Ecoepidemic predator–prey model with feeding satiation, prey herd behavior and abandoned infected prey

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A B S T R A C T
In this paper we analyse a predator–prey model where the prey population shows group defense and the prey individuals are affected by a transmissible disease. The resulting model is of the Rosenzweig–MacArthur predator–prey type with an SI (susceptible-infected) disease in the prey. Modeling prey group defense leads to a square root dependence in the Holling type II functional for the predator–prey interaction term. The system dynamics is investigated using simulations, classical existence and asymptotic stability analysis and numerical bifurcation analysis. A number of bifurcations, such as transcritical and Hopf bifurcations which occur commonly in predator–prey systems will be found. Because of the square root interaction term there is non-uniqueness of the solution and a singularity where the prey population goes extinct in a finite time. This results in a collapse initiated by extinction of the healthy or susceptible prey and thereafter the other population(s). When also a positive attractor exists this leads to bistability similar to what is found in predator–prey models with a strong Allee effect. For the two-dimensional disease-free (i.e. the purely demographic) system the region in the parameter space where bistability occurs is marked by a global bifurcation. At this bifurcation a heteroclinic connection exists between saddle prey-only equilibrium points where a stable limit cycle together with its basin of attraction, are destruct. In a companion paper (Gimmelli et al., 2015) the same model was formulated and analysed in which the disease was not in the prey but in the predator. There we also observed this phenomenon. Here we extend its analysis using a phase portrait analysis. For the three-dimensional ecoepidemic predator–prey system where the prey is affected by the disease, also tangent bifurcations including a cusp bifurcation and a torus bifurcation of limit cycles occur. This leads to new complex dynamics. Continuation by varying one parameter of the emerging quasi-periodic dynamics from a torus bifurcation can lead to its destruction by a collision with a saddle-cycle. Under other conditions the quasi-periodic dynamics changes gradually in a trajectory that lands on a boundary point where the prey go extinct in finite time after which a total collapse of the three-dimensional system occurs.

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1. Introduction

Recently the role of social behavior in the context of interacting populations has been introduced in predator–prey models. In the classical Rosenzweig–MacArthur model [19,20] both prey and predators have an homogeneous spatial distribution. The prey grows logistically in the absence of the predator and the natural predator mortality rate. The predator–prey interaction is described by a Holling type II functional response (the predation rate per predator which is a monotonic increasing prey-dependent hyperbolic relationship) where a handling time of the prey introduces feeding satiation. In [7] the predators are assumed to have a heterogeneous spatial distribution (for instance when they form a colony or school). Then the functional response depends on both predator and prey densities in a manner that reflects feeding interference between predators. This leads to a ratio-dependent or Beddington–DeAngelis type of functional response (see also [6]). In [10,21] on the other hand, the prey spatial distribution is heterogeneous giving group defense and the Holling type IV or Monod–Haldane functional response is used. This expression is also prey-dependent but the function is now not monotonically increasing. The predation rate per predator decreases for larger prey densities. Bate and Hilker [4] note that Holling type IV functional responses usually result in an upper threshold of prey density, beyond which the predator cannot survive. Further, in recent work

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the predators functional response is derived starting from first principles.

Here we study a different formulation with heterogeneous prey spatial distribution on the ground. The prey gather together in herds where only prey individuals that live close to the herds boundary on the ground are subject to hunting by predators. In [1,5,23,25] this feature has been taken into account in ecoepidemiological systems. These, besides ecological situations dealing with demographically interacting populations, consider also a transmissible disease in the system, see [16,24] for an introductory account.

In a parallel paper [8] an ecoepidemiological model in which the epidemics spread among the predators was proposed. Here we investigate a model where the prey are affected by a disease that propagates by contact. With respect to earlier formulations, these models exhibit the feature of feeding satiation, modeled via a Holling type II response function such as in the Rosenzweig–MacArthur model [20]. However, here the prey-dependent hyperbolic relationship is expressed as a function of the “square root” of the prey size instead of the prey size itself. It differs from the herd behavior model presented in [1], because it takes into account the feeding saturation phenomenon also explored in [8]. In the recent paper [4] a similar problem was studied but the predator-group defending prey functional response was the Holling type IV instead of the “square root” functional response.

The paper is organized as follows. In Section 2 we present the ecoepidemic model and the outline the methodology of the study. The two-dimensional models, the epidemic one, with infected prey population, and the purely demographic, i.e. disease-free, predator–prey model, are analysed respectively in Sections 3 and 4. Here we extend the analysis of [8] by a phase portrait analysis to study the total collapse of the system caused by a heteroclinic connection between the two prey-only saddle equilibria.

In Section 5 we move to the analysis of the full model where the prey is affected by the infectious disease, assuming that diseased individuals are left behind by the herd. We start with a classical existence and stability analysis of all equilibria in Sections 5.1 and 5.2. In Section 5.3, the numerical bifurcation analysis is carried out, completed for the special instance of codimension-two bifurcations. In addition to the bifurcations of the classical predator–prey models, i.e. transcritical, tangent (saddle node) and Hopf bifurcations, here also the torus (Neimark–Sacker) bifurcation occurs. A new phenomenon is represented by the abrupt destruction of the quasi-periodic dynamics on a torus similar to what was found in [3,6].

