Abstract—We create a novel optimisation technique inspired by natural ecosystems, where the optimisation works at two levels: a first optimisation, migration of genes which are distributed in a peer-to-peer network, operating continuously in time; this process feeds a second optimisation based on evolutionary computing that operates locally on single peers and is aimed at finding solutions to satisfy locally relevant constraints. We consider from the domain of computer science distributed evolutionary computing, with the relevant theory from the domain of theoretical biology, including the fields of evolutionary and ecological theory, the topological structure of ecosystems, and evolutionary processes within distributed environments. We then define ecosystem-oriented distributed evolutionary computing, imbied with the properties of self-organisation, scalability and sustainability from natural ecosystems, including a novel form of distributed evolutionary computing. Finally, we conclude with a discussion of the apparent compromises resulting from the hybrid model created, such as the network topology.

I. INTRODUCTION

In Biologically-Inspired Computing, one of the primary sources of inspiration from nature has been evolution [1]. Evolution has been clearly identified as the source of many diverse and creative solutions to problems in nature [2], [3]. However, it can also be useful as a problem-solving tool in artificial systems. Computer scientists and other theoreticians realised that the selection and mutation mechanisms that appear so effective in biological evolution could be abstracted to be implemented in a computational algorithm [1]. Evolutionary computing is now recognised as a sub-field of artificial intelligence (more particularly computational intelligence) that involves combinatorial optimisation problems [4].

The use of evolution for algorithm generation may be well understood within computer science under the auspices of evolutionary computing [5], but ecology is not. Ecosystems are considered to be robust, scalable architectures that can automatically solve complex, dynamic problems, possessing several properties that may be useful in automated systems. These properties include self-organisation, self-management, scalability, the ability to provide complex solutions, and the automated composition of these complex solutions [5]. So, potential exists to create an ecosystem-oriented form of evolutionary computing. We propose that an ecosystem inspired approach, would be more effective at greater scales than traditionally inspired approaches, because it would be built upon the scalable and self-organising properties of ecosystems [6].

In this paper, we give a brief introduction to the theory of distributed evolutionary computing (section II). We then introduce ecosystems, including relevant issues around networks and spacial dynamics (section III). We then describe what is required for ecosystem-oriented distributed evolution (section IV), before illustrating our model of ecosystem-oriented distributed evolutionary computing (section V). We then consider its emergent topology (section VI), before then discussing the apparent compromises resulting from the hybrid model created (section VII), and stating our conclusions (section VIII).

II. DISTRIBUTED EVOLUTIONARY COMPUTING

The fact that evolutionary computing manipulates a population of independent solutions actually makes it well suited for parallel and distributed computation architectures [7], i.e. evolution is well suited to the ecological spaces in which it occurs. The motivation for using parallel or distributed evolutionary algorithms is twofold: first, improving the speed of evolutionary processes by conducting concurrent evaluations of individuals in a population; second, improving the problem-solving process by overcoming difficulties that face traditional evolutionary algorithms, such as maintaining diversity to avoid premature convergence [8]. There are several variants of distributed evolutionary computing, with there being two main forms [7], [8]:

Fine-grained diffusion models [8] assign one individual per processor. A local neighbourhood topology is assumed, and individuals are allowed to mate only within their neighbourhood, called a deme [7]. The demes overlap by an amount that depends on their shape and size, and in this way create an implicit migration mechanism. Each processor runs an identical evolutionary algorithm which selects parents from the local neighbourhood, produces an offspring, and decides whether to replace the current individual with an offspring. However, even with the advent of multi-processor computers, multi-core processors, and large scale distributed computing (cloud computing) which provide the ability to execute

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multiple threads simultaneously, this approach would still prove impractical in supporting the number of populations necessary to create an ecosystem-orientated form of distributed evolutionary computing.

