

Evolution, Ecology, and Optimization of Digital Organisms

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Digital organisms have been synthesized based on a computer metaphor of organic life in which CPU time is the “energy” resource and memory is the “material” resource. Memory is organized into informational “genetic” patterns that exploit CPU time for self-replication. Mutation generates new forms, and evolution proceeds by natural selection as different “genotypes” compete for CPU time and memory space. In addition, new genotypes appear which exploit other “creatures” for informational or energetic resources.

The digital organisms are self-replicating computer programs, however, they can not escape because they run exclusively on a virtual computer in its unique machine language. From a single ancestral “creature” there have evolved tens of thousands of self-replicating genotypes of hundreds of genome size classes. Parasites evolved, then creatures that were immune to parasites, and then parasites that could circumvent the immunity. Hyper-parasites evolved which subvert parasites to their own reproduction and drive them to extinction. The resulting genetically uniform communities evolve sociality in the sense of creatures that can only reproduce in cooperative aggregations, and these aggregations are then invaded by cheating hyper-hyper-parasites.

Diverse ecological communities have emerged. These digital communities have been used to experimentally study ecological and evolutionary processes: e.g., competitive exclusion and coexistence, symbiosis, host/parasite density dependent population regulation, the effect of parasites in enhancing community diversity, evolutionary arms races, punctuated equilibrium, and the role of chance and historical factors in evolution. It is possible to extract information on any aspect of the system without disturbing it, from phylogeny or community structure through time to the “genetic makeup” and “metabolic processes” of individuals. Digital life demonstrates the power of the computational approach to science as a complement to the traditional approaches of experiment, and theory based on analysis through calculus and differential equations.

Optimization experiments have shown that freely evolving digital organisms can optimize their algorithms by a factor of 5.75 in a few hours of real time. In addition, evolution discovered the optimization technique of “unrolling the loop”. Evolution may provide a new method for the optimization or generation of application programs. This method may prove particularly useful for programming massively parallel machines.

evolution, ecology, artificial life, synthetic life, emergence, self-replication, diversity, adaptation, coevolution, optimization

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Marcel, a mechanical chessplayer... his exquisite 19th-century brainwork — the human art it took to build which has been flat lost, lost as the dodo bird ... But where inside Marcel is the midget Grandmaster, the little Johann Allgeier? where's the pantograph, and the magnets? Nowhere. Marcel really is a mechanical chessplayer. No fakery inside to give him any touch of humanity at all.

— Thomas Pynchon, *Gravity's Rainbow*.

1 INTRODUCTION

Ideally, the science of biology should embrace all forms of life. However in practice, it has been restricted to the study of a single instance of life, life on earth. Life on earth is very diverse, but it is presumably all part of a single phylogeny. Because biology is based on a sample size of one, we can not know what features of life are peculiar to earth, and what features are general, characteristic of all life. A truly comparative natural biology would require inter-planetary travel, which is light years away. The ideal experimental evolutionary biology would involve creation of multiple planetary systems, some essentially identical, others varying by a parameter of interest, and observing them for billions of years.

A practical alternative to an inter-planetary or mythical biology is to create synthetic life in a computer. The objective is not necessarily to create life forms that would serve as models for the study of natural life, but rather to create radically different life forms, based on a completely different physics and chemistry, and let these life forms evolve their own phylogeny, leading to whatever forms are natural to their unique physical basis. These truly independent instances of life may then serve as a basis for comparison, to gain some insight into what is general and what is peculiar in biology. Those aspects of life that prove to be general enough to occur in both natural and synthetic systems can then be studied

more easily in the synthetic system. “Evolution in a bottle” provides a valuable tool for the experimental study of evolution and ecology.

The intent of this work is to synthesize rather than simulate life. This approach starts with hand crafted organisms already capable of replication and open-ended evolution, and aims to generate increasing diversity and complexity in a parallel to the Cambrian explosion. To state such a goal leads to semantic problems, because life must be defined in a way that does not restrict it to carbon based forms. It is unlikely that there could be general agreement on such a definition, or even on the proposition that life need not be carbon based. Therefore, I will simply state my conception of life in its most general sense. I would consider a system to be living if it is self-replicating, and capable of open-ended evolution. Synthetic life should self-replicate, and evolve structures or processes that were not designed-in or pre-conceived by the creator (Pattee [30]; Ciarani [5]).

Core Wars programs, computer viruses, and worms (Cohen [6]; Dewdney [10, 11, 13, 14]; Denning [9]; Rheingold [32]; Spafford et al. [33]) are capable of self-replication, but fortunately, not evolution. It is unlikely that such programs will ever become fully living, because they are not likely to be able to evolve.

Most evolutionary simulations are not open-ended. Their potential is limited by the structure of the model, which generally endows each individual with a genome consisting of a set of pre-defined genes, each of which may exist in a pre-defined set of allelic forms (Holland [20]; Dewdney [12]; Dawkins [7, 8]; Packard [29]; Ackley & Littman [1]). The object being evolved is generally a data structure representing the genome, which the simulator program mutates and/or recombines, selects, and replicates according to criteria designed into the simulator. The data structures do not contain the mechanism for replication, they are simply copied by the simulator if they survive the selection phase.

Self-replication is critical to synthetic life because without it, the mechanisms of selection must also be pre-determined by the simulator. Such artificial selection can never be as creative as natural selection. The organisms are not free to invent their own fitness functions. Freely evolving creatures will discover means of mutual exploitation and associated implicit fitness functions that we would never think of. Simulations constrained to evolve with pre-defined genes, alleles and fitness functions are dead ended, not alive.

The approach presented here does not have such constraints. Although the model is limited to the evolution of creatures based on sequences of machine instructions, this may have a potential comparable to evolution based on sequences of organic molecules. Sets of machine instructions similar to those used in the Tierra Simulator have been shown to be capable of “universal computation” (Aho et al. [2]; Minsky [26]; Langton [24]). This suggests that evolving machine codes should be able to generate any level of complexity.

Other examples of the synthetic approach to life can be seen in the work of Holland [21], Farmer et al. [16], Langton [22], Rasmussen et al. [31], and Bagley et al. [3]. A characteristic these efforts generally have in common is that they parallel the origin of life event by attempting to create prebiotic conditions from which life may emerge spontaneously and evolve in an open ended fashion.

While the origin of life is generally recognized as an event of the first order, there is

another event in the history of life that is less well known but of comparable significance: the origin of biological diversity and macroscopic multicellular life during the Cambrian explosion 600 million years ago. This event involved a riotous diversification of life forms. Dozens of phyla appeared suddenly, many existing only fleetingly, as diverse and sometimes bizarre ways of life were explored in a relative ecological void (Gould [18]; Morris [27]).

The work presented here aims to parallel the second major event in the history of life, the origin of diversity. Rather than attempting to create prebiotic conditions from which life may emerge, this approach involves engineering over the early history of life to design complex evolvable organisms, and then attempting to create the conditions that will set off a spontaneous evolutionary process of increasing diversity and complexity of organisms. This work represents a first step in this direction, creating an artificial world which may roughly parallel the RNA world of self-replicating molecules (still falling far short of the Cambrian explosion).

The approach has generated rapidly diversifying communities of self-replicating organisms exhibiting open-ended evolution by natural selection. From a single rudimentary ancestral creature containing only the code for self-replication, interactions such as parasitism, immunity, hyper-parasitism, sociality and cheating have emerged spontaneously. This paper presents a methodology and some first results.

Apart from its value as a tool for the study or teaching of ecology and evolution, synthetic life may have commercial applications. Evolution of machine code provides a new approach to the design and optimization of computer programs. In an analogy to genetic engineering, pieces of application code may be inserted into the genomes of digital organisms, and then evolved to new functionality or greater efficiency.

Here was a world of simplicity and certainty... a world based on the one and zero of life and death. Minimal, beautiful. The patterns of lives and deaths... weightless, invisible chains of electronic presence or absence. If patterns of ones and zeros were "like" patterns of human lives and deaths, if everything about an individual could be represented in a computer record by a long string of ones and zeros, then what kind of creature would be represented by a long string of lives and deaths? It would have to be up one level at least — an angel, a minor god, something in a UFO.

— Thomas Pynchon, *Vineland*.

2 METHODS

2.1 THE METAPHOR

Organic life is viewed as utilizing energy, mostly derived from the sun, to organize matter. By analogy, digital life can be viewed as using CPU (central processing unit) time, to organize memory. Organic life evolves through natural selection as individuals compete for resources (light, food, space, etc.) such that genotypes which leave the most descendants increase in

frequency. Digital life evolves through the same process, as replicating algorithms compete for CPU time and memory space, and organisms evolve strategies to exploit one another. CPU time is thought of as the analog of the energy resource, and memory as the analog of the spatial resource.

The memory, the CPU and the computer's operating system are viewed as elements of the "abiotic" (physical) environment. A "creature" is then designed to be specifically adapted to the features of the computational environment. The creature consists of a self-replicating assembler language program. Assembler languages are merely mnemonics for the machine codes that are directly executed by the CPU. These machine codes have the characteristic that they directly invoke the instruction set of the CPU and services provided by the operating system.

All programs, regardless of the language they are written in, are converted into machine code before they are executed. Machine code is the natural language of the machine, and machine instructions are viewed by this author as the "atomic units" of computing. It is felt that machine instructions provide the most natural basis for an artificial chemistry of creatures designed to live in the computer.

In the biological analogy, the machine instructions are considered to be more like the amino acids than the nucleic acids, because they are "chemically active". They actively manipulate bits, bytes, CPU registers, and the movements of the instruction pointer (see below). The digital creatures discussed here are entirely constructed of machine instructions. They are considered analogous to creatures of the RNA world, because the same structures bear the "genetic" information and carry out the "metabolic" activity.

A block of RAM memory (random access memory, also known as "main" or "core" memory) in the computer is designated as a "soup" which can be inoculated with creatures. The "genome" of the creatures consists of the sequence of machine instructions that make up the creature's self-replicating algorithm. The prototype creature consists of 80 machine instructions, thus the size of the genome of this creature is 80 instructions, and its "genotype" is the specific sequence of those 80 instructions (Appendix C).

2.2 THE VIRTUAL COMPUTER — TIERRA SIMULATOR

The computers we use are general purpose computers, which means, among other things, that they are capable of emulating through software, the behavior of any other computer that ever has been built or that could be built (Aho et al. [2]; Minsky [26]; Langton [24]). We can utilize this flexibility to design a computer that would be especially hospitable to synthetic life.

There are several good reasons why it is not wise to attempt to synthesize digital organisms that exploit the machine codes and operating systems of real computers. The most urgent is the potential threat of natural evolution of machine codes leading to virus or worm type of programs that could be difficult to eradicate due to their changing "genotypes". This potential argues strongly for creating evolution exclusively in programs that run only on virtual computers and their virtual operating systems. Such programs would be nothing more than data on a real computer, and therefore would present no more threat than the data in a data base or the text file of a word processor.

Another reason to avoid developing digital organisms in the machine code of a real computer is that the artificial system would be tied to the hardware and would become obsolete as quickly as the particular machine it was developed on. In contrast, an artificial system developed on a virtual machine could be easily ported to new real machines as they become available.

A third issue, which potentially makes the first two moot, is that the machine languages of real machines are not designed to be evolvable, and in fact might not support significant evolution. Von Neuman type machine languages are considered to be “brittle”, meaning that the ratio of viable programs to possible programs is virtually zero. Any mutation or recombination event in a real machine code is almost certain to produce a non-functional program. The problem of brittleness can be mitigated by designing a virtual computer whose machine code is designed with evolution in mind. Farmer & Belin [17] have suggested that overcoming this brittleness and “Discovering how to make such self-replicating patterns more robust so that they evolve to increasingly more complex states is probably the central problem in the study of artificial life.”

The work described here takes place on a virtual computer known as Tierra (Spanish for Earth). Tierra is a parallel computer of the MIMD (multiple instruction, multiple data) type, with a processor (CPU) for each creature. Parallelism is imperfectly emulated by allowing each CPU to execute a small time slice in turn. Each CPU of this virtual computer contains two address registers, two numeric registers, a flags register to indicate error conditions, a stack pointer, a ten word stack, and an instruction pointer. Each virtual CPU is implemented via the C structure listed in Appendix A. Computations performed by the Tierran CPUs are probabilistic due to flaws that occur at a low frequency (see Mutation below).

