Convergence Properties of Solutions of a Length-Structured Density-Dependent Model for Fish*

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Abstract

We numerically study solutions of a length-structured matrix model for fish populations in which the probability that a fish grows into the next length class is a decreasing nonlinear function of the total biomass of the population. We make conjectures about the convergence properties of solutions, and give numerical simulations which support these conjectures. We also study the distribution of biomass in the different age classes as a function of the total biomass.

Keywords: Population dynamics, Nonlinearity, Stability, Equilibria, Fish populations

1 Introduction and Background

In this paper we numerically analyze a discrete-time, nonlinear length-based model for a fish population. This model has been mathematically studied (with proofs) in Callahan, et. al. [1], with a restrictive condition on the survival probabilities. In this paper we numerically study the solutions of this model when this condition on the survival probabilities is removed. We make conjectures that are stronger than the results proved in [1], and give numerical examples that make these conjectures plausible.

In [1] references are given for papers that give fish population models which incorporate length structure. These papers give justifications for incorporating length structure into a population model; for instance, "length structure is included because most population dynamics processes are related to size rather than the age of the fish" [4]. It is known that many fish species are less likely to grow when their habitat is crowded. Thus it makes sense to include in

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this model the condition that the probability that a fish moves to a larger length class in one time step is a decreasing function of the total amount of biomass in the habitat. This function should be positive, should take the value 1 when there is no biomass (since with no crowding fish are guaranteed to grow), and should approach 0 as the biomass gets infinite (since with "infinite" crowding fish can't grow). Incorporating such a nonlinearity into the model means that when the population gets too big, there's pressure to keep it down, and when the population is small, it grows unimpeded. Thus it is plausible that there are circumstances under which we expect the population to converge. In [1] it is proved that under certain conditions the biomass converges uniformly (that is, independent of the initial conditions), and we conjecture that in fact under those circumstances the population converges uniformly, even when the survival hypotheses used in [1] are not satisfied. We will back up this, and other conjectures, with numerical simulations.

The paper is organized as follows. In Sect. 2 we describe the discrete-time nonlinear length-structured model which we are studying. In this model the probability that a fish will grow into the next length class in one time step is a nonlinear function of the total biomass of the population. In Sect. 3 we make conjectures about the convergence properties of solutions to the model. In Sect. 4 we give numerical results that are consistent with these conjectures. We use as a case study an invasive White Perch species studied in [2, 1]. For the purposes of angling, a long-term population which is dominated by fish in the larger length classes is desirable, and in Sect. 5 we make connections between the distribution of biomass and the survival rates and total biomass.

Notation: Denote transpose by a T superscript, a row vector by \vec{v}^T , $[v_1, v_1, \ldots, v_n]$ or $[v_j]$. Denote a column vector by \vec{v} , $[v_1, v_1, \ldots, v_n]^T$ or $[v_j]^T$. The spectral radius of a square matrix A is denoted by $\rho(A)$, and is the modulus of the largest eigenvalue of A. The 1-norm of a vector \vec{v} is denoted by $||\vec{v}||$, and is the sum of the absolute values of the entries.

2 Mathematical Model

This model was introduced in [1], and is similar to models in [3, 5], and is related to age-structured models in [2, 6]. We start with n length classes of reproductively viable fish and denote the population in each class after t time steps by $P_1(t), P_2(t), \ldots, P_n(t)$. We define the population vector to be $\vec{P}(t) = [P_0(t), P_1(t), \ldots, P_n(t)]^T \in \mathbb{R}^{n+1}$. We assume that newborn fish cannot reproduce in their first time step of life and place them in a zeroth class with population $P_0(t)$. After one time step, surviving newborn fish enter class 1 with population $P_1(t)$. Let L_i be the average length of fish in class $i = 0, 1, \ldots, n$, so L_0 is the average length of newborn fish. The time step size is constant and might be determined by the behavior of the species or by the timing of the data collection. We assume that in each time step a surviving fish either stays in its length class or grows into the next length class but cannot skip beyond the next length class, so $L_{i+1} - L_i$ is the maximum a fish in class $i = 1, 2, \ldots n - 1$ can

grow in one time step.

