X_{cr}^{(6A)} > X_{cr}^{(4a)}\).

5. Model B

In this model a very interesting behavior is found by the MC simulations. One can observe four different phases and three subsequent phase transitions when \(X_s\) is increased. It is more surprising that the transition points are very close to each other. To demonstrate the main features Fig. 4 illustrates the average values of species concentrations in the stationary states reached by the system starting from a random initial state for different values of \(X_s\). Here the open (closed) symbols are used for the species with odd (even) labels to reduce the confusion coming from the coincidence of several data. Furthermore, the neutral pairs are denoted by the same type of symbols.

![Figure 4. Concentration of species vs. \(X_s\). The MC data are denoted by closed circles, open diamonds, closed squares, open circles, closed diamonds, and open squares for the species labelled from 0 to 5. The arrows indicate the three transition points.](image)

The cyclic defensive alliance \(\Phi_{135}^{(c)}\) is proved to be the most stable solution if \(X_s < X_{c1}^{(6B)} = 0.06155(5)\). In the next region \([X_{c1}^{(6B)} < X_s < X_{c2}^{(6B)} = 0.06545(5)]\)
the final stationary state will be equivalent to one of the three well-mixed states of neutral pairs \( \Phi_{03}^{(n)}, \Phi_{14}^{(n)}, \text{and} \Phi_{25}^{(n)} \). All the six species survive in the third region of \( X_s \) in such a way that \( \rho_0 = \rho_2 = \rho_4 \) and \( \rho_1 = \rho_3 = \rho_5 \). If \( X_s > X_{c3}^{(6B)} = 0.07372(2) \) then only the species with even labels are present and their coexistence with the same concentration is maintained by the cyclic invasions (state \( \Phi_{024}^{(c)} \)).

The transitions at \( X_{c1} \) and \( X_{c2} \) can be considered as a first order transition meanwhile the third one exhibits the features of the continuous (critical) transitions. The concentration of the species 1, 3, and 5 vanishes continuously when approaching \( X_{c3} \) from below. More precisely, our data are consistent with a power law decrease in the close vicinity of the transition point, that is \( \rho_1 = \rho_3 = \rho_5 \simeq (X_{c3} - X_s)^\beta \) with \( \beta = 0.37(4) \). Similar power law behavior (with the same exponent) is found for \( P_n \) as shown in Fig. 5. At the same time the MC data show diverging fluctuations for these quantities that is another relevant feature of the critical transitions. Unfortunately, we could not deduce an adequate \( (\gamma) \) exponent to describe this divergence because of the large statistical uncertainties in these quantities.

![Figure 5](image.png)

Figure 5. Log-log plot of the MC data (open squares) for the probability of neutral pairs vs. \( X_{c3}^{(6B)} - X_s \). The fitted power law is denoted by the straight line with a slope of 0.373.

In the presence of absorbing state(s) the extinction of a single species in the spatial models has very robust features and the corresponding transition belongs to the directed percolation (DP) universality class characterized by a higher exponent \( (\beta_{DP} = 0.58(2) \) for the two-dimensional systems) [28, 29, 30, 31]. In the present model, however, three species die out simultaneously and the background can mediate some interactions among the vanishing species. Evidently, further analysis is necessary to quantify the features of this transition as well as to clarify the reasons causing the distinct behavior.

In contrary to the model \( A \) here the formation of one the three equivalent well-mixed phases of neutral species is limited to a narrow region \( [X_{c1}^{(6B)} < X_s < X_{c2}^{(6B)} = 0.06545(5)] \). The formation of these structures differs significantly from those described
for model $A$ because the food web $B$ does not provide mutual protection for a neutral pair against another neutral pair. Instead, the well-mixed phases of the neutral pairs destroy cyclically each other ($\Phi_{03}^{(n)}$ destroys $\Phi_{14}^{(n)}$ destroys $\Phi_{25}^{(n)}$ destroys $\Phi_{03}^{(n)}$) via a complicated process. For example, the offspring of a single species 0 can occupy the whole territory of the phase $\Phi_{14}^{(c)}$. As a result, if the growing domains of the phases $\Phi_{03}^{(c)}$ and $\Phi_{14}^{(c)}$ meet along a common boundary then the domain $\Phi_{14}^{(c)}$ is transformed (via the invasion) into a domain of $\Phi_{0}^{(h)}$ that is also unstable against the invasions of species 2 and 5.