In the coarse-grained island models \[10, 7\], evolution occurs in multiple parallel sub-populations (islands), each running a local evolutionary algorithm, evolving independently with occasional migrations of highly fit individuals among sub-populations. An example island-model \[10, 7\] is visualised in Figure 1 in which there are different probabilities of going from island 1 to island 2, as there is of going from island 2 to island 1. This allows maximum flexibility for the migration process, and mirrors the naturally inspired quality that although two populations have the same physical separation, it may be easier to migrate in one direction than the other, i.e. fish migration is easier downstream than upstream. The migration of the island models is like the notion of migration in nature, being similar to the metapopulation models of theoretical ecology \[11\]. However, all the islands in this approach work on exactly the same problem, which makes it less analogous to ecosystems in which different locations can be environmentally different \[12\]. We will take advantage of this property later when defining ecosystem-orientated distributed evolutionary computing.

A. Other Work

While there have been past attempts to have ecologically-inspired forms of evolutionary computing, for example \[13, 14\], they have all lacked actual ecological dynamics. In the case of \[13\] they attempt to include a feature of population dynamics, which they incorrectly refer to as ecology-inspired. While we have previously considered \[15, 16, 17, 18, 19, 20\] the possibility of ecosystem-oriented distributed evolutionary computing (DEC), but only in very specific applications, so now we can consider it abstractly from which it can wide applicability.

III. Natural Ecosystems

An ecosystem is a natural unit made up of living (biotic) and non-living (abiotic) components, from whose interactions emerge a dynamically stable, self-perpetuating system. It is made up of one or more communities of organisms, consisting of species in their habitats, with their populations existing in their respective micro-habitats \[12\]. A community is a naturally occurring group of populations from different species that live together, and interact as a self-contained unit in the same habitat. A habitat is a distinct part of the environment \[12\], for example, a stream. Individual organisms migrate through the ecosystem into different habitats competing with other organisms for limited resources, with a population being the aggregate number of the individuals, of a particular species, inhabiting a specific habitat or micro-habitat \[12\]. A micro-habitat is a subdivision of a habitat that possesses its own unique properties, such as a micro-climate \[22\]. Evolution occurs to all living components of an ecosystem, with the evolutionary pressures varying from one population to the next depending on the environment that is the population’s habitat. A population, in its micro-habitat, comes to occupy a niche, which is the functional relationship of a population to the environment that it occupies. A niche results in the highly specialised adaptation of a population to its micro-habitat \[22\].

A. Networks and Spatial Dynamics

A key factor in the maintenance of diversity in ecosystems is spatial interactions, and several modelling systems have been used to represent these spatial interactions, including metapop-
Evolutionary computing is uniquely defined by the creation of multiple evolving populations in response to similar requests, whereas in the island-models of distributed evolutionary computing there are multiple evolving populations in response to only one request. So, different requests are evaluated on separate islands (populations), with their evolution accelerated by the sharing of solutions between the evolving populations (islands), because they are working to solve similar requests (problems). Furthermore, the islands (habitats) will be able to support multiple populations, like a natural ecosystem. This is shown in Figure 3 where the dashed yellow lines connecting the evolving populations indicate similarity in the requests being managed.

Our distributed evolutionary computing architecture favours the use of Pareto-sets for fitness determination, because Pareto optimisation for multi-objective problems is usually most effective with spatial distribution of the populations, as partial solutions (solutions to different niches) evolve in different parts of a distributed population (i.e. different populations in different habitats). By contrast, in a single population, individuals are always interacting with each other, via crossover, which does not allow for this type of specialisation.

It should be noted that our approach requires a sufficiently large user base, so that there can be communities within the user base, and therefore allow for similarity in the user requests. Assuming a user base of hundreds of users, then there would be hundreds of habitats, in which there will be potentially three or more times the number of populations at any one time. Then there will be thousands of genes (atomic algorithms) available to meet the requests for algorithms from the users. In such a scenario, there would be a sufficient number of users to find similarity within their requests, and therefore apply our novel form of distributed evolutionary computing.