We now set up our model. Let s_i be the survival rate of fish in class $i = 0, 1, \ldots n$ each year and f_i the fecundity of fish in class $i = 1, 2, \ldots n$ each year. We assume that the time step in the model is less than or equal to the time needed to reach maturation size. In [1] the following assumptions are made:

A: The survival rates satisfy $0 < s_i \le 1$ and the fecundities $f_i \ge 0$ and not all $f_i = 0$. The sequences $(f_j)_{j=0}^n$ and $(s_j)_{j=0}^n$ are nondecreasing.

The condition on $(f_j)_{j=0}^n$ is satisfied for most species since the fecundity of larger fish is greater because larger fish can hold more eggs. The assumption that the survivals $(s_j)_{j=0}^n$ are nondecreasing, is not as plausible, since angling can decrease the population of large fish more than the population of small fish. One of the purposes of the current paper is to study what happens when the survival condition in Assumption A is removed. Therefore, in this paper we will work with the modified assumptions:

A': The survival rates satisfy $0 < s_i \le 1$ and the fecundities $f_i \ge 0$ and not all $f_i = 0$. The sequence $(f_j)_{j=0}^n$ is nondecreasing.

Let p_t be the probability at time step t that a fish grows into the next length class in one time step. We use a model of the form

$$\vec{P}(t+1) = A_{p_t}\vec{P}(t)$$

where

$$A_{p_t} = \begin{bmatrix} 0 & f_1 & f_2 & f_3 & \cdots & f_{n-2} & f_{n-1} & f_n \\ s_0 & s_1(1-p_t) & 0 & 0 & \cdots & 0 & 0 & 0 \\ 0 & s_1p_t & s_2(1-p_t) & 0 & \cdots & 0 & 0 & 0 \\ 0 & 0 & s_2p_t & s_3(1-p_t) & \cdots & 0 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \vdots & \vdots \\ 0 & 0 & 0 & 0 & \cdots & s_{n-2}p_t & s_{n-1}(1-p_t) & 0 \\ 0 & 0 & 0 & 0 & \cdots & 0 & s_{n-1}p_t & s_n \end{bmatrix}.$$

$$(1)$$

Thus the population at time t+1 is a A_{p_t} times the population at time t. This equation is solvable once an initial population $\vec{P}(0)$ is specified. A matrix of this type is known as a "population projection matrix". We now give an interpretation of this matrix. When the top row is multiplied by the population vector, we get

$$\sum_{j=1}^{n} f_j P_j(t),$$

which can be interpreted as the number of newborns generated by the current population. Hence the 0th entry of $A_{p_t}\vec{P}(t)$ is the number of newborns generated from the current population. The jth entry of $A_{p_t}\vec{P}(t)$ for $j=2,\ldots n$ is

$$s_{i-1}p_tP_{i-1} + s_i(1-p_t)P_i$$
.

Since p_t is the probability that a fish will move to the next length class, this can be interpreted as the number of fish that survived and got bigger from the (j-1)th length class, plus the number of fish that survived and didn't get bigger from the jth length class. When j=1, the jth entry of $A_{p_t}\vec{P}(t)$ is just how many survive from the 0th class.

As in [1, 2, 6], we assume that p_t is a strictly decreasing function of the population biomass. That is, the more biomass there is in the population, the less likely a fish is to move to the next length class. Furthermore, we expect the probability to be near zero when the biomass is very large, and we expect the probability to be 1 when the biomass is zero. Let B(t) denote the population biomass at time step t. We need to identify a function g so that $p_t = g(B(t))$.

In [1, 2, 6] that function is

$$g(y) = \frac{1}{1 + b_{growth} \ y}$$

where b_{growth} is a scaling parameter. Note that that g is strictly decreasing on $[0,\infty)$ (since more biomass means lower probability), g(0)=1 (since no biomass means probability 1) and $\lim_{y\to\infty}g(y)=0$ (since large biomass means small probability).