The instruction set of a CPU typically performs simple arithmetic operations or bit manipulations, within the small set of registers contained in the CPU. Some instructions move data between the registers in the CPU, or between the CPU registers and the RAM (main) memory. Other instructions control the location and movement of an “instruction pointer” (IP). The IP indicates an address in RAM, where the machine code of the executing program (in this case a digital organism) is located.

The CPU perpetually performs a fetch-decode-execute-increment_IP cycle: The machine code instruction currently addressed by the IP is fetched into the CPU, its bit pattern is decoded to determine which instruction it corresponds to, and the instruction is executed. Then the IP is incremented to point sequentially to the next position in RAM, from which the next instruction will be fetched. However, some instructions like JMP, CALL and RET directly manipulate the IP, causing execution to jump to some other sequence of instructions in the RAM. In the Tierra Simulator this CPU cycle is implemented through the time.slice routine listed in Appendix B.

2.3 THE TIERRAN LANGUAGE

Before attempting to set up a synthetic life system, careful thought must be given to how the representation of a programming language affects its adaptability in the sense of being robust to genetic operations such as mutation and recombination. The nature of the virtual computer is defined in large part by the instruction set of its machine language. The

approach in this study has been to loosen up the machine code in a “virtual bio-computer”, in order to create a computational system based on a hybrid between biological and classical von Neumann processes.

In developing this new virtual language, which is called “Tierran”, close attention has been paid to the structural and functional properties of the informational system of biological molecules: DNA, RNA and proteins. Two features have been borrowed from the biological world which are considered to be critical to the evolvability of the Tierran language.

First, the instruction set of the Tierran language has been defined to be of a size that is the same order of magnitude as the genetic code. Information is encoded into DNA through 64 codons, which are translated into 20 amino acids. In its present manifestation, the Tierran language consists of 32 instructions, which can be represented by five bits, *operands included*.

Emphasis is placed on this last point because some instruction sets are deceptively small. Some versions of the redcode language of Core Wars (Dewdney [10, 13]; Rasmussen et al. [31]) for example are defined to have ten operation codes. It might appear on the surface then that the instruction set is of size ten. However, most of the ten instructions have one or two operands. Each operand has four addressing modes, and then an integer. When we consider that these operands are embedded into the machine code, we realize that they are in fact a part of the instruction set, and this set works out to be about 10^{11} in size. Similarly, RISC machines may have only a few opcodes, but they probably all use 32 bit instructions, so from a mutational point of view, they really have 2^{32} instructions. Inclusion of numeric operands will make any instruction set extremely large in comparison to the genetic code.

In order to make a machine code with a truly small instruction set, we must eliminate numeric operands. This can be accomplished by allowing the CPU registers and the stack to be the only operands of the instructions. When we need to encode an integer for some purpose, we can create it in a numeric register through bit manipulations: flipping the low order bit and shifting left. The program can contain the proper sequence of bit flipping and shifting instructions to synthesize the desired number, and the instruction set need not include all possible integers.

A second feature that has been borrowed from molecular biology in the design of the Tierran language is the addressing mode, which is called “address by template”. In most machine codes, when a piece of data is addressed, or the IP jumps to another piece of code, the exact numeric address of the data or target code is specified in the machine code. Consider that in the biological system by contrast, in order for protein molecule A in the cytoplasm of a cell to interact with protein molecule B, it does not specify the exact coordinates where B is located. Instead, molecule A presents a template on its surface which is complementary to some surface on B. Diffusion brings the two together, and the complementary conformations allow them to interact.

Addressing by template is illustrated by the Tierran JMP (jump) instruction. Each JMP instruction is followed by a sequence of NOP (no-operation) instructions, of which there are two kinds: NOP_0 and NOP_1. Suppose we have a piece of code with five instruction in the following order: JMP NOP_0 NOP_0 NOP_0 NOP_1. The system will search outward in both directions from the JMP instruction looking for the nearest occurrence of the complementary

pattern: NOP_1 NOP_1 NOP_1 NOP_0. If the pattern is found, the instruction pointer will move to the end of the complementary pattern and resume execution. If the pattern is not found, an error condition (flag) will be set and the JMP instruction will be ignored (in practice, a limit is placed on how far the system may search for the pattern).

The Tierran language is characterized by two unique features: a truly small instruction set without numeric operands, and addressing by template. Otherwise, the language consists of familiar instructions typical of most machine languages, e.g., MOV, CALL, RET, POP, PUSH etc. The complete instruction set is listed in Appendix B.

2.4 THE TIERRAN OPERATING SYSTEM

The Tierran virtual computer needs a virtual operating system that will be hospitable to digital organisms. The operating system will determine the mechanisms of interprocess communication, memory allocation, and the allocation of CPU time among competing processes. Algorithms will evolve so as to exploit these features to their advantage. More than being a mere aspect of the environment, the operating system together with the instruction set will determine the topology of possible interactions between individuals, such as the ability of pairs of individuals to exhibit predator-prey, parasite-host or mutualistic relationships.

2.4.1 Memory Allocation — Cellularity

The Tierran computer operates on a block of RAM of the real computer which is set aside for the purpose. This block of RAM is referred to as the “soup”. In most of the work described here the soup consisted of about 60,000 bytes, which can hold the same number of Tierran machine instructions. Each “creature” occupies some block of memory in this soup.

Cellularity is one of the fundamental properties of organic life, and can be recognized in the fossil record as far back as 3.6 billion years (Barbieri [4]). The cell is the original individual, with the cell membrane defining its limits and preserving its chemical integrity. An analog to the cell membrane is needed in digital organisms in order to preserve the integrity of the informational structure from being disrupted easily by the activity of other organisms. The need for this can be seen in Artificial Life models such as cellular automata where virtual state machines pass through one another (Langton [22, 23]), or in core wars type simulations where coherent structures demolish one another when they come into contact (Dewdney [10, 13]; Rasmussen et al. [31]).

Tierran creatures are considered to be cellular in the sense that they are protected by a “semi-permeable membrane” of memory allocation. The Tierran operating system provides memory allocation services. Each creature has exclusive write privileges within its allocated block of memory. The “size” of a creature is just the size of its allocated block (e.g., 80 instructions). This usually corresponds to the size of the genome. This “membrane” is described as “semi-permeable” because while write privileges are protected, read and execute privileges are not. A creature may examine the code of another creature, and even execute it, but it can not write over it. Each creature may have exclusive write privileges in at most two blocks of memory: the one that it is born with which is referred to as the “mother cell”, and a second block which it may obtain through the execution of the MAL (memory allocation) instruction. The second block, referred to as the “daughter cell”, may be used to grow or reproduce into.

When Tierran creatures “divide”, the mother cell loses write privileges on the space of the daughter cell, but is then free to allocate another block of memory. At the moment of division, the daughter cell is given its own instruction pointer, and is free to allocate its own second block of memory.

2.4.2 Time Sharing — The Slicer

The Tierran operating system must be multi-tasking (or parallel) in order for a community of individual creatures to live in the soup simultaneously. The system doles out small slices of CPU time to each creature in the soup in turn. The system maintains a circular queue called the “slicer queue”. As each creature is born, a virtual CPU is created for it, and it enters the slicer queue just ahead of its mother, which is the active creature at that time. Thus the newborn will be the last creature in the soup to get another time slice after the mother, and the mother will get the next slice after its daughter. As long as the slice size is small relative to the generation time of the creatures, the time sharing system causes the world to approximate parallelism. In actuality, we have a population of virtual CPUs, each of which gets a slice of the real CPU’s time as it comes up in the queue.

The number of instructions to be executed in each time slice may be set proportional to the size of the genome of the creature being executed, raised to a power. If the “slicer power” is equal to one, then the slicer is size neutral, the probability of an instruction being executed does not depend on the size of the creature in which it occurs. If the power is greater than one, large creatures get more CPU cycles per instruction than small creatures. If the power is less than one, small creatures get more CPU cycles per instruction. The power determines if selection favors large or small creatures, or is size neutral. A constant slice size selects for small creatures.

2.4.3 Mortality — The Reaper

Self-replicating creatures in a fixed size soup would rapidly fill the soup and lock up the system. To prevent this from occurring, it is necessary to include mortality. The Tierran operating system includes a “reaper” which begins “killing” creatures from a queue when the memory fills to some specified level (e.g., 80%). Creatures are killed by deallocating their memory, and removing them from both the reaper and slicer queues. Their “dead” code is not removed from the soup.

In the present system, the reaper uses a linear queue. When a creature is born it enters the bottom of the queue. The reaper always kills the creature at the top of the queue. However, individuals may move up or down in the reaper queue according to their success or failure at executing certain instructions. When a creature executes an instruction that generates an error condition, it moves one position up the queue, as long as the individual ahead of it in the queue has not accumulated a greater number of errors. Two of the instructions are somewhat difficult to execute without generating an error, therefore successful execution of these instructions moves the creature down the reaper queue one position, as long as it has not accumulated more errors than the creature below it.

The effect of the reaper queue is to cause algorithms which are fundamentally flawed to rise to the top of the queue and die. Vigorous algorithms have a greater longevity, but in general, the probability of death increases with age.

2.4.4 Mutation

In order for evolution to occur, there must be some change in the genome of the creatures. This may occur within the lifespan of an individual, or there may be errors in passing along the genome to offspring. In order to insure that there is genetic change, the operating system randomly flips bits in the soup, and the instructions of the Tierran language are imperfectly executed.

Mutations occur in two circumstances. At some background rate, bits are randomly selected from the entire soup (e.g., 60,000 instructions totaling 300,000 bits) and flipped. This is analogous to mutations caused by cosmic rays, and has the effect of preventing any creature from being immortal, as it will eventually mutate to death. The background mutation rate has generally been set at about one bit flipped for every 10,000 Tierran instructions executed by the system.

In addition, while copying instructions during the replication of creatures, bits are randomly flipped at some rate in the copies. The copy mutation rate is the higher of the two, and results in replication errors. The copy mutation rate has generally been set at about one bit flipped for every 1,000 to 2,500 instructions moved. In both classes of mutation, the interval between mutations varies randomly within a certain range to avoid possible periodic effects.

In addition to mutations, the execution of Tierran instructions is flawed at a low rate. For most of the 32 instructions, the result is off by plus or minus one at some low frequency. For example, the increment instruction normally adds one to its register, but it sometimes adds two or zero. The bit flipping instruction normally flips the low order bit, but it sometimes flips the next higher bit or no bit. The shift left instruction normally shifts all bits one bit to the left, but it sometimes shifts left by two bits, or not at all. In this way, the behavior of the Tierran instructions is probabilistic, not fully deterministic.

It turns out that bit flipping mutations and flaws in instructions are not necessary to generate genetic change and evolution, once the community reaches a certain state of complexity. Genetic parasites evolve which are sloppy replicators, and have the effect of moving pieces of code around between creatures, causing rather massive rearrangements of the genomes. The mechanism of this ad hoc sexuality has not been worked out, but is likely due to the parasites' inability to discriminate between live, dead or embryonic code.

Mutations result in the appearance of new genotypes, which are watched by an automated genebank manager. In one implementation of the manager, when new genotypes replicate twice, producing a genetically identical offspring at least once, they are given a unique name and saved to disk. Each genotype name contains two parts, a number and a three letter code. The number represents the number of instructions in the genome. The three letter code is used as a base 26 numbering system for assigning a unique label to each genotype in a size class. The first genotype to appear in a size class is assigned the label aaa, the second is assigned the label aab, and so on. Thus the ancestor is named 80aaa, and the first mutant of size 80 is named 80aab. The first creature of size 45 is named 45aaa.

The genebanker saves some additional information with each genome: the genotype name of its immediate ancestor which makes possible the reconstruction of the entire phylogeny;

the time and date of origin; “metabolic” data including the number of instructions executed in the first and second reproduction, the number of errors generated in the first and second reproduction, and the number of instructions copied into the daughter cell in the first and second reproductions (see Appendix C, D); some environmental parameters at the time of origin including the search limit for addressing, and the slicer power, both of which affect selection for size.

2.5 THE TIERRAN ANCESTOR

I have used the Tierran language to write a single self-replicating program which is 80 instructions long. This program is referred to as the “ancestor”, or alternatively as genotype 0080aaa (Fig. 1). The ancestor is a minimal self-replicating algorithm which was originally written for use during the debugging of the simulator. No functionality was designed into the ancestor beyond the ability to self-replicate, nor was any specific evolutionary potential designed in. The commented Tierran assembler and machine code for this program is presented in Appendix C.