V. Computational Model

A. Gene

The gene, $g$, represents the base unit of the evolutionary computing processes, in the same way that genes are the functional unit in evolution, and so are atomic (irreducible) algorithms. Collections of genes are functionally analogous to the organisms of ecosystems, including their behaviour of migration. While evolutionary computing will be used for combinatorial optimisation to evolve optimal set of genes, $\{g_1, ..., g_k\}$, in response to a user $u$ request $r$ for an algorithm. So, the genes $g$ will migrate through the habitat network $(H = \{h_1, ..., h_c\})$ adapting to find niches where they are useful in fulfilling other user requests for algorithms $r$. Therefore, interacting and evolving, adapting over time $t$ to the environment, and thereby serving the ever-changing requirements imposed by a user base $U = \{u_1, ..., u_v\}$.

The genes $g$ migrate through the interconnected habitats $h$ combining with one another in populations $P$ to meet user requests for algorithms $r$. The migration path from the current habitat $h$ is dependent on the connections $c$ between the habitats $h$, the migration probabilities $c = \{p(h_1), ..., p(h_c)\}$. The migration of a gene $g$ is initially triggered by deployment to its user’s habitat $h$, for distribution to other users who will potentially make use of the algorithm that the gene $g$
represents. So, upon deployment a gene \( g \) is copied to the gene-pool \( \text{g}_{\text{r}} \) of the user’s habitat \( h \), and from there the migration of the gene \( g \) occurs, which involves migrating (copying) the gene \( g \) probabilistically \( (p(h) > 0) \) to all the connected habitats \( h \). The gene \( g \) is copied rather than moved, because the gene \( g \) may also be of use to the providing user \( u \). The copying of a gene \( g \) to a connected habitat \( h \) depends on the associated migration probability. If the probability were one \( (p(h) = 1) \), then it would definitely be sent. When migration occurs, depending on the probabilities associated with the habitat \( h \) connections \( c \), an exact copy of the gene \( g \) is made at a connected habitat \( h \). The successful use of the migrated gene \( g \), in response to user requests for applications \( r \), will lead to further migration (distribution) and therefore availability of the gene \( g \) to other users \( u \).

So, the connections \( c \) joining the habitats \( h \) are reinforced by successful gene \( g \) migration. The success of the migration, the migration feedback, leads to the reinforcing and creation of migration links between the habitats \( c \), just as the failure of migration leads to the weakening and negating of migration links between the habitats \( h \). The success of migration is determined by the usage of the genes \( g \) at the habitats \( h \) to which they migrate. When a gene-set \( \{g_i, ..., g_j\} \) is found and used in responding to a user request, then the appropriate connection probabilities can be updated. If the gene-set \( \{g\} \) was fully or partly evolved elsewhere, then where the set or subsets were created needs to be passed on to the connection probabilities, because the value in a gene-set \( \{g\} \) is the unique combination it provides of the individual genes \( g \) contained within. So, it is necessary to manage the feedback to the connection probabilities for migrating gene-sets \( \{g_i, ..., g_j\} \), and not just the individual genes \( g \) contained within the sequence, including the partial use of a gene-set, \( \{g\} \), in a newly evolved one, \( \{g_i, ..., g_j\} \). Specifically, the mechanism for migration feedback needs to know the habitats \( h \) where migrating gene-sets \( \{g\} \) were created, to create new connections or reinforce existing connections to these habitats \( h \). The global effect of the gene \( g \) migration and migration feedback on the habitat network \( H \) is the clustering of habitats \( h \) around the communities present within the user base \( U \).