Following [1, 2, 6], we assume that the mass of a fish of length L_i is $W_i = \alpha L_i^3$ where α is the mass-length coefficient. The population biomass at time step t is then approximated by

$$B(t) = \sum_{i=0}^{n} W_i P_i(t) \tag{2}$$

We study the nonlinear dynamical system

$$\vec{P}(t+1) = A_{p_t}\vec{P}(t), \ p_t = g(B(t)), \ \vec{P}(0) = \vec{P}_0.$$
 (3)

This system gives the population at time step t+1 as the matrix multiplication of matrix A_{p_t} with the population $\vec{P}(t)$, where the probability of growth p_t is updated at each time step.

3 Mathematical Conjectures

In this section we give some conjectures about the solutions to (3).

Definition 3.1 A vector is positive if all of its entries are positive. A vector is nonnegative if all its entries are nonnegative. A nonnegative vector is nonzero if not all its entries are zero.

In the study of dynamical systems, the *equilibria* play a central role. An equilibrium is a vector that is unchanged by the system from a time step to the next.

Definition 3.2 1. We say that a vector $\vec{P}^* = [P_1^*, P_2^*, \dots P_n]^T$ is an equilibrium for (3) if

$$\vec{P}^* = A_{p^*} \vec{P}^*, \ p^* = g(B^*), B^* = \sum_{i=0}^n W_i P_i^*$$

2. We say that an equilibrium \vec{P}^* for (3) is globally attracting if for every nonnegative nonzero \vec{P}_0 ,

 $\lim_{t \to \infty} \vec{P}_{p_t} = \vec{P}^*.$

The goal of this paper is to use simulations to make the following conjecture plausible.

Conjecture 3.1 1. If $\rho(A_1) < 1$, then the zero population $\vec{0}$ is globally attracting.

- 2. If $\rho(A_0) > 1$, then $\lim_{t \to \infty} ||\vec{P}(t)|| = \infty$ for all nonzero nonnegative initial states $\vec{P}(0)$.
- 3. If $\rho(A_0) < 1 < \rho(A_1)$, then the system has a unique nonzero positive equilibrium \vec{P}^* which is globally attracting.

Before discussing what these conjectures mean, we will discuss the role of the spectral radius in predicting population dynamics. Consider first the simpler linear model

$$\vec{P}(t+1) = A\vec{P}(t). \tag{4}$$

for a constant matrix A. It is well known from linear algebra that the long term behavior of $\vec{P}(t)$ is determined by $\rho(A)$. Roughly speaking, if $\rho(A) < 1$, then all solutions of (4) go to zero; if $\rho(A) > 1$, then all solutions of (4) go to infinity in norm; if $\rho(A) = 1$, all solutions converge, but not uniformly. The moral is that the growth, decay or convergence of the population is determined by $\rho(A)$ in the linear case. In our nonlinear model, the analysis uses $\rho(A_0)$ and $\rho(A_1)$:

For Case 1 of Conjecture 3.1: A_1 describes what happens when the probability of a fish moving to the next length class in one time step is always 1 - this is in some sense the "best case scenario" for the fish. If $\rho(A_1) < 1$, that means that in the best case scenario the population goes to zero, so it is plausible that the solution of (3) goes to zero as well.

For Case 2 of Conjecture 3.1: A_0 describes what happens when the probability of a fish moving to the next length class in one time step is always 0 - this is in some sense the "worst case scenario" for the fish. If $\rho(A_0) > 1$, that means that in the worst case scenario the population goes to infinity, so it is plausible that the solution of (3) goes to infinity as well.

In Case 1 the population is endangered, while Case 2 is unlikely to happen in a physical habitat. If the population is not endangered and the population is known to stay finite, the nonlinear probability of growth kicks in, and Case 3 is likely to occur. The conjectures for Case 1 and Case 2 were proved in [1] in

the case where the survivals are a nondecreasing function of size class. In this paper we numerically verify that this appears to be true when this condition is removed. For Case 3, in [1] the global convergence of the biomass is proved, but not the global attractivity of the equilibrium. In this paper we numerically verify that the global attractivity of the equilibrium is true.