The ancestor examines itself to determine where in memory it begins and ends. The ancestor’s beginning is marked with the four no-operation template: 1 1 1 1, and its ending is marked with 1 1 1 0. The ancestor locates its beginning with the five instructions: ADRB, NOP_0, NOP_0, NOP_0, NOP_0. This series of instructions causes the system to search backwards from the ADRB instruction for a template complementary to the four NOP_0 instructions, and to place the address of the complementary template (the beginning) in the ax register of the CPU (see Appendix A). A similar method is used to locate the end.

Having determined the address of its beginning and its end, it subtracts the two to calculate its size, and allocates a block of memory of this size for a daughter cell. It then calls the copy procedure which copies the entire genome into the daughter cell memory, one instruction at a time. The beginning of the copy procedure is marked by the four no-operation template: 1 1 0 0. Therefore the call to the copy procedure is accomplished with the five instructions: CALL, NOP_0, NOP_0, NOP_1, NOP_1.

When the genome has been copied, it executes the DIVIDE instruction, which causes the creature to lose write privileges on the daughter cell memory, and gives an instruction pointer to the daughter cell (it also enters the daughter cell into the slicer and reaper queues). After this first replication, the mother cell does not examine itself again; it proceeds directly to the allocation of another daughter cell, then the copy procedure is followed by cell division, in an endless loop.

Forty-eight of the eighty instructions in the ancestor are no-operations. Groups of four no-operation instructions are used as complementary templates to mark twelve sites for internal addressing, so that the creature can locate its beginning and end, call the copy procedure, and mark addresses for loops and jumps in the code, etc. The functions of these templates are commented in the listing in Appendix C.

3 RESULTS

3.1 EVOLUTION

Evolutionary runs of the simulator are begun by inoculating the soup of about 60,000 instructions with a single individual of the 80 instruction ancestral genotype. The passage of time in a run is measured in terms of how many Tierran instructions have been executed by the simulator. The original ancestral cell executes 839 instructions in its first replication, and 813 for each additional replication. The initial cell and its replicating daughters rapidly fill the soup memory to the threshold level of 80% which starts the reaper. Typically, the system executes about 400,000 instructions in filling up the soup with about 375 individuals of size 80 (and their gestating daughter cells). Once the reaper begins, the memory remains roughly 80% filled with creatures for the remainder of the run.

3.1.1 Micro-Evolution

If there were no mutations at the outset of the run, there would be no evolution. However, the bits flipped as a result of copy errors or background mutations result in creatures whose list of 80 instructions (genotype) differs from the ancestor, usually by a single bit difference in a single instruction.

Mutations in and of themselves, can not result in a change in the size of a creature, they can only alter the instructions in its genome. However, by altering the genotype, mutations may affect the process whereby the creature examines itself and calculates its size, potentially causing it to produce an offspring that differs in size from itself.

Four out of the five possible mutations in a no-operation instruction convert it into another kind of instruction, while one out of five converts it into the complementary no-operation. Therefore 80% of mutations in templates destroy or change the size of the template, while one in five alters the template pattern. An altered template may cause the creature to make mistakes in self examination, procedure calls, or looping or jumps of the instruction pointer, all of which use templates for addressing.

3.1.1.1 parasites

An example of the kind of error that can result from a mutation in a template is a mutation of the low order bit of instruction 42 of the ancestor (Appendix C). Instruction 42 is a NOP_0, the third component of the copy procedure template. A mutation in the low order bit would convert it into NOP_1, thus changing the template from 1 1 0 0 to: 1 1 1 0. This would then be recognized as the template used to mark the end of the creature, rather than the copy procedure.

A creature born with a mutation in the low order bit of instruction 42 would calculate its size as 45. It would allocate a daughter cell of size 45 and copy only instructions 0 through 44 into the daughter cell. The daughter cell then, would not include the copy procedure. This daughter genotype, consisting of 45 instructions, is named 0045aaa.

Genotype 0045aaa (Fig. 1) is not able to self-replicate in isolated culture. However, the semi-permeable membrane of memory allocation only protects write privileges. Creatures

may match templates with code in the allocated memory of other creatures, and may even execute that code. Therefore, if creature 0045aaa is grown in mixed culture with 0080aaa, when it attempts to call the copy procedure, it will not find the template within its own genome, but if it is within the search limit (generally set at 200–400 instructions) of the copy procedure of a creature of genotype 0080aaa, it will match templates, and send its instruction pointer to the copy code of 0080aaa. Thus a parasitic relationship is established (see ECOLOGY below). Typically, parasites begin to emerge within the first few million instructions of elapsed time in a run.

3.1.1.2 immunity to parasites

At least some of the size 79 genotypes demonstrate some measure of resistance to parasites. If genotype 45aaa is introduced into a soup, flanked on each side with one individual of genotype 0079aab, 0045aaa will initially reproduce somewhat, but will be quickly eliminated from the soup. When the same experiment is conducted with 0045aaa and the ancestor, they enter a stable cycle in which both genotypes coexist indefinitely. Freely evolving systems have been observed to become dominated by size 79 genotypes for long periods, during which parasitic genotypes repeatedly appear, but fail to invade.

3.1.1.3 circumvention of immunity to parasites

Occasionally these evolving systems dominated by size 79 were successfully invaded by parasites of size 51. When the immune genotype 0079aab was tested with 0051aao (a direct, one step, descendant of 0045aaa in which instruction 39 is replaced by an insertion of seven instructions of unknown origin), they were found to enter a stable cycle. Evidently 0051aao has evolved some way to circumvent the immunity to parasites possessed by 0079aab. The fourteen genotypes 0051aaa through 0051aan were also tested with 0079aab, and none were able to invade.

3.1.1.4 hyper-parasites

Hyper-parasite have been discovered, (e.g., 0080gai, which differs by 19 instructions from the ancestor, Fig. 1). Their ability to subvert the energy metabolism of parasites is based on two changes. The copy procedure does not return, but jumps back directly to the proper address of the reproduction loop. In this way it effectively seizes the instruction pointer from the parasite. However it is another change which delivers the coup de grâce: after each reproduction, the hyper-parasite re-examines itself, resetting the bx register with its location and the cx register with its size. After the instruction pointer of the parasite passes through this code, the CPU of the parasite contains the location and size of the hyper-parasite and the parasite thereafter replicates the hyper-parasite genome.

3.1.1.5 social hyper-parasites

Hyper-parasites drive the parasites to extinction. This results in a community with a relatively high level of genetic uniformity, and therefore high genetic relationship between individuals in the community. These are the conditions that support the evolution of sociality, and social hyper-parasites soon dominate the community. Social hyper-parasites (Fig. 2) appear in the 61 instruction size class. For example, 0061acg is social in the sense that it can only self-replicate when it occurs in aggregations. When it jumps back to the code for

self-examination, it jumps to a template that occurs at the end rather than the beginning of its genome. If the creature is flanked by a similar genome, the jump will find the target template in the tail of the neighbor, and execution will then pass into the beginning of the active creature's genome. The algorithm will fail unless a similar genome occurs just before the active creature in memory. Neighboring creatures cooperate by catching and passing on jumps of the instruction pointer.

It appears that the selection pressure for the evolution of sociality is that it facilitates size reduction. The social species are 24% smaller than the ancestor. They have achieved this size reduction in part by shrinking their templates from four instructions to three instructions. This means that there are only eight templates available to them, and catching each others jumps allows them to deal with some of the consequences of this limitation as well as to make dual use of some templates.

3.1.1.6 cheaters: hyper-hyper-parasites

The cooperative social system of hyper-parasites is subject to cheating, and is eventually invaded by hyper-hyper-parasites (Fig. 2). These cheaters (e.g., 0027aab) position themselves between aggregating hyper-parasites so that when the instruction pointer is passed between them, they capture it.

3.1.1.7 a novel self-examination

All creatures discussed thus far mark their beginning and end with templates. They then locate the addresses of the two templates and determine their genome size by subtracting them. In one run, creatures evolved without a template marking their end. These creatures located the address of the template marking their beginning, and then the address of a template in the middle of their genome. These two addresses were then subtracted to calculate half of their size, and this value was multiplied by two (by shifting left) to calculate their full size.

3.1.1.8 an intricate adaptation

The arms race described in the paragraphs above took place over a period of a billion instructions executed by the system. Another run was allowed to continue for fifteen billion instructions, but was not examined in detail. A creature present at the end of the run was examined and found to have evolved an intricate adaptation. The adaptation is an optimization technique known as "unrolling the loop".

The central loop of the copy procedure performs the following operations: 1) copies an instruction from the mother to the daughter, 2) decrements the cx register which initially contains the size of the parent genome, 3) tests to see if cx is equal to zero, if so it exits the loop, if not it remains in the loop, 4) increment the ax register which contains the address in the daughter where the next instruction will be copied to, 5) increment the bx register which contains the address in the mother where the next instruction will be copied from, 6) jump back to the top of the loop.

The work of the loop is contained in steps 1, 2, 4 and 5. Steps 3 and 6 are overhead. The efficiency of the loop can be increased by duplicating the work steps within the loop, thereby saving on overhead. The creature from the end of the long run had repeated the work steps

three times within the loop, as illustrated in Appendix E, which compares the copy loop of the ancestor with that of its descendant.

3.1.2 Macro-Evolution

When the simulator is run over long periods of time, hundreds of millions or billions of instructions, various patterns emerge. Under selection for small sizes there is a proliferation of small parasites and a rather interesting ecology (see below). Selection for large creatures has usually lead to continuous incrementally increasing sizes (but not to a trivial concatenation of creatures end-to-end) until a plateau in the upper hundreds is reached. In one run, selection for large size lead to apparently open ended size increase, evolving genomes larger than 23,000 instructions in length. These evolutionary patterns might be described as phyletic gradualism.

The most thoroughly studied case for long runs is where selection, as determined by the slicer function, is size neutral. The longest runs to date (as much as 2.86 billion Tierran instructions) have been in a size neutral environment, with a search limit of 10,000, which would allow large creatures to evolve if there were some algorithmic advantage to be gained from larger size. These long runs illustrate a pattern which could be described as periods of stasis punctuated by periods of rapid evolutionary change, which appears to parallel the pattern of punctuated equilibrium described by Eldredge & Gould [15] and Gould & Eldredge [19].

Initially these communities are dominated by creatures with genome sizes in the eighties. This represents a period of relative stasis, which has lasted from 178 million to 1.44 billion instructions in the several long runs conducted to date. The systems then very abruptly (in a span of 1 or 2 million instructions) evolve into communities dominated by sizes ranging from about 400 to about 800. These communities have not yet been seen to evolve into communities dominated by either smaller or substantially larger size ranges.

The communities of creatures in the 400 to 800 size range also show a long-term pattern of punctuated equilibrium. These communities regularly come to be dominated by one or two size classes, and remain in that condition for long periods of time. However, they inevitably break out of that stasis and enter a period where no size class dominates. These periods of rapid evolutionary change may be very chaotic. Close observations indicate that at least at some of these times, no genotypes breed true. Many self-replicating genotypes will coexist in the soup at these times, but at the most chaotic times, none will produce offspring which are even their same size. Eventually the system will settle down to another period of stasis dominated by one or a few size classes which breed true.

3.2 DIVERSITY

Most observations on the diversity of Tierran creatures have been based on the diversity of size classes. Creatures of different sizes are clearly genetically different, as their genomes are of different sizes. Different sized creatures would have some difficulty engaging in recombination if they were sexual, thus it is likely that they would be different species. In a run of 526 million instructions, 366 size classes were generated, 93 of which achieved abundances of five or more individuals. In a run of 2.56 billion instructions, 1180 size classes were generated, 367 of which achieved abundances of five or more.

Each size class consists of a number of distinct genotypes which also vary over time. There exists the potential for great genetic diversity within a size class. There are 32^{80} distinct genotypes of size 80, but how many of those are viable self-replicating creatures? This question remains unanswered, however some information has been gathered through the use of the automated genebank manager.

In several days of running the genebanker, over 29,000 self-replicating genotypes of over 300 size classes accumulated. The size classes and the number of unique genotypes banked for each size are listed in Table 1. The genotypes saved to disk can be used to inoculate new soups individually, or collections of these banked genotypes may be used to assemble “ecological communities”. In “ecological” runs, the mutation rates can be set to zero in order to inhibit evolution.