So, a gene’s life-cycle begins with deployment to its user’s habitat \( h \) for distribution within the habitat network \( H \). The gene \( g \) is then migrated to any habitats connected to their user’s habitat \( (p(h_i) > 0) \), to make it available in other habitats where it could potentially be useful. The gene \( g \) is then available to the local evolutionary optimisation, to be used in evolving the optimal gene-sets(s) in response to a user request \( r \). The optimal gene-set \( \{g_i, ..., g_j\} \) is then registered at the habitat \( h \), being stored in the habitat’s gene-pool \( \text{g}_{\text{p}} \). If a gene-set solution \( \{g\} \) is then executed, an attempt is made to migrate (copy) it to every other connected habitat \( h \), success depending on the probability associated with the connection. A gene \( g \) can also be deleted if after several successive user requests at a habitat \( h \) it remains unused; it will have a small number of escape migrations, in which it is not copied, but is randomly moved to another connected habitat \( h \). If the gene \( g \) fails to find a niche before running out of escape migrations, then it will be deleted.

In biology a gene-pool is all the genes in a population [22], but here it is the set of genes \( \{g_i, ..., g_j\} \) available at a habitat \( h \).
The escape range is the number of escape migrations available to a gene $g$ upon the risk of death (deletion). If a gene $g$ migrates to a habitat $h$ and is not used after several user requests $r$, then it will have the opportunity to migrate (move not copy) randomly to another connected habitat $h$. After this happens several times the gene $g$ will be deleted (die). The escape range will be dynamically responsive to the size of the habitat $h$ cluster that the gene $g$ exists within. This creates a dynamic time-to-live [26] for the genes $g$, such that those that are used more will live longer and distribute further than those that are used less.

B. Population

A population in a natural ecosystem is all the members of a species that occupy a particular area at a given time [22]. Our population, $P$, is also all the members of a species that occupy a particular area at a given time, like an island from the island-models of distributed evolutionary computing [10]. So, the population $P$ represents evolving populations in our ecosystem-oriented distributed evolutionary computing. The population $P$ is instantiated to evolve an optimal algorithm (set of genes $\{g_1, ..., g_j\}$) to an individual user request $r$, from the set of genes (gene-pool, $g_p = \{g_1, ..., g_j\}$) available at the habitat $h$ where it is instantiated. The solution is found by evolving the population of solutions $\{(g_1, ..., g_j), ..., (g_k, ..., g_j)\}$, with the evolutionary process of a population $P$ providing a combinatorial optimisation [35] of the genes $g$ available, when responding to a user request $r$. We shall assume that the genes are not mutated as they are atomic by definition. So, as in genetic algorithms [37], mutations will occur by switching genes $g$ in and out of the gene-sets $\{g\}$ that define individuals. Recombination (crossover) will occur by performing a crossing of two individuals (gene-sets). Therefore, an evolving population $P$ consists of the selection pressure $s$ defined by the user $u$ request $r$, initialised with the gene-pool $\{g\}$ of the habitat $h$ where it is instantiated. Then, the evolving population $P$ becomes a collection of gene-sets $\{g\}$ evolving to the selection pressure $s$ created from the user request $r$.

$$population = P(r, g_p, g_s) = \{(g_1, ..., g_j), ..., (g_k, ..., g_j)\}. \quad (1)$$

The evolutionary optimisation is accelerated in the following three ways: first, the habitat $h$ provides an optimal subset of the genes $g$ available globally $G$ in the ecosystem $E$, which is localised to the specific user $u$ it represents; second, making use of gene-sets, $\{g_1, ..., g_j\}$, previously evolved in response to the user’s earlier requests $r_e$; and third, taking advantage of relevant gene-sets evolved elsewhere in response to similar requests $r_s$ by other users. The population $P$ then proceeds to evolve the optimal gene-set(s) $\{g\}$ that fulfils the user request $r$, and as the genes $g$ are the base unit for evolution, it searches the available gene-set combination space $P(g_p)$. For an evolved gene-set $\{g\}$ that is executed (instantiated) by the user $u$, it then migrates to other peers (habitats $h$) becoming hosted where it is useful, to combine with other genes $g$ in other populations $P$ to assist in responding to other user requests for algorithms $r$.