4 Numerical Simulations

We have done extensive simulations which illustrate Conjecture 3.1. We will give examples to illustrate each of the cases in this Conjecture. As a case study we will use the parameters for the white perch population studied in [1, 2]. Please see these papers for a discussion of the species. We will use the simulation parameters L_i , f_i , and α given in Section 4 of [1] (with n=8, so there are nine stages in the population vector). We will be varying the survival probabilities s_j to get the three cases in Conjecture 3.1. Since Cases 1 and 2 of Conjecture 3.1 have been established in [1] when (s_j) is nondecreasing, our examples will use survivals that are not nondecreasing.

1. For case 1, we'll use survival rates

In this case $\rho(A_1) = 0.777272 < 1$, so this satisfies the hypotheses of case 1. The population dynamics for this model (with an arbitrarily chosen initial population) are given in Figure 1. The top graph represents the first size class, and the bottom graph represents the last size class.

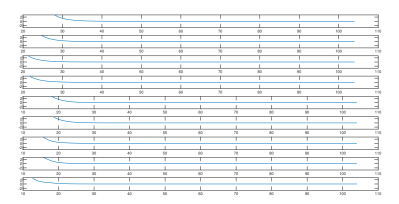


Figure 1: Example for case 1 of Conjecture 1, $\rho(A_1) < 1$; the population appears to converge to the predicted zero population.

For us to say that the population converges to zero, the population for each size class is compared after each time step with the previous time step and the absolute value of the difference is be calculated. If the difference was less than 2^{-1021} (which is the smallest float in Matlab) we conclude convergence. We repeated this simulation for many randomly chosen initial populations, and for every initial condition the population appeared to converge to zero.

2. For case 2, we use the following survival rates:

$$[.99, 1, .97, .96, .95, .94, .93, .92, .91]$$

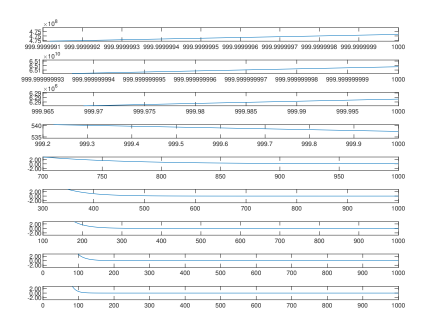


Figure 2: Example for case 2, $\lim_{t\to\infty} \|\vec{P}(t)\| = \infty$ and p_t approaches 0.

In this case we see that $\rho(A_0)=1.007207>1$. We check convergence to infinity by setting a large upper bound, and showing that the total population eventually exceeds that upper bound. We did simulations with 100,000 time steps, and found clear evidence that the population in the first two stages goes to infinity, the population in the third stage goes to 6.2985×10^6 , and the population in the larger stages goes to zero. We only show 1000 time steps in Figure 2, in order to get a clearer picture. As expected, the probability p_t appears to converge to zero, but does not get to zero because the biomass does not get to infinity. Also, we see in this example an extreme case of *stunting*, where the population of the smaller fish dominates as the population gets more and more crowded.

3. For case 3, we use survival rates:

[.7.7.7.7.4.4.4.4.4]

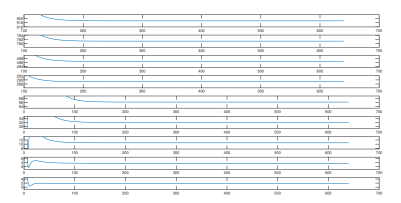


Figure 3: Example for case 3, $\rho(A_0) < 1 < \rho(A_1)$; the population converges to a unique limiting population.

In this case $\rho(A_0) = 0.707258$ and $\rho(A_1) = 1.149390$, so $\rho(A_0) < 1 < \rho(A_1)$, and the hypotheses for case 3 are satisfied. We test convergence to the limiting population in the same way that we tested convergence to zero in case 1, using the tolerance 2^{-1021} . It is easy to check that the limiting population is in fact an equilibrium.

Part of Cases 1 and 3 of Conjecture 3.1 is that the convergence is independent of the initial population. To illustrate this, we give in the Appendix two charts one for Case 1 and one for Case 3 - where the top rows gives 15 initial populations vectors, and the bottom row gives the corresponding limiting population vectors. We see from these that the limiting population is indeed independent of the initial population (up to at least seven significant figures) in these examples. Thus in Case 1 it appears that the zero vector is in fact globally attracting, and in Case 3 it appears that the positive equilbrium is in fact globally attracting. We found the same type of results for other examples.