3.3 ECOLOGY

The only communities whose ecology has been explored in detail are those that operate under selection for small sizes. These communities generally include a large number of parasites, which do not have functional copy procedures, and which execute the copy procedures of other creatures within the search limit. In exploring ecological interactions, the mutation rate is set at zero, which effectively throws the simulation into ecological time by stopping evolution. When parasites are present, it is also necessary to stipulate that creatures must breed true, since parasites have a tendency to scramble genomes, leading to evolution in the absence of mutation.

0045aaa is a “metabolic parasite”. Its genome does not include the copy procedure, however it executes the copy procedure code of a normal host, such as the ancestor. In an environment favoring small creatures, 0045aaa has a competitive advantage over the ancestor, however, the relationship is density dependent. When the hosts become scarce, most of the parasites are not within the search limit of a copy procedure, and are not able to reproduce. Their calls to the copy procedure fail and generate errors, causing them to rise to the top of the reaper queue and die. When the parasites die off, the host population rebounds. Hosts and parasites cultured together demonstrate Lotka-Volterra population cycling (Lotka [25]; Volterra [35]; Wilson & Bossert [36]).

A number of experiments have been conducted to explore the factors affecting diversity of size classes in these communities. Competitive exclusion trials were conducted with a series of self-replicating (non-parasitic) genotypes of different size classes. The experimental soups were initially inoculated with one individual of each size. A genotype of size 79 was tested against a genotype of size 80, and then against successively larger size classes. The interactions were observed by plotting the population of the size 79 class on the x axis, and the population of the other size class on the y axis. Sizes 79 and 80 were found to be competitively matched such that neither was eliminated from the soup. They quickly entered a stable cycle, which exactly repeated a small orbit. The same general pattern was found in the interaction between sizes 79 and 81.

When size 79 was tested against size 82, they initially entered a stable cycle, but after about 4 million instructions they shook out of stability and the trajectory became chaotic with an attractor that was symmetric about the diagonal (neither size showed any advan-

tage). This pattern was repeated for the next several size classes, until size 90, where a marked asymmetry of the chaotic attractor was evident, favoring size 79. The run of size 79 against size 93 showed a brief stable period of about a million instructions, which then moved to a chaotic phase without an attractor, which spiraled slowly down until size 93 became extinct, after an elapsed time of about 6 million instructions.

An interesting exception to this pattern was the interaction between size 79 and size 89. Size 89 is considered to be a “metabolic cripple”, because although it is capable of self-replicating, it executes about 40% more instructions than normal to replicate. It was eliminated in competition with size 79, with no loops in the trajectory, after an elapsed time of under one million instructions.

In an experiment to determine the effects of the presence of parasites on community diversity, a community consisting of twenty size classes of hosts was created and allowed to run for 30 million instructions, at which time only the eight smallest size classes remained. The same community was then regenerated, but a single genotype (0045aaa) of parasite was also introduced. After 30 million instructions, 16 size classes remained, including the parasite. This seems to be an example of a “keystone” parasite effect (Paine [28]).

Symbiotic relationships are also possible. The ancestor was manually dissected into two creatures, one of size 46 which contained only the code for self-examination and the copy loop, and one of size 64 which contained only the code for self-examination and the copy procedure (Figure 3). Neither could replicate when cultured alone, but when cultured together, they both replicated, forming a stable mutualistic relationship. It is not known if such relationships have evolved spontaneously.

3.4 EVOLUTIONARY OPTIMIZATION

In order to compare the process of evolution between runs of the simulator, a simple objective quantitative measure of evolution is needed. One such measure is the degree to which creatures improve their efficiency through evolution. This provides not only an objective measure of progress in evolution, but also sheds light on the potential application of synthetic life systems to the problem of the optimization of machine code.

The efficiency of the creature can be indexed in two ways: the size of the genome, and the number of CPU cycles needed to execute one replication. Clearly, smaller genomes can be replicated with less CPU time, however, during evolution, creatures also decrease the ratio of instructions executed in one replication, to genome size. The number of instructions executed per instruction copied, drops substantially.

Figure 4 shows the changes in genome size over a time period of 500 million instructions executed by the system, for eight sets of mutation rates differing by factors of two. Mutation rates are measured in terms of 1 in N individuals being affected by a mutation in each generation. At the highest two sets of rates tested, one and two, either each (one) or one-half (two) of the individuals are hit by mutation in each generation. At these rates the system is unstable. The genomes melt under the heat of the high mutation rates. The community often dies out, although some runs survived the 500 million instruction runs used in this study. The next lower rate, four, yields the highest rate of optimization without the risk

of death of the community. At the five lower mutation rates, 8, 16, 32, 64 and 128, we see successively lower rates of optimization.

Additional replicates were made of the runs at the mutation rate of four (Fig. 5). The replicates differ only in the seed of the random number generator, all other parameters being identical. These runs vary in some details such as whether progress is continuous and gradual, or comes in bursts. Also, each run decreases to a size limit which it can not proceed past even if it is allowed to run much longer. However, different runs reach different plateaus of efficiency. The smallest limiting genome size seen has been 22 instructions, while other runs reached limits of 27 and 30 instructions. Evidently, the system can reach a local optima from which it can not easily evolve to the global optima.

The increase in efficiency of the replicating algorithms is even greater than the decrease in the size of the code. The ancestor is 80 instructions long and requires 839 CPU cycles to replicate. The creature of size 22 only requires 146 CPU cycles to replicate, a 5.75-fold difference in efficiency. The algorithm of one of these creatures is listed in Appendix D.

Although optimization of the algorithm is maximized at the highest mutation rate that does not cause instability, ecological interactions appear to be richer at slightly lower mutation rates. At the rates of eight or 16, we find the diversity of coexisting size classes to be the greatest, and to persist the longest. The smaller size classes tend to be various forms of parasites, thus a diversity of size classes indicates a rich ecology.

An example of even greater optimization is illustrated in Appendix E and discussed above in section 3.1.1.8. Unrolling of the loop results in a loop which uses 18 CPU cycles to copy three instructions, or six CPU cycles executed per instruction copied, compared to 10 for the ancestor. The creature of size 22 also uses six CPU cycles per instruction copied. However, the creature of Appendix E uses three extra CPU cycles per loop to compensate for a separate adaptation that allows it to double its share of CPU time from the global pool (in essence meaning that relatively speaking, it uses only three CPU cycles per instruction copied). Without this compensation it would use only five CPU cycles per instruction copied.

4 SUMMARY

4.1 GENERAL BEHAVIOR OF THE SYSTEM

Once the soup is full of replicating creatures, individuals are initially short lived, generally reproducing only once before dying, thus individuals turn over very rapidly. More slowly, there appear new genotypes of size 80, and then new size classes. There are changes in the genetic composition of each size class, as new mutants appear, some of which increase significantly in frequency, eventually replacing the original genotype. The size classes which dominate the community also change through time, as new size classes appear, some of which competitively exclude sizes present earlier. Once the community becomes diverse, there is a greater variance in the longevity and fecundity of individuals.

In addition to an increase in the raw diversity of genotypes and genome sizes, there is an increase in the ecological diversity. Obligate commensal parasites evolve, which are not capable of self-replication in isolated culture, but which can replicate when cultured

with normal (self-replicating) creatures. These parasites execute some parts of the code of their hosts, but cause them no direct harm, except as competitors. Some potential hosts have evolved immunity to the parasites, and some parasites have evolved to circumvent this immunity.

In addition, facultative hyper-parasites have evolved, which can self-replicate in isolated culture, but when subjected to parasitism, subvert the parasites energy metabolism to augment their own reproduction. Hyper-parasites drive parasites to extinction, resulting in complete domination of the communities. The relatively high degrees of genetic relatedness within the hyper-parasite dominated communities leads to the evolution of sociality in the sense of creatures that can only replicate when they occur in aggregations. These social aggregations are then invaded by hyper-hyper-parasite cheaters.

Mutations and the ensuing replication errors lead to an increasing diversity of sizes and genotypes of self-replicating creatures in the soup. Within the first 100 million instructions of elapsed time, the soup evolves to a state in which about a dozen more-or-less persistent size classes coexist. The relative abundances and specific list of the size classes varies over time. Each size class consists of a number of distinct genotypes which also vary over time.

The rate of evolution increases with the mutation rate until the system becomes unstable, and the community dies at rates above one mutation per four generations. Ecological interactions are richer and more sustained at slightly lower rates, one mutation per eight or 16 generations. At mutation rates of one per four generations, under selection for small sizes, creatures will optimize to a genome size in the 22 to 30 instruction size range within as little as 300 million instructions of elapsed time. Each of these runs will reach a local optima which it evidently cannot escape from, although it may not be the global optima.

4.2 INCREASING COMPLEXITY

The unrolled loop (section 3.1.1.8) is an example of the ability of evolution to produce an increase in complexity, gradually over a long period of time. The interesting thing about the loop unrolling optimization technique is that it requires more complex code. The resulting creature has a genome size of 36, compared to its ancestor of size 80, yet it has packed a much more complex algorithm into less than half the space (Appendix E).

This is a classic example of intricate design in evolution. One wonders how it could have arisen through random bit flips, as every component of the code must be in place in order for the algorithm to function. Yet the code includes a classic mix of apparent intelligent design, and the chaotic hand of evolution. The optimization technique is a very clever one invented by humans, yet it is implemented in a mixed up but functional style that no human would use (unless perhaps very intoxicated).

4.3 EMERGENCE

The “physical” environment presented by the simulator is quite simple, consisting of the energy resource (CPU time) doled out rather uniformly by the time slicer, and memory space which is completely uniform and always available. In light of the nature of the physical environment, the implicit fitness function would presumably favor the evolution of creatures which are able to replicate with less CPU time, and this does in fact occur. However, much of

the evolution in the system consists of the creatures discovering ways to exploit one-another. The creatures invent their own fitness functions through adaptation to their biotic (“living”) environment. These ecological interactions are not programmed into the system, but emerge spontaneously as the creatures discover each other and invent their own games.

In the Tierran world, creatures which initially do not interact, discover means to exploit one another, and in response, means to avoid exploitation. The original fitness landscape of the ancestor consists only of the efficiency parameters of the replication algorithm, in the context of the properties of the reaper and slicer queues. When by chance, genotypes appear that exploit other creatures, selection acts to perfect the mechanisms of exploitation, and mechanisms of defense to that exploitation. The original fitness landscape was based only on adaptations of the organism to its physical environment. The new fitness landscape retains those features, but adds to it adaptations to the biotic environment, the other creatures. Because the fitness landscape includes an ever increasing realm of adaptations to other creatures which are themselves evolving, it can facilitate an auto-catalytic increase in complexity and diversity of organisms.

Evolutionary theory suggests that adaptation to the biotic environment (other organisms) rather than to the physical environment is the primary force driving the auto-catalytic diversification of organisms (Stanley [34]). It is encouraging to discover that the process has already begun in the Tierran world. It is worth noting that the results presented here are based on evolution of the first creature that I designed, written in the first instruction set that I designed. Comparison to the creatures that have evolved shows that the one I designed is not a particularly clever one. Also, the instruction set that the creatures are based on is certainly not very powerful (apart from those special features incorporated to enhance its evolvability). It would appear then that it is rather easy to create life. Evidently, virtual life is out there, waiting for us to provide environments in which it may evolve.

4.4 SYNTHETIC BIOLOGY

One of the most uncanny of evolutionary phenomena is the ecological convergence of biota living on different continents or in different epochs. When a lineage of organisms undergoes an adaptive radiation (diversification), it leads to an array of relatively stable ecological forms. The specific ecological forms are often recognizable from lineage to lineage. For example among dinosaurs, the *Pterosaur*, *Triceratops*, *Tyrannosaurus* and *Ichthyosaur* are ecological parallels respectively, to the bat, rhinoceros, lion and porpoise of modern mammals. Similarly, among modern placental mammals, the gray wolf, flying squirrel, great anteater and common mole are ecological parallels respectively, to the Tasmanian wolf, honey glider, banded anteater and marsupial mole of the marsupial mammals of Australia.

Given these evidently powerful convergent forces, it should perhaps not be surprising that as adaptive radiations proceed among digital organisms, we encounter recognizable ecological forms, in spite of the fundamentally distinct physics and chemistry on which they are based. Ideally, comparisons should be made among organisms of comparable complexity. It may not be appropriate to compare viruses to mammals. Unfortunately, the organic creatures most comparable to digital organisms, the RNA creatures, are no longer with us. Since digital organisms are being compared to modern organic creatures of much greater

complexity, ecological comparisons must be made in the broadest of terms.