There is a huge body of work and continuing research regarding theoretical approaches to evolutionary computing [5], including the extensive use of genetic algorithms for practical real-world problem solving [38]. Full use should make of the current state-of-the-art, and future developments in the area of evolutionary computing [39] in determining the optimal evolutionary computing approaches to have in the populations $P$ of our ecosystem-oriented distributed evolutionary computing $E$.

C. Habitat

The habitats $h$ are functionally analogous to the habitats of a biological ecosystem [22], providing a distributed environment for gene migration and evolution to occur, extending the island-model of distributed evolutionary computing [10] for the connectivity between habitats $h$. There will be a habitat $h$ for each user $u$, through which they will submit their requests $r$ for algorithms. So, the habitat $h$ represents a location at which a gene $g$ can move to, and so the collection of genes $g$ at the habitat $h$ defines the gene-pool $g_p$ with which to seed populations $P$ to solve user requests $r$ for algorithms. Supporting this functionality, habitats $h$ have the following core functions:

- Provide a subset of the genes $g$ available globally $G$, relevant to the user $u$ that it represents, the gene-pool $g_p$.
- Accelerate, via its gene-pool $g_p$, the populations $P$ instantiated to evolve optimal gene-sets $\{g_i, ..., g_j\}$ in response to user requests for algorithms $r$.
- Manage the inter-habitat connections for gene migration, $c = \{p(h_1), ..., p(h_i)\}$.

The collection of genes $g_p$ at each habitat $h$ (peer) will change over time, as the more successful genes $g$ spread through the habitat network, $H = \{h_1, ..., h_i\}$, of the ecosystem $E$, and as the less successful genes $g$ are deleted. Successive user requests $r$ over time $t$ to their dedicated habitats $h$ makes this process possible, because the continuous and varying user requests for applications, $R = \{r_1, ..., r_j\}$, provide a dynamic evolutionary pressure, $S = \{s_1, ..., s_j\}$, on the genes globally $G = \{g_1, ..., g_j\}$, which have to evolve to better satisfy those requests. So, the genes $g$ will recombine and evolve over time, constantly seeking to increase their effectiveness for the user base $U = \{u_1, ..., u_i\}$. Additionally, it also stores gene-sets, $g_s$ evolved from the habitat’s populations $P$, and gene-sets that migrate to the habitat $h$ from other users’ habitats, because they can potentially accelerate future populations $P$ instantiated to respond to user requests $r$.

A habitat consists of the connections to other habitats $c = \{p(h_1), ..., p(h_i)\}$; the genes available (gene-pool $g_p$), previously evolved gene-sets $g_s$ available, and current populations $P$ being evolved, $habitat = h = \{c, g_p, g_s, \{P_1, ..., P_j\}\}$. \quad (2)

The landscape, in energy-centric biological ecosystems, defines the connectivity between habitats [12]. Connectivity
of nodes in the digital world is generally not defined by geography or spatial proximity, but by information or semantic proximity. For example, connectivity in a peer-to-peer network is based primarily on bandwidth and information content, and not geography. The island-models of distributed evolutionary computing use an information-centric model for the connectivity of nodes (islands) [10]. However, because it is generally defined for one-time use (to evolve a solution to one problem and then stop) it usually has a fixed connectivity between the nodes, and therefore a fixed topology [7]. So, supporting ecosystem-oriented distributed evolutionary computing, with dynamic multi-objective selection pressures (fitness landscapes [40] with many peaks), requires a re-configurable network topology, such that habitat connectivity can be dynamically adapted based on the observed migration paths of the genes between the users within the habitat network $H$. So, based on the island-models of distributed evolutionary computing [10], each connection between the habitats is bi-directional and there is a probability associated with moving in either direction across the connection, with the connection probabilities c affecting the rate of migration of the genes. However, additionally, the connection probabilities will be updated by the success or failure of gene migration using the concept of Hebbian learning [41]: the habitats $h$ which do not successfully exchange genes $g$ will become less strongly connected, and the habitats $h$ which do successfully exchange genes $g$ will achieve stronger connections. This leads to a topology that adapts over time, resulting in a network that supports and resembles the connectivity of the user base $U$. We will further discuss a resulting topology in the next section.