We have done many more simulations, with many choices of survivals, and many initial conditions. All of them are consistent with Conjecture 3.1. Of course, that is not a proof of Conjecture 3.1, but it does make the Conjecture very plausible.

5 Distribution of biomass in the limiting population

In [1] we studied (in Case 3 and examples where the survivals were nondecreasing) the relationship between the limiting total population biomass and the distribution of biomass in the nine stages. It was found, and proved mathematically, that as the total biomass got larger, the biomass became more concentrated in the lower stages. In this paper we would like to illustrate that we expect this to happen even when the survivals are not nondecreasing. In our case study, we use all of the parameter values from the previous section, except use the survivals

$$(.99 - \alpha, .98 - \alpha, .97 - \alpha, .96 - \alpha, .95 - \alpha, .94 - \alpha, .93 - \alpha, .92 - \alpha, .91 - \alpha)$$

where α is a parameter which varies from 0 to .91. The total limiting population biomass is very large when α is very close to 0, and is very close to 0 when α is close to .91; since we cannot take infinitely many time steps, the total biomass never gets to zero. We show the total biomass on the curve in Figure 4, with value of α on the x-axis and the biomass scale shown on the right y-axis. To illustrate the biomass distribution with a heat map, we take 9 equally-spaced values of α , and above each value of α there are colored boxes which indicate what percentage of the biomass is in each of the nine length stages. The stages are shown on the left y-axis, the the smallest stage at the top and the largest stage at the bottom (note that we label the stages 1 through 9 instead of 0 through 8). The scale for the heat map is on the top of Figure 4.

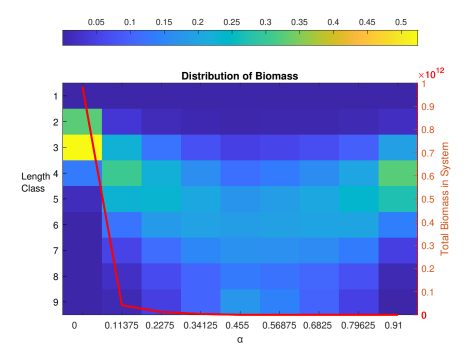


Figure 4: Heat map for biomass distribution as a function of α .

In all cases a small fraction of the biomass is in the newborn fish class. When α is near zero, the total biomass is very high, and is skewed towards the P_1 class. As α increase, we see that the biomass percentage shifts towards the larger classes of fish until $\alpha=.455$. Between $\alpha=.455$ and $\alpha=.56875$, the total biomass in the system falls to near zero, and stays there for larger values of α . Calculating the spectral radius of A_1 and A_0 where $\alpha=.455$, yields $\rho(A_1)=1.000507$ and $\rho(A_0)=0.532369$, which is Case 3. When $\alpha=.568750$, $\rho(A_1)=0.807852$ and $\rho(A_0)=0.418629$, which is Case 1. Thus the system changes from Case 3 to Case 1 for some α between .455 and .56875. Just looking at α from 0 to .455, we're in the most interesting case, and we see that smaller biomass corresponds to a larger percentage of large fish. This is consistent with the results in [1], even though that paper doesn't apply to our survival parameters.

For larger values of α , we're in Case 3, where where the population crashes; however, in practice, we'll never get to zero biomass, so it is still of theoretical interest how the population is distributed, and the heat maps shows a limiting biomass distribution. We find that as α gets closer to 1 (so the survivals get smaller), the population starts moving towards smaller fish, which was not expected.

6 Conclusions

In this paper we study a discrete-time length-structured model for invasive fish which incorporates a nonlinear growth probability. This model was proposed in [1], and studied there with the restrictive condition that the survival parameters be nondecreasing. We numerically studied the case where the survival parameters do not satisfy this restrictive condition. We give Conjecture 3.1 about the long-term behavior of the solutions - this conjecture is stronger than the results proved in [1]. We then do numerical simulations to test this conjecture, and all simulations are consistent with the conjecture. Finally, we study the effect of the total biomass on the length distribution of the limiting population.