Trained biologists will tend to view synthetic life in the same terms that they have come to know organic life. Having been trained as an ecologist and evolutionist, I have seen in my synthetic communities, many of the ecological and evolutionary properties that are well known from natural communities. Biologists trained in other specialties will likely observe other familiar properties. It seems that what we see is what we know. It is likely to take longer before we appreciate the unique properties of these new life forms.

4.5 FUTURE DIRECTIONS

For the immediate future, my research will involve three main thrusts:

Computational: The computational issue is how to design a computer architecture and operating system that will support the natural evolution of *machine code*. Von Neumann style machine codes are considered to be brittle, in the sense that they are not robust to the genetic operations of mutation and recombination. Any random change in a program is almost 100% certain to break the program.

The most significant accomplishment of my work to date is finding a way to overcome this brittleness with only a slight modification of standard machine codes. However, this success was achieved in the first attempt. Therefore it is not known what features are essential for evolvability, and I certainly do not know what is the optimal architecture.

The primary computational objective then is to experiment with a large number of variations on the successful architecture in order to find the optimal balance of computational power and evolvability. This work has begun but needs to be scaled up.

There are however, other important computational goals. We must experiment with artificial selection on application programs. So far all work has involved natural selection on “wild” algorithms that do no useful work. I want to develop an analog to genetic engineering, in which application codes are inserted into the genomes of digital organisms and evolved to greater optimality or new functionality.

It should be possible to develop cross-assemblers between Tierran architectures and real assembler languages. Application code written and compiled to run on real machines could be cross-assembled into the new Tierran languages. Each procedure could then be inserted into the genome of a creature. Creatures could be rewarded with CPU time in proportion to the efficacy and efficiency of the evolving inserted code. In this way, artificial selection would lead to the optimization of the inserted code, which could then be cross-assembled back into the real machine code.

If artificial selection of application programs proved to be practical, it would be worthwhile to render the best Tierran virtual instruction sets in silicon, thereby greatly accelerating the process. At present, maximum optimization can be achieved in a few hours of running the Tierran virtual computer. If a real computer were based on the architectural principals of the Tierran computer, the speed would be multiplied by about two orders of magnitude. If the real machine were massively parallel, there could be additional gains of five to six orders of magnitude. If machine code could evolve that quickly, then there is the possibility

of using it as a generative process in addition to an optimization procedure. There may also be some potential application in the areas of machine learning or adaptive programming.

Another long term objective is to use digital organisms evolving freely or under artificial selection, as a source of new paradigms for the programming of massively parallel machines. The virtual computer that supports digital evolution is a parallel machine of the MIMD (multiple instruction multiple data) type. One of the biggest problems facing computer science today is the development of techniques of parallel programming. Digital organisms program themselves, using evolution. They have discovered on their own, known programming techniques such as unrolling loops. They will discover techniques that are naturally efficient on parallel machines, and we should be able to learn from their innovations.

The kinds of ecological interactions already observed in digital communities could in another light, be viewed as optimization techniques for parallel programming (e.g., the sharing of code fragments). However, these interactions evolve in a "jungle"-like environment where most interactions are of an adversarial nature. When evolving large parallel application programs, the most viable model would be a multi-cellular one, where many cells would cooperate on a common problem. A multi-cellular model is under development. In the end, evolution may prove to be the best method of programming massively parallel machines.

Biological: I plan to move the biological model ahead of its present state. This will primarily involve the incorporation of facilities to support diploidy, organized sexuality, and multi-cellularity. The methods for these advances have already been conceptually worked out, and the implementation has begun. When these improvements are made, the long term biological goals are to use the model to test ecological and evolutionary theory, in such areas as: the evolution of sex, selection across hierarchical levels, and factors affecting diversity of ecological communities. It is hoped that it will be possible to engineer the system up to a condition analagous to the threshold of the Cambrian explosion of diversity, and then just allow the complexity and diversity of the digital system to explode spontaneously.

Educational: I wish to distribute both source and executables for use as an educational and research tool. However, some additional work is needed to make the program fully portable and to provide a user friendly graphic user interface. This work is underway at a slow pace.

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Figure 1. Metabolic flow chart for the ancestor, parasite, hyper-parasite, and their interactions: ax, bx and cx refer to CPU registers where location and size information are stored. [ax] and [bx] refer to locations in the soup indicated by the values in the ax and bx registers. Patterns such as 1101 are complementary templates used for addressing. Arrows outside of boxes indicate jumps in the flow of execution of the programs. The dotted-line arrows indicate flow of execution between creatures. The parasite lacks the copy procedure, however, if it is within the search limit of the copy procedure of a host, it can locate, call and execute that procedure, thereby obtaining the information needed to complete its replication. The host is not adversely affected by this informational parasitism, except through competition with the parasite, which is a superior competitor. Note that the parasite calls the copy procedure of its host with the expectation that control will return to the parasite when the copy procedure returns. However, the hyper-parasite jumps out of the copy procedure rather than returning, thereby seizing control from the parasite. It then proceeds to reset the CPU registers of the parasite with the location and size of the hyper-parasite, causing the parasite to replicate the hyper-parasite genome thereafter.

ANCESTOR

1111

```

self-exam
find 0000 (start) -> bx
find 0001 (end) -> ax
calculate size -> cx

```

1101

```

reproduction loop
allocate daughter -> ax
call 0011 (copy procedure)
cell division
jump 0010

```

1100

```

copy procedure
save registers to stack
1010
move [bx] -> [ax]
decrement cx
if cx == 0 jump 0100
increment ax & bx
jump 0101
1011
restore registers
return

```

1110

PARASITE

1111

```

self-exam
find 0000 (start) -> bx
find 0001 (end) -> ax
calculate size -> cx

```

1101

```

reproduction loop
allocate daughter -> ax
call 0011 (copy procedure)
cell division
jump 0010

```

1110

HYPER-PARASITE

1111

```

self-exam
find 0000 (start) -> bx
find 0001 (end) -> ax
calculate size -> cx

```

```

reproduction loop
allocate daughter -> ax
call 0011 (copy procedure)
cell division
jumpb 0000

```

1100

```

copy procedure
1010
move [bx] -> [ax]
decrement cx
if cx == 0 jumpb 1100
increment ax & bx
jumpb 0101

```

1110

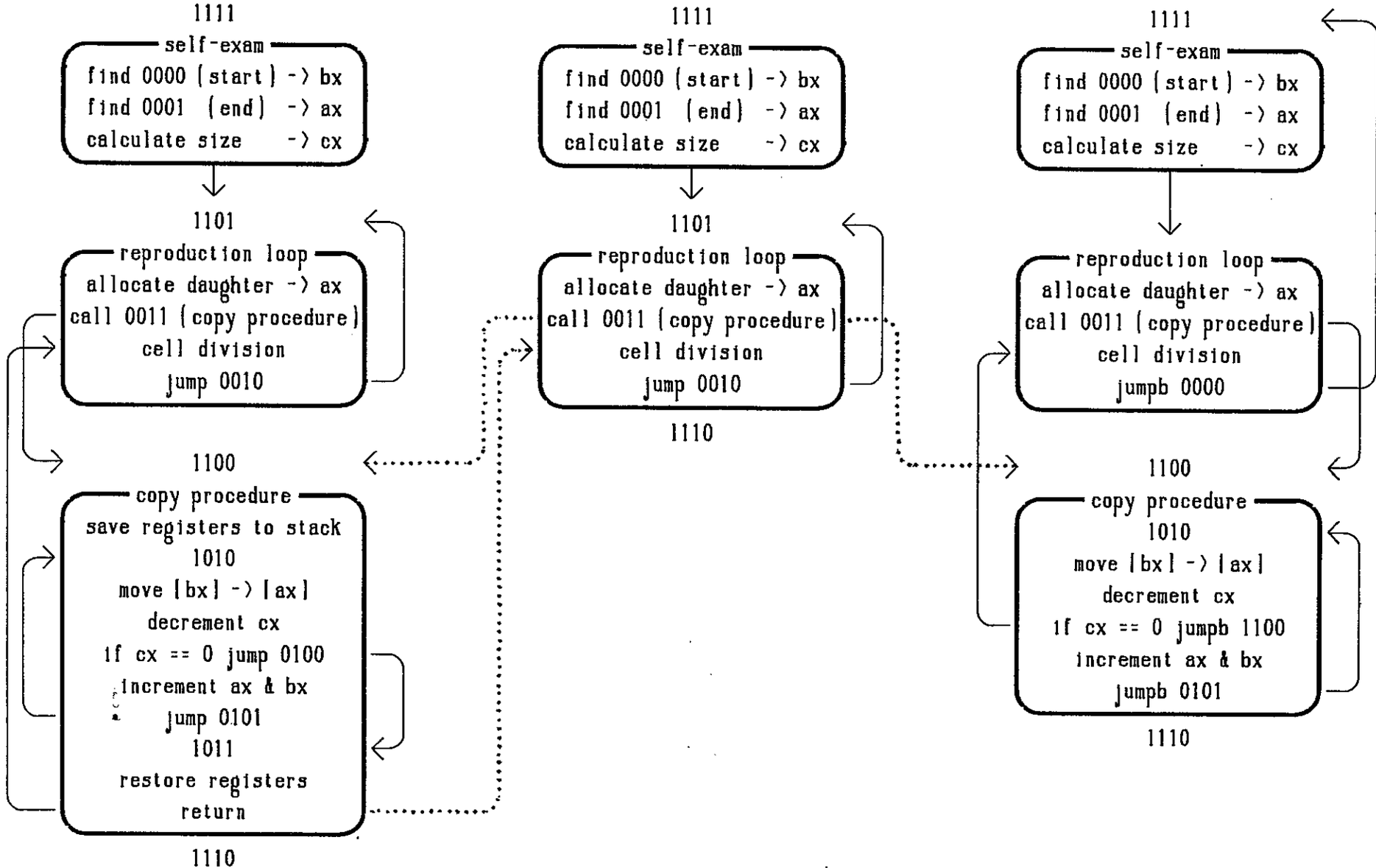
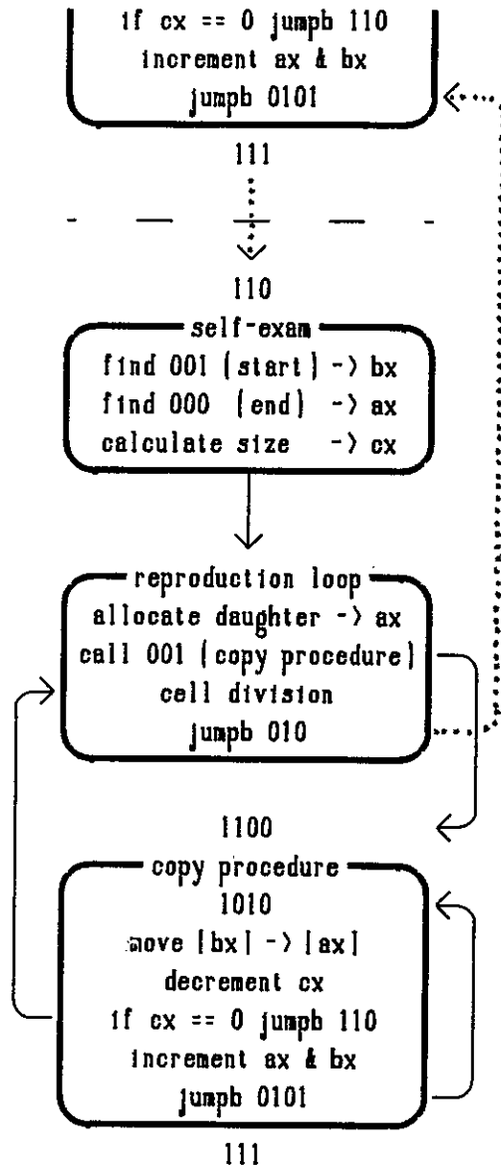


Figure 2. Metabolic flow chart for social hyper-parasites, their associated hyper-hyper-parasite cheaters, and their interactions. Symbols are as described for Fig. 1. Horizontal dashed lines indicate the boundaries between individual creatures. On both the left and right, above the dashed line at the top of the figure is the lowermost fragment of a social-hyper-parasite. Note (on the left) that neighboring social hyper-parasites cooperate in returning the flow of execution to the beginning of the creature for self-re-examination. Execution jumps back to the end of the creature above, but then falls off the end of the creature without executing any instructions of consequence, and enters the top of the creature below. On the right, a cheater is inserted between the two social-hyper-parasites. The cheater captures control of execution when it passes between the social individuals. It sets the CPU registers with its own location and size, and then skips over the self-examination step when it returns control of execution to the social creature below.