When a new user $u$ joins, a habitat $h$ needs to be created for them, and most importantly connected to the correct cluster(s) within the habitat network $H = \{h_1, ..., h_i\}$. A new user’s habitat $h_u$ can be connected randomly to the habitat network $H$, as it will dynamically reconnect based upon the user’s behaviour. User profiling can also be used to help connect a new user’s habitat $h_u$ to the optimal part of the network $H$, by finding a similar user $u_s$, or asking the user to identify a similar user $u_s$, and then cloning their habitat’s $h_u$ connections $c$ (the set of probabilities for connecting to other habitats). Also, when a new habitat $h$ is created, its gene-pool $g_h$ should be created by merging the gene-pools of the habitats to which it is initially connected.

### D. Ecosystem

Our ecosystem-oriented distributed evolutionary computing supports the automatic re-combining of numerous genes, $g$, simple atomic algorithms, by their interaction in evolving populations, $P$, to meet user, $u$, requests, $r$, for algorithms, in a scalable architecture of distributed interconnected habitats, $h$. The sharing of genes $g$ between habitats $h$ ensures the system is scalable, while maintaining a high evolutionary specialisation for each user, $u$. The network of interconnected habitats, $H$, is equivalent to the abiotic environment of biological ecosystems [12], combined with the genes $g$ within the populations $P$ and the gene migration for distributed evolutionary computing, with the environmental selection pressures $S$ provided by a user base $\{u\}$, then the union of the habitats $h$ creates the ecosystem $E$. The continuous and varying user requests for algorithms provide a dynamic evolutionary pressure on the populations of genes, which have to evolve to better fulfill those user requests. So, local and global optimisations concurrently operate to determine solutions to satisfy different optimisation problems. Therefore, the local search is improved through this twofold process to yield better local optima faster.

\[
ecosystem = E = (H, U, R, S).
\]

### VI. Topology

Our ecosystem-oriented form of distributed evolutionary computing allows for the connectivity of the habitats $h$ to adapt to the connectivity within the user base $U$, with a cluster of habitats representing a community within the user base. If a user $u$ is a member of more than one community, the user’s habitat $h_u$ will be in more than one cluster. This leads to a network topology that will be discovered with time, and which reflects the connectivity within the user base $U$. Similarities in requests by different users will reinforce behavioural patterns, and lead to clustering of the habitats within the ecosystem, which can occur over geography, language, etc. This will form communities for more effective information sharing, the creation of niches, and will improve the responsiveness of the system.

The connections between the habitats are self-managed, through the migration between the habitats. Essentially, successful migration will reinforce habitat connections, thereby increasing the probability of future migration along these connections. If a successful multi-hop migration occurs, then a new link between the start and end habitats can be formed. Unsuccessful migrations will lead to connections (migration probabilities) decreasing, until finally the connection is closed. So, connections between habitats $h$ are reconfigured depending on the connectivity of the user base $U$, the habitat clustering will therefore be parallel to the user communities, as shown in Figure 5. The communities will cluster over defining properties of the user base, such as language, nationality, geography, etc. So, the system will take on a topology similar to that of the user base.

Landscape connectivity plays an important part in ecosystems. When the density of habitats within an environment falls below a critical threshold, widespread species may fragment into isolated populations. Fragmentation can have several consequences. Within populations, these effects include loss of genetic diversity and detrimental inbreeding [42]. At a broader scale, isolated populations may diverge genetically, leading to speciation. Fragmentation of the habitat network...
H could occur, but only if dictated by the structure of the user base $U$. The issue of greater concern is when individual habitats $h$ become totally disconnected, which could only occur under certain conditions. One condition is that the evolved solutions (algorithms) consistently fail to satisfy user requests $r$. Another condition is when the solutions they share are undesirable to the users that are within the migration range of these solutions. These scenarios could arise because the habitat $h$ is located within the wrong cluster, in which case the user can be asked to join another cluster within the habitat network, assuming the user base $U$ is of sufficient size to provide a viable alternative.