Within the second region of $X_s$ the six species can coexist for a sufficiently long time in such a structure where the species with even labels are in minority. The formation of the mixed phases of the neutral species begins in those patches where two species of minorities are missing due to the fluctuations. According to the previous analysis of the four-species subsystem $b$ the spontaneous formation of the well-mixed phase of the corresponding neutral species is permitted because the condition $X_s > X_{cr}^{(db)}$ is fulfilled within the given region of $X_s$. By this means sufficiently large domains of the states $\Phi_{03}^{(n)}$, $\Phi_{14}^{(n)}$, and $\Phi_{25}^{(n)}$ happen to form by chance. During their expansion these domains meet one of their rivals and then one of them is destroyed as described above.

![Figure 6](image.png)

**Figure 6.** Time dependence of the probability of neutral (dashed line) and predator-prey (solid line) pairs for $X_s = 0.0654$ and $L = 2000$. The MC data are smoothed out by averaging over 2000 MCS.

Figure 6 shows the variation of $P_n(t)$ and $P_{pp}(t)$ for a given $X_s$ close to the second transition point. At the beginning these curves are smooth. Later, however, one can observe larger and larger variations reflecting the destruction of domains with types $\Phi_{03}^{(n)}$, $\Phi_{14}^{(n)}$, and $\Phi_{25}^{(n)}$ whose average size increases monotonously. The series of these larger and larger invasion processes is ended when only one of these phases prevail the whole system.

It is conjectured that this (rock-scissors-paper like) cyclic dominance between the two-species associations plays a crucial role in the maintenance of the six-species phase.
6. Model C

This model has eight equivalent three-species cyclic associations as mentioned above. The simulations for a system size of $L = 400$ have justified that any of them can be the final stationary state independently of the value of $X_s$.

It is expected that this ”mono-domain” state is developed via a domain growing process. This phenomenon can be described by the variation of the probability of neutral pairs vanishing in the bulk of ”mono-domain” states. The results of MC simulations (for $L = 2500$) are plotted in Fig. 7 where the data are averaged over a sampling time increased from 2 to 2000 MCS.

![Figure 7](image-url)

Figure 7. Log-log plot of the probability of neutral pairs vs. time for $X_s = 0.05$ (upper solid line) and $X_s = 0$ (central solid line). The lower solid line shows the same quantity in the model A for $X_s = 0$. To compare the MC results with the theoretical expectations, the upper dashed line represents the function $P_n(t) = 0.12/\ln(t)$ while the lower one shows an algebraic decay, $P_n(t) = 0.25/\sqrt{t}$.

Figure 7 shows that the very slow domain growing process for $X_s = 0$ becomes even slower for $X_s = 0.05$. Indeed we cannot exclude the blocking of coarsening process as it happens in the voter model for higher dimensions ($d > 2$) [32]. Keeping in mind the large statistical uncertainties (for large times) these data remind us to the growing processes observed for the two-dimensional voter model where the concentration of domain walls vanishes as $1/\ln(t)$ plotted also in Fig. 7 [33, 32]. The voter model exemplifies the coarsening process in a broad class of models undergoing a phase ordering without surface tension [34]. In the present model the definition of the boundary separating two cyclic three-species phases is confusing because these phases may involve one or two common species (compare $\Phi_{015}^{(c)}$ with $\Phi_{012}^{(c)}$ or $\Phi_{123}^{(c)}$).

Strikingly different growth process is found for model A (discussed previously) where the system has only two cyclic three-species phases composed from disjoint sets.
of species (Φ_{024}^{(c)} or Φ_{135}^{(c)}). In order to visualize the relevant differences Fig. 7 shows also the MC results obtained in the model A for X_s = 0 and L = 4000. These numerical results indicate that here the asymptotic behavior of \( P_n(t) \) is close to the theoretical prediction (\( P_n(t) \sim 1/\sqrt{t} \)) characterizing the growth controlled by the reduction of an interfacial energy \[25\] \[27\].

Evidently, further research is required to clarify those ingredients of these models that are responsible for the different growth processes. Furthermore, here it is worth mentioning that the logarithmic growth supports the maintenance of biodiversity for a very long time if the system is sufficiently large.