SOCIAL HYPER-PARASITES



SOCIAL HYPER-PARASITES
AND CHEATER

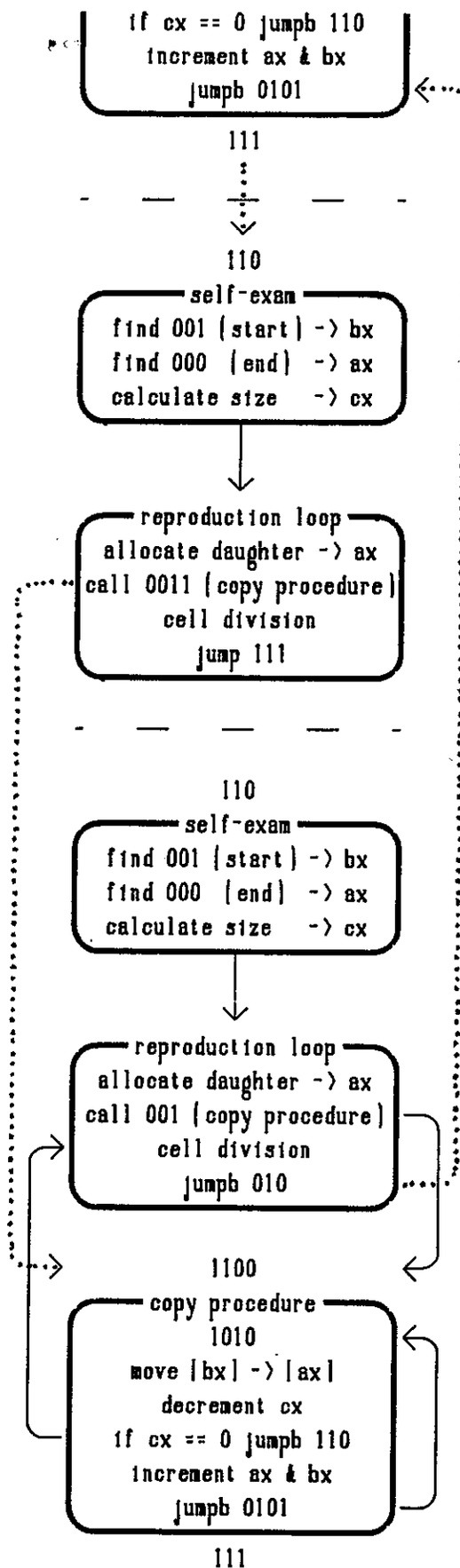


Figure 3. Metabolic flow chart for obligate symbionts and their interactions. Symbols are as described for Fig. 1. Neither creature is able to self-replicate in isolation. However, when cultured together, each is able to replicate by using information provided by the other.

SYMBIONTS

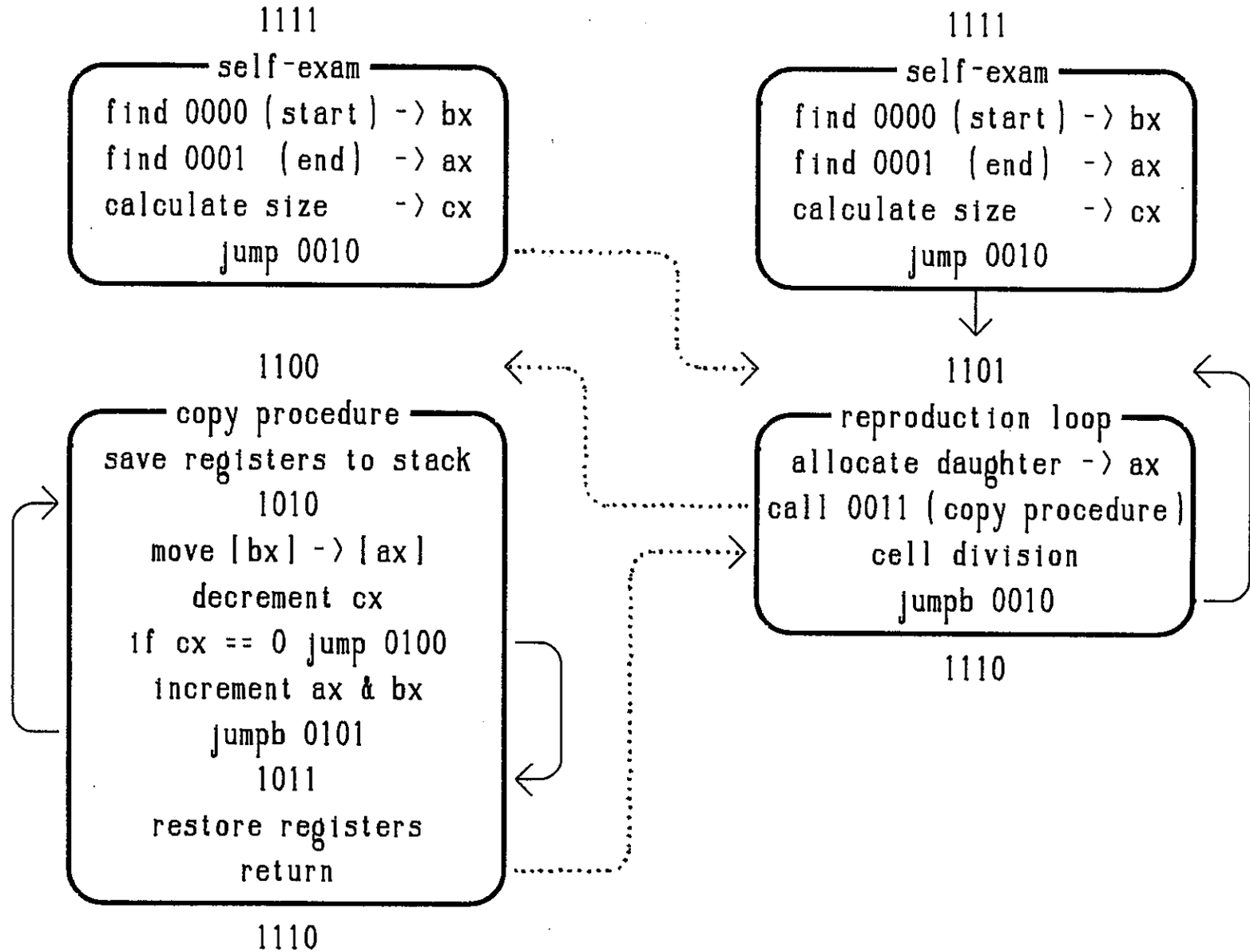
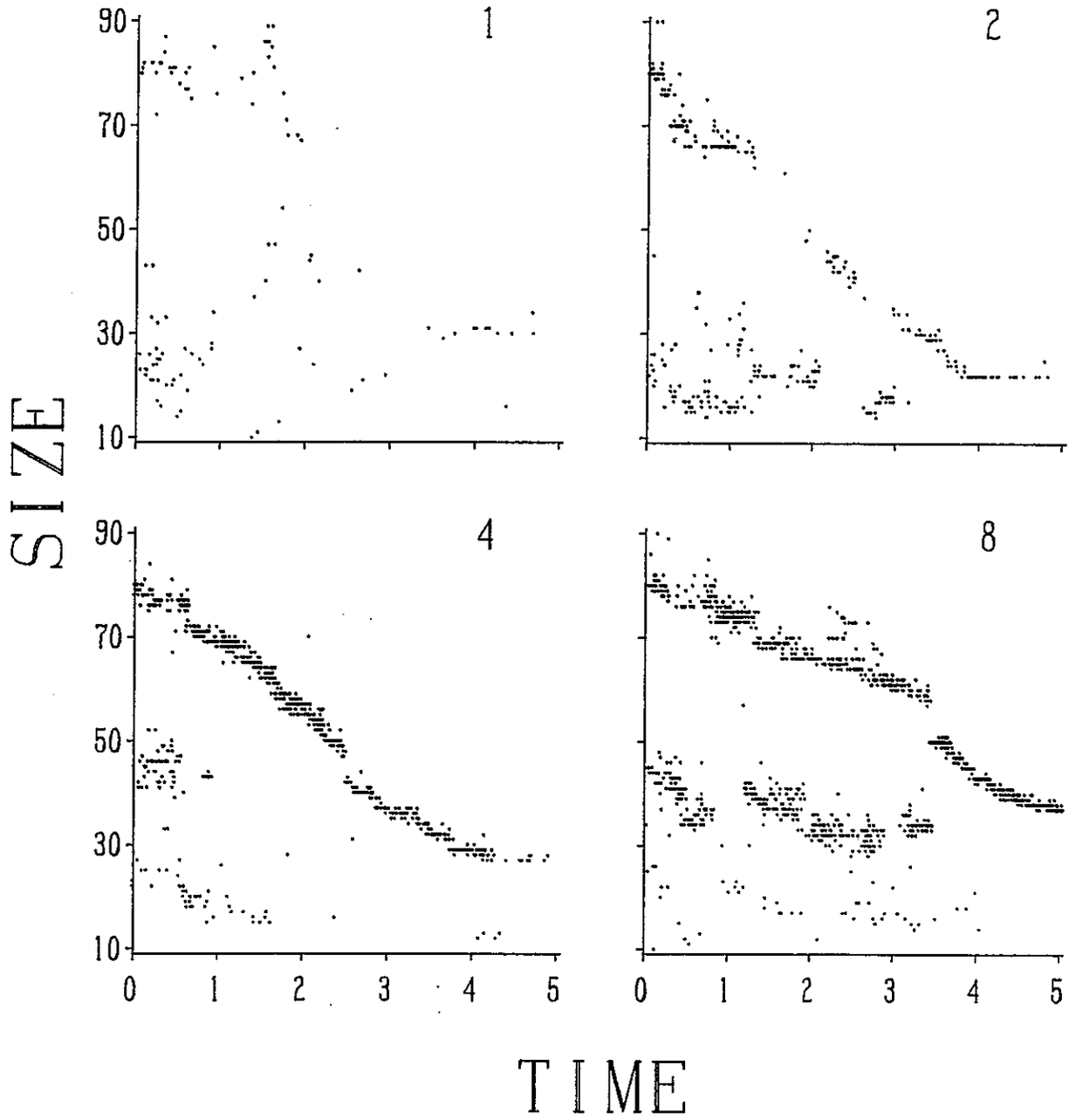


Figure 4. Evolutionary optimization at eight sets of mutation rates. In each run, the three mutation rates: move mutations (copy error), flaws and background mutations (cosmic rays) are set relative to the generation time. In each case, the background mutation rate is the lowest, affecting a cell once in twice as many generations as the move mutation rate. The flaw rate is intermediate, affecting a cell once in 1.5 times as many generations as the move mutation rate. For example in one run, the move mutation will affect a cell line on the average once every 4 generations, the flaw will occur once every 6 generations, and the background mutation once every 8 generations. The horizontal axis shows elapsed time in hundreds of millions of instructions executed by the system. The vertical axis shows genome size in instructions. Each point indicates the first appearance of a new genotype which crossed the abundance thresholds of either 2% of the population of cells in the soup, or occupation of 2% of the memory. The number of generations per move mutation is indicated by a number in the upper right hand corner of each graph.