VII. DISCUSSION

Ecosystem-oriented distributed evolutionary computing proved significantly different to natural ecosystems with respect to the information-centric dynamically re-configurable network topology. A re-configurable network topology is required to support the constantly changing multi-objective information-centric selection pressures $S$ of the user base $U$. Hence, using the concept of Hebbian learning [41], habitat connectivity is dynamically adapted based on the observed migration paths of the genes within the habitat network. We would argue that this difference is not compromises, but a unique feature that arises from mimicking natural ecosystems. Biomimicry is best when not slavish imitation; it is inspiration using the principles which nature has demonstrated to be successful design strategies [43]. So while the topology of natural ecosystems is energy-centric, the topology of ecosystem-oriented distributed evolutionary computing is information-centric.

Our ecosystem-oriented distributed evolutionary computing proposes a fundamental paradigm shift, from a pull-oriented approach to a push-oriented approach. So, instead of the pull-oriented approach of generating algorithms only upon request [46], a push-oriented approach is followed of distributing and composing algorithms pre-emptively, as well as upon request. Given the increasing need to tailor software to specific uses and users, and the need to deal with increasing software complexity, we consider this shift from pull to push will become paramount.

We have previously considered a case study, Digital Business Ecosystems [44], [17], where the genes $g$ where mapped to a hybrid construct of agents and services to create a Digital Ecosystem [20], [17] capable of supporting a business ecosystem (a networks of Small and Medium sized Enterprises). This included experimental simulation results, comparing to service-oriented computing [20] and natural ecosystems [19], [17]. We also considered an abstraction of this Digital Ecosystem, based within Complex Adaptive Systems, for applicability to other classes of systems that could be considered as, or augmented to, ‘eco’-systems [45].

VIII. CONCLUSION

We have defined the fundamentals for a new class of system, ecosystem-oriented distributed evolutionary computing, created through combining understanding from theoretical ecology, evolutionary theory, and distributed evolutionary computing. By comparing and contrasting the relevant theoretical ecology we have examined how some ecological features may emerge in distributed evolutionary computing. Specifically, we suggested that ecosystem-oriented distributed evolutionary computing, like natural ecosystems, will usually consist of self-replicating individuals that interact both with one another and with an external environment [12]. Population dynamics and evolution, and spatial and network interactions, will all influence the behaviour of these systems. Many of these properties can be understood via well-known ecological models [47], [29]. These models provide a theoretical basis for the occurrence of self-organisation, in artificial and biological ecosystems, resulting from the interactions among the individuals and their environment, leading to complex non-linear behaviour [47], [29], [6]; and it is this property that provides the underlying potential for scalable problem-solving in artificial environments.

We have strived such that the word ecosystem is more than just a metaphor, instead creating an artificial counterpart of natural ecosystems, and therefore potentially having their desirable properties, such as scalability and self-organisation. The stability, convergence and equilibrium of the evolving populations of our ecosystem-oriented distributed evolutionary computing have been analysed here [43], [49], while their complexity and self-organisation have been considered here [50], [51], [52], and the overall diversity of the ecosystem has been considered here [53]. This, and other work, will help to provide theoretical bounds on the control parameters. There is also an obviously desirability to accelerate and optimise our ecosystem-oriented distributed evolutionary computing, for which we have explored the application of augmentations that interact with the ecosystem dynamics [54].

The ever-increasing challenge of software complexity in creating progressively more sophisticated and distributed applications, makes the design and maintenance of efficient and flexible systems a growing challenge [55], for which current software development techniques have hit a complexity wall [56]. In response we have created ecosystem-oriented distributed evolutionary computing, possessing properties of self-organisation, scalability and sustainability from ecosystems [6]. So, overcoming the challenge by allowing the automated
search for new algorithms in a scalable architecture, through the evolution of software in a distributed network.

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