These results are significant because population managers care about the long-term behavior of populations. This study makes it plausible that if the populations are bounded (which is expected in real habitats), then the population either goes to zero, or the population approaches some equilibrium which is independent of the initial population. The independence of initial population is important because it implies that the limiting population can be computed even if the initial population is unknown.

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7 Appendix

In this Appendix we illustrate the part of Conjecture 3.1 that says that in Cases 1 and 3 the limiting population is independent of the initial population.

Figure 5 illustrates the convergence to zero in Case 1, and Figure 6 illustrates the convergence to an equilibrium in Case 3. The top rows represent the randomly chosen initial populations vectors, and the bottom rows represent the corresponding limiting population vector.

		Test 1	Test 2	Test 3	Test 4	Test 5	Test 6	Test 7	Test 8	Test 9	Test 10	Test 11	Test 12	Test 13	Test 14	Test 15
		Initial Pop	ulation ve	ctors												
Population	Population Class 0		2353.124	1812.296	4904.518	2883.791	1862.671	3334.658	1944.419	4498.567	614.075	3474.026	93.06387	1878.461	2000.399	2699.525
Population	n Class 1	4613.723	2803.567	3940.567	1433.102	129.2874	2965.923	4668.628	2273.709	2251.968	2036.592	4171.845	3373.882	2732.769	4159.357	476.8635
Population	n Class 2	4001.86	1345.458	3901.479	4004.101	2232.655	4362.763	4054.75	1233.436	1028.362	1376.435	3048.148	2192.544	2809.601	671.6917	732.5743
Population	n Class 3	1429.734	3745.092	3342.561	4480.557	3231.51	4667.508	2422.741	3922.115	4498.255	3583.349	2873.686	2189.101	1979.111	302.3339	3155.706
Population	n Class 4	2718.316	2519.439	667.5193	2987.633	2606.015	3342.321	3783.746	4414.188	3812.928	1416.922	1630.211	585.1841	1990.654	421.2353	4296.602
Population	n Class 5	4923.881	3234.048	107.7794	4420.084	1861.563	1033.882	2085.237	4568.558	4412.432	4480.994	2282.123	4073.408	2576.836	819.4916	4871.108
Population	n Class 6	3578.39	1538.728	2799.204	4718.658	4685.673	3269.253	4858.93	2791.425	1424.751	4132.894	3568.978	1624.277	3287.653	1621.1	2854.192
Population	Population Class 7		693.6232	1504.095	2745.79	4147.664	360.2578	4939.874	2994.341	3366.13	1950.133	4422.025	1231.141	4754.576	1508.634	4984.251
Population	n Class 8	2166.303	2377.865	4697.049	3641.934	4245.427	2033.635	4320.738	744.3836	3321.4	2489.515	3604.278	1713.566	3611.743	58.40496	2767.708
		Limiting Population Vectors														
Population	n Class 0	1.81E-12	1.83E-12	1.71E-12	1.64E-12	2.22E-12	1.67E-12	1.81E-12	1.59E-12	1.58E-12	1.68E-12	1.61E-12	2.08E-12	2.25E-12	2.22E-12	1.67E-12
Population	n Class 1	5.22E-13	5.27E-13	4.92E-13	4.74E-13	6.41E-13	4.82E-13	5.24E-13	4.58E-13	4.56E-13	4.85E-13	4.66E-13	6.01E-13	6.50E-13	6.42E-13	4.83E-13
Population	n Class 2	1.51E-13	1.52E-13	1.42E-13	1.37E-13	1.85E-13	1.39E-13	1.51E-13	1.32E-13	1.32E-13	1.40E-13	1.35E-13	1.74E-13	1.88E-13	1.85E-13	1.39E-13
Population	n Class 3	1.74E-13	1.76E-13	1.64E-13	1.58E-13	2.14E-13	1.61E-13	1.75E-13	1.53E-13	1.52E-13	1.62E-13	1.56E-13	2.01E-13	2.17E-13	2.14E-13	1.61E-13
Population	n Class 4	5.03E-14	5.08E-14	4.74E-14	4.56E-14	6.17E-14	4.64E-14	5.04E-14	4.41E-14	4.39E-14	4.67E-14	4.49E-14	5.79E-14	6.26E-14	6.18E-14	4.65E-14
Population	n Class 5	5.81E-14	5.87E-14	5.48E-14	5.27E-14	7.13E-14	5.36E-14	5.82E-14	5.10E-14	5.08E-14	5.40E-14	5.19E-14	6.69E-14	7.23E-14	7.14E-14	5.37E-14
Population	n Class 6	1.68E-14	1.69E-14	1.58E-14	1.52E-14	2.06E-14	1.55E-14	1.68E-14	1.47E-14	1.47E-14	1.56E-14	1.50E-14	1.93E-14	2.09E-14	2.06E-14	1.55E-14
Population	n Class 7	1.94E-14	1.96E-14	1.83E-14	1.76E-14	2.38E-14	1.79E-14	1.94E-14	1.70E-14	1.69E-14	1.80E-14	1.73E-14	2.23E-14	2.41E-14	2.38E-14	1.79E-14
Population	n Class 8	7.87E-15	7.94E-15	7.42E-15	7.14E-15	9.66E-15	7.26E-15	7.89E-15	6.90E-15	6.87E-15	7.31E-15	7.02E-15	9.06E-15	9.79E-15	9.67E-15	7.28E-15