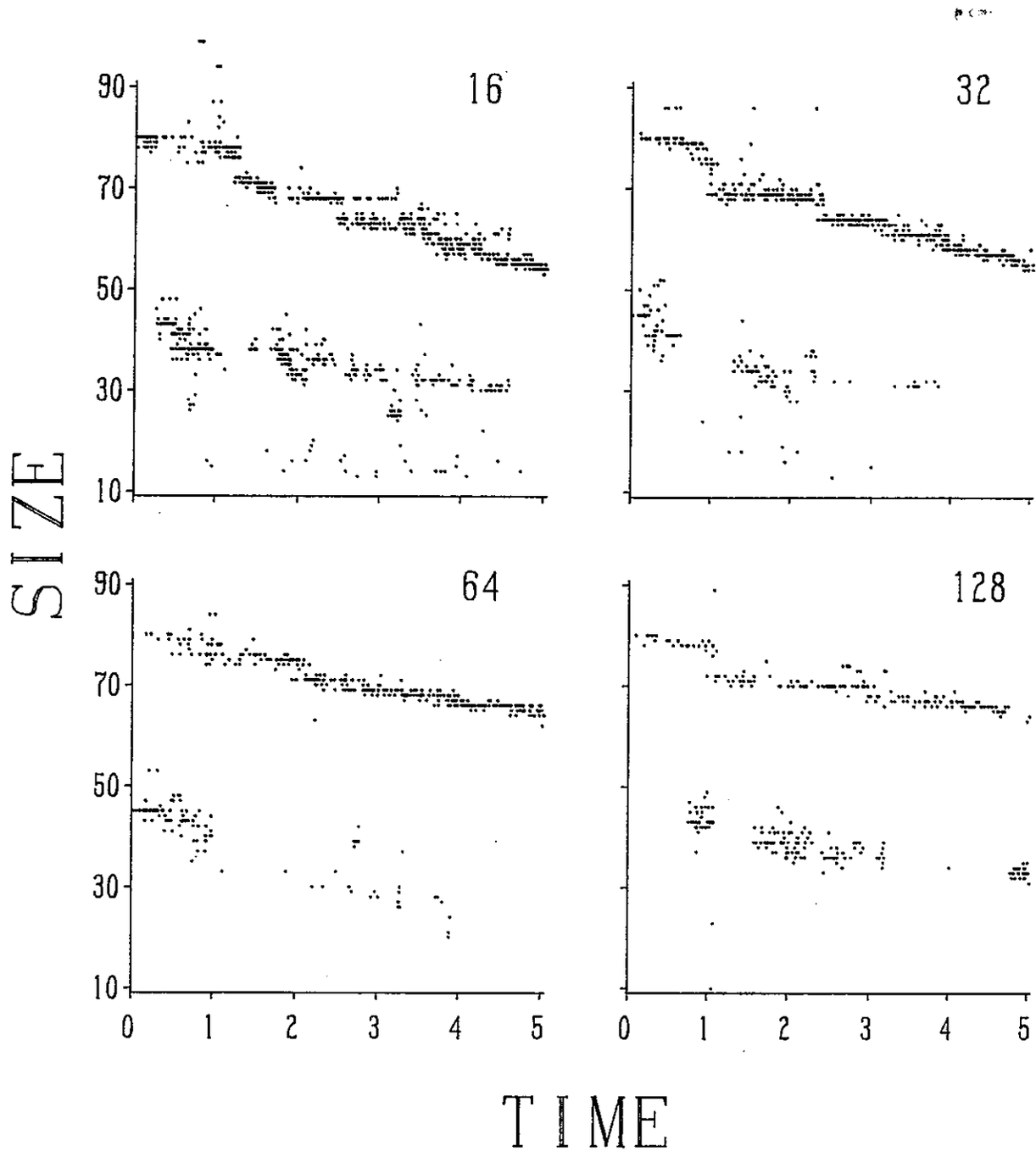


Figure 5. Variation in evolutionary optimization under constant conditions. Based on a mutation rate of four generations per move mutation, all other parameters as in Fig. 4. The plots are otherwise as described for Fig. 4.

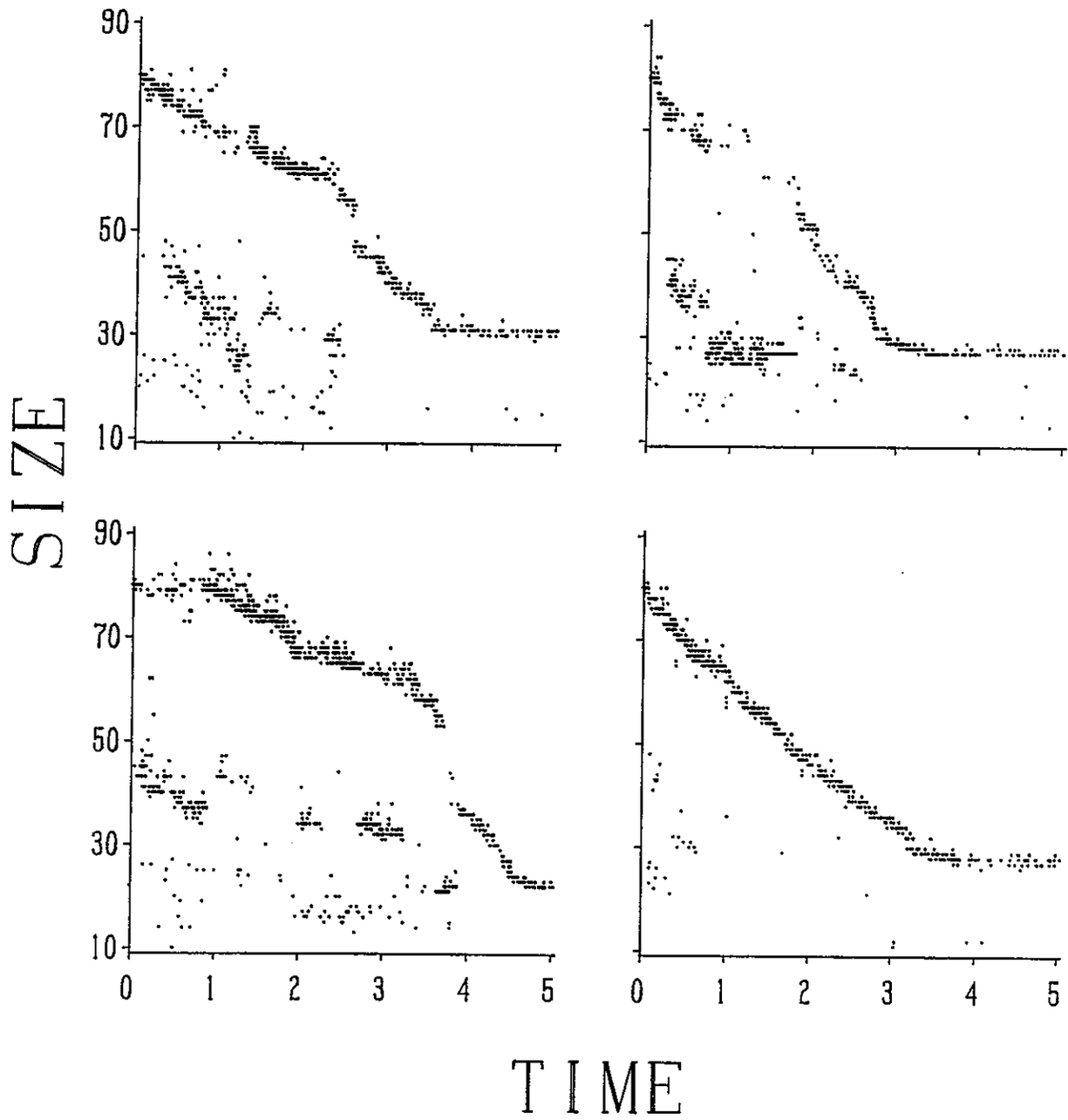


Table 1: Genebank. Table of numbers of size classes in the genebank. Left column is size class, right column is number of self-replicating genotypes of that size class. 305 sizes, 29,275 genotypes.

0034 1	0092 362	0150 2	0205 5	0418 1	5213 2
0041 2	0093 261	0151 1	0207 3	0442 10	5229 4
0043 12	0094 241	0152 2	0208 2	0443 1	5254 1
0044 7	0095 211	0153 1	0209 1	0444 61	5888 36
0045 191	0096 232	0154 2	0210 9	0445 1	5988 1
0046 7	0097 173	0155 3	0211 4	0456 2	6006 2
0047 5	0098 92	0156 77	0212 4	0465 6	6014 1
0048 4	0099 117	0157 270	0213 5	0472 6	6330 1
0049 8	0100 77	0158 938	0214 47	0483 1	6529 1
0050 13	0101 62	0159 836	0218 1	0484 8	6640 1
0051 2	0102 62	0160 3229	0219 1	0485 3	6901 5
0052 11	0103 27	0161 1417	0220 2	0486 9	6971 1
0053 4	0104 25	0162 174	0223 3	0487 2	7158 2
0054 2	0105 28	0163 187	0226 2	0493 2	7293 3
0055 2	0106 19	0164 46	0227 7	0511 2	7331 1
0056 4	0107 3	0165 183	0231 1	0513 1	7422 70
0057 1	0108 8	0166 81	0232 1	0519 1	7458 1
0058 8	0109 2	0167 71	0236 1	0522 6	7460 7
0059 8	0110 8	0168 9	0238 1	0553 1	7488 1
0060 3	0111 71	0169 15	0240 3	0568 6	7598 1
0061 1	0112 19	0170 99	0241 1	0578 1	7627 63
0062 2	0113 10	0171 40	0242 1	0581 3	7695 1
0063 2	0114 3	0172 44	0250 1	0582 1	7733 1
0064 1	0115 3	0173 34	0251 1	0600 1	7768 2
0065 4	0116 5	0174 15	0260 2	0683 1	7860 25
0066 1	0117 3	0175 22	0261 1	0689 1	7912 1
0067 1	0118 1	0176 137	0265 2	0757 6	8082 3
0068 2	0119 3	0177 13	0268 1	0804 2	8340 1
0069 1	0120 2	0178 3	0269 1	0813 1	8366 1
0070 7	0121 60	0179 1	0284 16	0881 6	8405 5
0071 5	0122 9	0180 16	0306 1	0888 1	8406 2
0072 17	0123 3	0181 5	0312 1	0940 2	8649 2
0073 2	0124 11	0182 27	0314 1	1006 6	8750 1
0074 80	0125 6	0184 3	0316 2	1016 1	8951 1
0075 56	0126 11	0185 21	0318 3	1077 5	8978 3
0076 21	0127 1	0186 9	0319 2	1116 1	9011 3
0077 28	0130 3	0187 3	0320 23	1186 1	9507 3
0078 409	0131 2	0188 11	0321 5	1294 7	9564 3
0079 850	0132 5	0190 20	0322 21	1322 7	9612 1
0080 7399	0133 2	0192 12	0330 1	1335 1	9968 1
0081 590	0134 7	0193 4	0342 5	1365 11	10259 31

0082 384	0135 1	0194 4	0343 1	1631 1	10676 1
0083 886	0136 1	0195 11	0351 1	1645 3	11366 5
0084 1672	0137 1	0196 19	0352 3	2266 1	11900 1
0085 1531	0138 1	0197 2	0386 1	2615 2	12212 2
0086 901	0139 2	0198 3	0388 2	2617 9	15717 3
0087 944	0141 6	0199 35	0401 3	2671 7	16355 1
0088 517	0143 1	0200 1	0407 1	3069 3	17356 3
0089 449	0144 4	0201 84	0411 22	4241 1	18532 1
0090 543	0146 1	0203 1	0412 3	5101 15	23134 14
0091 354	0149 1	0204 1	0416 1	5157 9	

APPENDIX A

Structure definition to implement the Tierra virtual CPU. The complete source code for the Tierra Simulator can be obtained by contacting the author by email.

```
struct cpu { /* structure for registers of virtual cpu */
    int  ax; /* address register */
    int  bx; /* address register */
    int  cx; /* numerical register */
    int  dx; /* numerical register */
    char fl; /* flag */
    char sp; /* stack pointer */
    int  st[10]; /* stack */
    int  ip; /* instruction pointer */
};
```