7. Model D

For this food web the system has a four-species cyclic defensive alliance (Φ_{0234}^{(c)} as shown in Fig. 1) that is stable if \( X_s = 0 \) \[19\]. According to the simulations for weak rate of site exchange the species 1 and 5 die out within a short transient time. In the corresponding four-species subsystem (see Sect. 3) the cyclic invasions can sustain their coexistence if \( X_s < X_{cr}^{(4a)} \). For higher \( X_s \) the spatial distribution of these four species decomposes into two types of two-species domains (Φ_{03}^{(n)} and Φ_{24}^{(n)}) as described above. If \( X_s \) is increased then this process becomes faster, and above a threshold value the domains of phases Φ_{03}^{(n)} and Φ_{24}^{(n)} are built up by these processes before their mortal enemy (species 1 and 5) would die out. One can easily check that the species 1 (5) can occupy the whole territory of the phases Φ_{03}^{(n)} (Φ_{24}^{(n)}). Thus, for a sufficiently intensive mixing the species 1 and 5 will have an enhanced chance to survive because they are fed by the emerging domains of Φ_{03}^{(n)} and Φ_{24}^{(n)}. For large \( X_s \) this mechanism can be so effective that finally the phase Φ_{15}^{(n)} will dominate the whole system as shown in Fig. 8. Notice that the well mixed phase Φ_{15}^{(n)} is also a defensive alliance because species 1 and 5 protect each other against the different invaders.

Figure 8 shows that four types of final stationary states can be distinguished when \( X_s \) is increased. In the first region (\( X_s < X_{c1}^{(6D)} = X_{cr}^{(4a)} \)) the species of the phase Φ_{0234}^{(c)} take place with the same concentration meanwhile \( P_n \) and \( P_{pp} \) vary with \( X_s \) as plotted in Fig. 8. Starting from a random initial state this system evolves into one of the neutral-pair phases Φ_{03}^{(n)} and Φ_{24}^{(n)} if \( X_{c1}^{(6D)} < X_s < X_{c2}^{(6D)} \) where the value of \( X_{c2}^{(6D)} \) depends on the size \( L \) as it will be discussed below. Within the third region (\( X_{c2}^{(6D)} < X_s < X_{c3}^{(6D)} \)) the six species coexist in such a way that \( \rho_0 = \rho_2 = \rho_3 = \rho_4 > \rho_1 = \rho_5 > 0 \). If \( X_s > X_{c3}^{(6D)} = 0.0581(1) \) then only the species 1 and 5 will survive with forming a phase Φ_{15}^{(n)}.

In this model the first and the third transitions exhibit a sudden change both in concentrations and in pair probabilities (\( P_n \) and \( P_{pp} \)). However, the classification of the second transition (as well as the numerical value of \( X_{c2}^{(6D)} \)) is difficult because of its unusual features. Namely, \( \rho_1 \) and \( \rho_5 \) decrease monotonously when \( X_s \) is decreased within the third region. At the same time we have observed a dramatic increase in the fluctuation of concentration for the majority species (see Fig. 9) meanwhile
the corresponding average concentrations are close to 1/4 (see Fig. 8). To avoid the undesired effects of these fluctuations the simulations were performed for such a large system sizes (the linear size is increased up to $L = 2600$) where the current value of concentrations were significantly larger than the amplitude of fluctuation characterized by their standard deviation dependent on size ($\sqrt{\chi/N}$). For smaller sizes the system evolves into the phase $\Phi_{03}^{(n)}$ or $\Phi_{24}^{(n)}$.

Figure 8. Concentration of species vs. $X_s$ in the model $D$. The MC data are denoted by symbols with fourfold symmetry (pluses, $X_s$, open squares, and diamonds) for those species belonging to four-species cyclic defensive alliance, and by open circles for the species 1 and 5. The arrows indicates the critical values of $X_s$ where transitions appear.

Figure 9 demonstrates clearly the absence of fluctuations for the missing species as
well as for the phases composed of neutral species. The MC data refer to an exponential increase in $\chi_0$, $\chi_2$, $\chi_3$, and $\chi_4$ if $X_s$ approaches $X_{c2}$ from above.