Figure 5: 15 initial populations and their respective limiting populations of case 1 of conjecture 1, all initial populations lead to limiting populations that are indistinguishable from zero.

		Test 1	Test 2	Test 3	Test 4	Test 5	Test 6	Test 7	Test 8	Test 9	Test 10	Test 11	Test 12	Test 13	Test 14	Test 15
		Initial Pop	oulation ve	ctors												
Population Class 0		353.4217	2353.124	1812.296	4904.518	2883.791	1862.671	3334.658	1944.419	4498.567	614.075	3474.026	93.06387	1878.461	2000.399	2699.525
Population	opulation Class 1		2803.567	3940.567	1433.102	129.2874	2965.923	4668.628	2273.709	2251.968	2036.592	4171.845	3373.882	2732.769	4159.357	476.8635
Population	Class 2	4001.86	1345.458	3901.479	4004.101	2232.655	4362.763	4054.75	1233.436	1028.362	1376.435	3048.148	2192.544	2809.601	671.6917	732.5743
Population	Class 3	1429.734	3745.092	3342.561	4480.557	3231.51	4667.508	2422.741	3922.115	4498.255	3583.349	2873.686	2189.101	1979.111	302.3339	3155.706
Population	Class 4	2718.316	2519.439	667.5193	2987.633	2606.015	3342.321	3783.746	4414.188	3812.928	1416.922	1630.211	585.1841	1990.654	421.2353	4296.602
Population	Class 5	4923.881	3234.048	107.7794	4420.084	1861.563	1033.882	2085.237	4568.558	4412.432	4480.994	2282.123	4073.408	2576.836	819.4916	4871.108
Population	Class 6	3578.39	1538.728	2799.204	4718.658	4685.673	3269.253	4858.93	2791.425	1424.751	4132.894	3568.978	1624.277	3287.653	1621.1	2854.192
Population	Class 7	4194.848	693.6232	1504.095	2745.79	4147.664	360.2578	4939.874	2994.341	3366.13	1950.133	4422.025	1231.141	4754.576	1508.634	4984.251
Population	Class 8	2166.303	2377.865	4697.049	3641.934	4245.427	2033.635	4320.738	744.3836	3321.4	2489.515	3604.278	1713.566	3611.743	58.40496	2767.708
	Limiting Populati		opulation	Vectors												
Population	Class 0	1.81E-12	1.83E-12	1.71E-12	1.64E-12	2.22E-12	1.67E-12	1.81E-12	1.59E-12	1.58E-12	1.68E-12	1.61E-12	2.08E-12	2.25E-12	2.22E-12	1.67E-12
Population	Class 1	5.22E-13	5.27E-13	4.92E-13	4.74E-13	6.41E-13	4.82E-13	5.24E-13	4.58E-13	4.56E-13	4.85E-13	4.66E-13	6.01E-13	6.50E-13	6.42E-13	4.83E-13
Population	Class 2	1.51E-13	1.52E-13	1.42E-13	1.37E-13	1.85E-13	1.39E-13	1.51E-13	1.32E-13	1.32E-13	1.40E-13	1.35E-13	1.74E-13	1.88E-13	1.85E-13	1.39E-13
Population	Class 3	1.74E-13	1.76E-13	1.64E-13	1.58E-13	2.14E-13	1.61E-13	1.75E-13	1.53E-13	1.52E-13	1.62E-13	1.56E-13	2.01E-13	2.17E-13	2.14E-13	1.61E-13
Population	Class 4	5.03E-14	5.08E-14	4.74E-14	4.56E-14	6.17E-14	4.64E-14	5.04E-14	4.41E-14	4.39E-14	4.67E-14	4.49E-14	5.79E-14	6.26E-14	6.18E-14	4.65E-14
Population	Class 5	5.81E-14	5.87E-14	5.48E-14	5.27E-14	7.13E-14	5.36E-14	5.82E-14	5.10E-14	5.08E-14	5.40E-14	5.19E-14	6.69E-14	7.23E-14	7.14E-14	5.37E-14
Population	Class 6	1.68E-14	1.69E-14	1.58E-14	1.52E-14	2.06E-14	1.55E-14	1.68E-14	1.47E-14	1.47E-14	1.56E-14	1.50E-14	1.93E-14	2.09E-14	2.06E-14	1.55E-14
Population	Class 7	1.94E-14	1.96E-14	1.83E-14	1.76E-14	2.38E-14	1.79E-14	1.94E-14	1.70E-14	1.69E-14	1.80E-14	1.73E-14	2.23E-14	2.41E-14	2.38E-14	1.79E-14
Population	Class 8	7.87E-15	7.94E-15	7.42E-15	7.14E-15	9.66E-15	7.26E-15	7.89E-15	6.90E-15	6.87E-15	7.31E-15	7.02E-15	9.06E-15	9.79E-15	9.67E-15	7.28E-15