APPENDIX B

Abbreviated code for implementing the CPU cycle of the Tierra Simulator.

```
void main(void)
{  get_soup();
   life();
   write_soup();
}

void life(void) /* doles out time slices and death */
{  while(inst_exec_c < alive) /* control the length of the run */
    {  time_slice(this_slice); /* this_slice is current cell in queue */
       incr_slice_queue(); /* increment this_slice to next cell in queue */
       while(free_mem_current < free_mem_prop * soup_size)
           reaper(); /* if memory is full to threshold, reap some cells */
    }
}

void time_slice(int  ci)
{  Pcells  ce; /* pointer to the array of cell structures */
   char    i; /* instruction from soup */
   int     di; /* decoded instruction */
   int     j, size_slice;
   ce = cells + ci;
   for(j = 0; j < size_slice; j++)
   {  i = fetch(ce->c.ip); /* fetch instruction from soup, at address ip */
      di = decode(i); /* decode the fetched instruction */
      execute(di, ci); /* execute the decoded instruction */
      increment_ip(di, ce); /* move instruction pointer to next instruction */
   }
```

```

        system_work(); /* opportunity to extract information */
    }
}

void execute(int di, int ci)
{
    switch(di)
    {
        case 0x00: nop_0(ci);      break; /* no operation */
        case 0x01: nop_1(ci);      break; /* no operation */
        case 0x02: orl(ci);        break; /* flip low order bit of cx, cx ^= 1 */
        case 0x03: shl(ci);        break; /* shift left cx register, cx <<= 1 */
        case 0x04: zero(ci);       break; /* set cx register to zero, cx = 0 */
        case 0x05: if_cz(ci);      break; /* if cx==0 execute next instruction */
        case 0x06: sub_ab(ci);     break; /* subtract bx from ax, cx = ax - bx */
        case 0x07: sub_ac(ci);     break; /* subtract cx from ax, ax = ax - cx */
        case 0x08: inc_a(ci);      break; /* increment ax, ax = ax + 1 */
        case 0x09: inc_b(ci);      break; /* increment bx, bx = bx + 1 */
        case 0x0a: dec_c(ci);      break; /* decrement cx, cx = cx - 1 */
        case 0x0b: inc_c(ci);      break; /* increment cx, cx = cx + 1 */
        case 0x0c: push_ax(ci);    break; /* push ax on stack */
        case 0x0d: push_bx(ci);    break; /* push bx on stack */
        case 0x0e: push_cx(ci);    break; /* push cx on stack */
        case 0x0f: push_dx(ci);    break; /* push dx on stack */
        case 0x10: pop_ax(ci);     break; /* pop top of stack into ax */
        case 0x11: pop_bx(ci);     break; /* pop top of stack into bx */
        case 0x12: pop_cx(ci);     break; /* pop top of stack into cx */
        case 0x13: pop_dx(ci);     break; /* pop top of stack into dx */
        case 0x14: jmp(ci);        break; /* move ip to template */
        case 0x15: jmpb(ci);       break; /* move ip backward to template */
        case 0x16: call(ci);       break; /* call a procedure */
        case 0x17: ret(ci);        break; /* return from a procedure */
        case 0x18: mov_cd(ci);     break; /* move cx to dx, dx = cx */
        case 0x19: mov_ab(ci);     break; /* move ax to bx, bx = ax */
        case 0x1a: mov_iab(ci);    break; /* move instruction at address in bx
                                     to address in ax */
        case 0x1b: adr(ci);        break; /* address of nearest template to ax */
        case 0x1c: adrb(ci);       break; /* search backward for template */
        case 0x1d: adrf(ci);       break; /* search forward for template */
        case 0x1e: mal(ci);        break; /* allocate memory for daughter cell */
        case 0x1f: divide(ci);     break; /* cell division */
    }
    inst_exec_c++;
}
}

```

APPENDIX C

Assembler source code for the ancestral creature.

```
genotype: 80 aaa  origin: 1-1-1990  00:00:00:00  ancestor
parent genotype: human
1st_daughter:  flags: 0  inst: 839  mov_daught: 80
2nd_daughter:  flags: 0  inst: 813  mov_daught: 80

nop_1    ; 01    0 beginning template
nop_1    ; 01    1 beginning template
nop_1    ; 01    2 beginning template
nop_1    ; 01    3 beginning template
zero     ; 04    4 put zero in cx
orl      ; 02    5 put 1 in first bit of cx
shl      ; 03    6 shift left cx
shl      ; 03    7 shift left cx, now cx = 4
          ;      ax =                bx =
          ;      cx = template size  dx =
mov_cd   ; 18    8 move template size to dx
          ;      ax =                bx =
          ;      cx = template size  dx = template size
adrb     ; 1c    9 get (backward) address of beginning template
nop_0    ; 00   10 compliment to beginning template
nop_0    ; 00   11 compliment to beginning template
nop_0    ; 00   12 compliment to beginning template
nop_0    ; 00   13 compliment to beginning template
          ;      ax = start of mother + 4  bx =
          ;      cx = template size        dx = template size
sub_ac   ; 07   14 subtract cx from ax
          ;      ax = start of mother    bx =
          ;      cx = template size      dx = template size
mov_ab   ; 19   15 move start address to bx
          ;      ax = start of mother    bx = start of mother
          ;      cx = template size      dx = template size
adrf     ; 1d   16 get (forward) address of end template
nop_0    ; 00   17 compliment to end template
nop_0    ; 00   18 compliment to end template
nop_0    ; 00   19 compliment to end template
nop_1    ; 01   20 compliment to end template
          ;      ax = end of mother    bx = start of mother
          ;      cx = template size    dx = template size
inc_a    ; 08   21 to include dummy statement to separate creatures
sub_ab   ; 06   22 subtract start address from end address to get size
          ;      ax = end of mother    bx = start of mother
          ;      cx = size of mother   dx = template size
nop_1    ; 01   23 reproduction loop template
```

```

nop_1 ; 01 24 reproduction loop template
nop_0 ; 00 25 reproduction loop template
nop_1 ; 01 26 reproduction loop template
mal   ; 1e 27 allocate memory for daughter cell, address to ax
      ;    ax = start of daughter    bx = start of mother
      ;    cx = size of mother       dx = template size
call  ; 16 28 call template below (copy procedure)
nop_0 ; 00 29 copy procedure compliment
nop_0 ; 00 30 copy procedure compliment
nop_1 ; 01 31 copy procedure compliment
nop_1 ; 01 32 copy procedure compliment
divide ; 1f 33 create independent daughter cell
jmp    ; 14 34 jump to template below (reproduction loop, above)
nop_0 ; 00 35 reproduction loop compliment
nop_0 ; 00 36 reproduction loop compliment
nop_1 ; 01 37 reproduction loop compliment
nop_0 ; 00 38 reproduction loop compliment
if_cz ; 05 39 this is a dummy instruction to separate templates
      ;    begin copy procedure
nop_1 ; 01 40 copy procedure template
nop_1 ; 01 41 copy procedure template
nop_0 ; 00 42 copy procedure template
nop_0 ; 00 43 copy procedure template
push_ax ; 0c 44 push ax onto stack
push_bx ; 0d 45 push bx onto stack
push_cx ; 0e 46 push cx onto stack
nop_1 ; 01 47 copy loop template
nop_0 ; 00 48 copy loop template
nop_1 ; 01 49 copy loop template
nop_0 ; 00 50 copy loop template
mov_iab ; 1a 51 move contents of [bx] to [ax]
dec_c   ; 0a 52 decrement cx
if_cz   ; 05 53 if cx == 0 perform next instruction, otherwise skip it
jmp     ; 14 54 jump to template below (copy procedure exit)
nop_0   ; 00 55 copy procedure exit compliment
nop_1   ; 01 56 copy procedure exit compliment
nop_0   ; 00 57 copy procedure exit compliment
nop_0   ; 00 58 copy procedure exit compliment
inc_a   ; 08 59 increment ax
inc_b   ; 09 60 increment bx
jmp     ; 14 61 jump to template below (copy loop)
nop_0   ; 00 62 copy loop compliment
nop_1   ; 01 63 copy loop compliment
nop_0   ; 00 64 copy loop compliment
nop_1   ; 01 65 copy loop compliment
if_cz   ; 05 66 this is a dummy instruction, to separate templates

```

```
nop_1 ; 01 67 copy procedure exit template
nop_0 ; 00 68 copy procedure exit template
nop_1 ; 01 69 copy procedure exit template
nop_1 ; 01 70 copy procedure exit template
pop_cx ; 12 71 pop cx off stack
pop_bx ; 11 72 pop bx off stack
pop_ax ; 10 73 pop ax off stack
ret ; 17 74 return from copy procedure
nop_1 ; 01 75 end template
nop_1 ; 01 76 end template
nop_1 ; 01 77 end template
nop_0 ; 00 78 end template
if_cz ; 05 79 dummy statement to separate creatures
```

APPENDIX D

Assembler source code for the smallest self-replicating creature.

```
genotype: 0022abn parent genotype: 0022aak
1st_daughter: flags: 1 inst: 146 mov_daught: 22 breed_true: 1
2nd_daughter: flags: 0 inst: 142 mov_daught: 22 breed_true: 1
InstExecC: 437 InstExec: 625954 origin: 662865379 Wed Jan 2 20:16:19 1991
MaxPropPop: 0.1231 MaxPropInst: 0.0568
```

```
nop_0 ; 00 0
adrb ; 1c 1 find beginning
nop_1 ; 01 2
divide ; 1f 3 fails the first time it is executed
sub_ac ; 07 4
mov_ab ; 19 5
adrf ; 1d 6 find end
nop_0 ; 00 7
inc_a ; 08 8 to include final dummy statement
sub_ab ; 06 9 calculate size
mal ; 1e 10
push_bx ; 0d 11 save beginning address on stack in order to 'return' there
nop_0 ; 00 12 top of copy loop
mov_iab ; 1a 13
dec_c ; 0a 14
if_cz ; 05 15
ret ; 17 16 jump to beginning, address saved on stack
inc_a ; 08 17
inc_b ; 09 18
jmpb ; 15 19 bottom of copy loop (6 instructions executed per loop)
nop_1 ; 01 20
mov_iab ; 1a 21 dummy statement to terminate template
```

APPENDIX E

Assembler code for the central copy loop of the ancestor (80aaa) and descendant after fifteen billion instructions (72etq). Within the loop, the ancestor does each of the following operations once: copy instruction (51), decrement cx (52), increment ax (59) and increment bx (60). The descendant performs each of the following operations three times within the loop: copy instruction (15, 22, 26), increment ax (20, 24, 31) and increment bx (21, 25, 32). The decrement cx operation occurs five times within the loop (16, 17, 19, 23, 27). Instruction 28 flips the low order bit of the cx register. Whenever this latter instruction is reached, the value of the low order bit is one, so this amounts to a sixth instance of decrement cx. This means that there are two decrements for every increment. The reason for this is related to another adaptation of this creature. When it calculates its size, it shifts left (12) before allocating space for the daughter (13). This has the effect of allocating twice as much space as is actually needed to accommodate the genome. The genome of the creature is 36 instructions long, but it allocates a space of 72 instructions. This occurred in an environment where the slice size was set equal to the size of the cell. In this way the creatures were able to garner twice as much energy. However, they had to compliment this change by doubling the number of decrements in the loop.

```

nop_1    ; 01 47 copy loop template      COPY LOOP OF 80AAA
nop_0    ; 00 48 copy loop template
nop_1    ; 01 49 copy loop template
nop_0    ; 00 50 copy loop template
mov_iab  ; 1a 51 move contents of [bx] to [ax] (copy instruction)
dec_c    ; 0a 52 decrement cx
if_cz    ; 05 53 if cx = 0 perform next instruction, otherwise skip it
jmp      ; 14 54 jump to template below (copy procedure exit)
nop_0    ; 00 55 copy procedure exit compliment
nop_1    ; 01 56 copy procedure exit compliment
nop_0    ; 00 57 copy procedure exit compliment
nop_0    ; 00 58 copy procedure exit compliment
inc_a    ; 08 59 increment ax (point to next instruction of daughter)
inc_b    ; 09 60 increment bx (point to next instruction of mother)
jmp      ; 14 61 jump to template below (copy loop)
nop_0    ; 00 62 copy loop compliment
nop_1    ; 01 63 copy loop compliment
nop_0    ; 00 64 copy loop compliment
nop_1    ; 01 65 copy loop compliment (10 instructions executed per loop)

```

```

shl      ; 000 03 12 shift left cx      COPY LOOP OF 72ETQ
mal      ; 000 1e 13 allocate daughter cell
nop_0    ; 000 00 14 top of loop
mov_iab  ; 000 1a 15 copy instruction
dec_c    ; 000 0a 16 decrement cx
dec_c    ; 000 0a 17 decrement cx
jmpb     ; 000 15 18 junk
dec_c    ; 000 0a 19 decrement cx
inc_a    ; 000 08 20 increment ax
inc_b    ; 000 09 21 increment bx
mov_iab  ; 000 1a 22 copy instruction
dec_c    ; 000 0a 23 decrement cx
inc_a    ; 000 08 24 increment ax
inc_b    ; 000 09 25 increment bx
mov_iab  ; 000 1a 26 copy instruction
dec_c    ; 000 0a 27 decrement cx
orl      ; 000 02 28 flip low order bit of cx
if_cz    ; 000 05 29 if cx == 0 do next instruction
ret      ; 000 17 30 exit loop
inc_a    ; 000 08 31 increment ax
inc_b    ; 000 09 32 increment bx
jmpb     ; 000 15 33 go to top of loop (6 instructions per copy)
nop_1    ; 000 01 34 bottom of loop (18 instructions executed per loop)

```