Within the region of large fluctuations the time dependence of concentrations shows the occurrence and destruction of domains with the structure $\Phi^{(n)}_0$ and $\Phi^{(n)}_2$. These domains are invaded by the sparsely dispersed species 1 and 5. The resultant phase, however, is not stable at the given $X_s$ and sooner or later the phases $\Phi^{(n)}_0$ or $\Phi^{(n)}_2$ are formed again. In some sense the evolution of the spatio-temporal pattern is similar to those described by the mentioned spatial rock-scissors-paper games [11, 13, 17] as well as in the forest-fire models [35, 36, 37]. In the present case two cycles interfere in a complex way because cyclic dominance emerges between the different set of associations and this mechanism supports the maintenance of biodiversity with more associations (and species).

Due to the mentioned difficulties we couldn’t determine accurately how concentrations $\rho_1$ and $\rho_5$ vanish when approaching $X_{c2}^{(6D)}$ from above. In the light of the MC results one can conclude a weakly first order phase transition (sudden jump in $\rho_1 = \rho_5$) accompanied with an increase in $\chi_1 = \chi_5$ as illustrated in Fig. 9.

8. Conclusions

The numerical analysis of the four different six-species, spatial predator-prey models has highlighted the large variety of states and phenomena (transitions) that can occur in self-sustaining, multi-species ecological systems. Due to their simplicity the present models have allowed us to consider quantitatively some interesting phenomena. Instead of recalling the curious features found for these oversimplified models henceforth I wish to concentrate on those general messages concluded from these examples.

One of the relevant messages is related to the increase in the degree of freedom. Many recent works are focused on spatial systems with one, two, or three species. For larger number of species the present models exemplify how the simple dynamical rules build up more complex objects (alliances) that can be considered as additional species participating in the evolutionary competition. Shortly, the degree of freedom (here the number of species) is increased by the spontaneously occurring alliances having particular composition and spatial structure. One can believe that the food web of these systems should be extended by these complex objects (with a suitable invasion rates) to have a better picture about the whole spatial system.

In the context of game theory (for details see the refs. [1, 2, 3, 4, 6]), the interaction of species is described by the payoff matrix (the adjacency matrix of the food web in the present models) that determines the Nash equilibrium from which the unilateral deviation yields loss. For many evolutionary (dynamical) rules the Nash equilibrium represents an evolutionary stable state under the mean-field conditions. The Nash equilibrium is not necessary unique and it can be a mixed state (strategy). In the spatial models with short range interactions several species can be absent from a given territory and the Nash equilibrium for the corresponding subsystem can be distinct from
those characterizing the complete system. The noises affect the stability of these spatial states \[38\] and modify the average velocity of invasions along the interfaces separating the competing phases \[21\]. The above results justify that the invasion processes between the “stable” states of subsystems can play a crucial role in the formation of the final stationary states.

For many cases the final stationary state is reached through a coarsening process where the alliances (degenerated Nash equilibria) form growing domains. The corresponding domain growth process is affected by the additional phases (and mechanisms) along interfaces. It is not yet known how these effects modify the growth process yielding significantly slower (logarithmical) domain growth.

Most of the (first order) transitions occur at such a value of \(X_{s}\) where the invasion rate between two dominant alliances vanishes as detailed previously \[21\]. Remarkable exceptions are the fourth transition in model \(B\) and the second one in model \(D\). In both cases more than one species die out simultaneously and their extinction is accompanied by an enhanced concentration fluctuation for the surviving species and the resultant spatio-temporal structures can mediate unusual interactions between the vanishing species. These transitions are distinct inherently and their classification requires further time-consuming numerical investigations.

The spontaneous formation of complex spatio-temporal structures is one of the most interesting predictions of these simple models because it sustains an abundance of species. Furthermore, this feature mixes the concept of the part and whole. At the same time the corresponding system can be interpreted as a set of objects that create objects in a way characterizing the living systems \[39\]. In the light of the present results one can easily imagine that the choice of more complex food webs can result in the emergence of more and more complex objects (e.g., alliances of alliances with particular spatio-temporal structures). Besides it these models imply the possibility that the evolution of food web by mutations \[40, 41, 42\] and the evolution of the spatial ecological system are strongly related to each other while the spatial inhomogeneity can be enhanced in the system.

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