Figure 6: 15 initial populations and their respective limiting populations of case 1 of conjecture 1, all initial populations lead to limiting populations that are indistinguishable from zero.

1		Test 1	Test 2	Test 3	Test 4	Test 5	Test 6	Test 7	Test 8	Test 9	Test 10	Test 11	Test 12	Test 13	Test 14	Test 15
2			ulation vec		1621.4	rest 5	resco	rest /	resco	resco	rest 10	rest 11	rest 12	rest 15	rest 14	Test 15
3	Population Class 0		2664.128		4616 808	2074 48	2520 201	1186 /18	1077 576	1676 784	4518.603	4522 611	1/12 2708	212 1557	2502 075	866 0421
	Population Class 1					1311.059		2294.244			4454.613					1954.689
	Population Class 2	2542.543				3014.215					1670.815					
	Population Class 3										3493.729					4016.822
	Population Class 4					1108.734					989.0491			4090.743		
	Population Class 5		3112.375								152.7047					
	Population Class 6	3221.591				1483.379					3720.371					
	Population Class 7	1893.047				1593.892					2500.112			749.3272		
	Population Class 8										2399.611			3298.026		3284.299
12	r opulation class o	Limiting Population vectors			20 101000	EZECIOO I	EGDETGES	00001070	2000.000	00701200	20001022	1102100	0.100.100	OLDOIGEO	000,000	OLO IILDI
	Population Class 0				388192.4	388192.4	388192.4	388192.4	388192.4	388192.4	388192.4	388192.4	388192.4	388192.4	388192.4	388192.4
	Population Class 1										11037584					_
	Population Class 2	1187950								1187950						
	Population Class 3	103528				103528										
	Population Class 4	7554,574									7554.574					7554.574
	Population Class 5	472,728														
	Population Class 6	25.82288									25.82288					25.82288
	Population Class 7	1.248516									1.248516					
21	Population Class 8	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708	0.062708

Figure 7: 15 initial populations and their respective limiting populations of case 3 of conjecture 1, all initial conditions lead to the same limiting vector to seven significant figures.