In Section 6 the results of all particular cases will be compared with the results of the ecoepidemic model with the infected predator population, instead of the prey, analysed in [8] and a final discussion concludes the paper. Assuming that the carrying capacity is sufficiently high to support coexistence of prey and predator, due to the weakening of the prey population by infection, the predator feeding on the prey population can persist for higher predators natural mortality rates.

2. Modeling and analysis approach

2.1. The model

We consider the model presented in [23], which we briefly illustrate again here for the convenience of the reader, to better emphasize the changes in that main model. The basic ecological model is an adapted Rosenzweig–MacArthur model first discussed in [19] where both prey and predators have an homogeneous spatial distribution. Mathematically, the consumption rate of the prey by the predator is expressed via a hyperbolic relationship.

In our case the spatial distribution of the prey population, forming a herd and occupying a certain portion of the ground, is heterogeneous. The prey individuals most subject to hunting are those close to the herd boundaries. The area occupied by the herd is proportional to the prey population and therefore to the size of the herd itself. The prey density on the herd perimeter is therefore proportional to the square root of the size of the herd and thus in the hyperbolic relationship of the standard Holling type II term, the prey size is here replaced by a square root of the prey size. The prey population grows logistically in the absence of the predator. In the absence of the prey, the predators die exponentially fast.

In order to model the spread of the disease, the prey population is divided into two classes consisting of healthy and diseased individuals. The latter are assumed to be too weak to reproduce and to compete for resources. Therefore the basic two-population demographic predator–prey model is extended into a three-dimensional predator–susceptible prey–infected prey model. As in the classical two compartmental SI-model the law of mass action is used to formulate the infection rate of the susceptible by infected prey, assuming possible contacts among all the individuals of the herd. The infected prey are assumed to be too weak both to reproduce and to compete for resources, i.e. they do not appear in the logistic reproduction function for the healthy prey. The infected prey are further assumed to drift away from the herd when become infected; this for instance occurs for elephants. But in the process, they are still able to infect other individuals in the herd. Once alone, they can easily be hunted by the predators. In view of the ease of these captures, we assume that the predators never get tired of hunting sick isolated prey individuals, this implying that in this case the hunting term is bilinear, i.e. a mass action term, as in the classical Lotka–Volterra model. On the other hand, as stated above, we assume that they can become saturated by hunting the healthy prey in the herd, observing that this hunt requires more effort than that on the infected prey. Thus, mathematically, this is better modelled by a Holling type II response function. The predators’ different attitudes in the prey capture therefore determine the different choices for the functional responses among healthy and infected prey.

The model where the state variables and parameters are overlined in order to be able to introduce re-scaled versions later, reads

\[
\frac{dR}{dt} = rR \left( 1 - \frac{R}{K} \right) - \lambda R \bar{I} - \frac{a\sqrt{RF}}{1 + T\bar{a}\sqrt{R}},
\]

\[
\frac{dI}{dt} = \bar{I} \left( \lambda R - b\bar{F} - \mu \right).
\]

\[
\frac{dF}{dt} = \bar{F} \left( -\frac{e\bar{a}\sqrt{R}}{1 + T\bar{a}\sqrt{R}} + \frac{e\bar{b}\bar{I} - \bar{m}}{1 + T\bar{a}\sqrt{R}} \right).
\]

The system consists of the equation for healthy prey, \( \bar{R}(t) \), reproducing logistically and being subject to the negative effects of hunting as well as to the infection process. The infected prey, \( \bar{I}(t) \), do not reproduce so that they are absent in the logistic growth term in the first equation, nor do they contribute to the population pressure on the susceptible prey, because we assume them to be too weak for that. The spread of the infection is modeled via a bilinear term with rate parameter \( \lambda \). The disease is un-dercoverable, i.e. once entered into this class, an infected individual only exits it by dying at rate \( \mu \), incorporating natural plus disease-related effects or possibly by predation modeled with the Holling type I functional response with rate parameter \( b \). Note that here we disregard the possible healthy prey population pressure on the infected prey, i.e. we do not introduce a term of the type \( e\bar{R}\bar{I} \) into the second equation, assuming that the mortality is already represented by the linear term. Note also that the infected prey are assumed to be left behind by the herd, so that they are hunted on a one-to-one basis by the predators. Hence, they are also an “easy
prey, not too difficult to capture because they are weakened by the disease, we assume that predators never get fed up with them and this explains the Holling type I functional response model formulation. Besides the infected prey, the predators gain from hunting healthy prey. This is expressed by a saturating Holling type II model containing the square root term for herd behavior. We thus distinguish hunting rates on healthy prey, indicated by \( \sigma \) and on infected prey, expressed by the parameter \( \overline{b} \). The conversion factor of both healthy and infected prey into new predators is the same \( \overline{e} \).

As in the companion paper [8] we avoid the possible singularity appearing in the Jacobian matrix by letting \( \overline{F} = \overline{\sqrt{\alpha}} \). The non-dimensionalized model is obtained using the following substitutions

\[
P(t) = \alpha \overline{F}(\tau), \quad F(t) = \beta \overline{F}(\tau), \quad I(t) = \gamma \overline{I}(\tau), \quad t = \delta \tau.
\]

The system becomes

\[
\frac{dP}{d\tau} = \frac{1}{2\delta} \left[ \lambda \left( 1 - \frac{\mu^2}{\alpha^2} \right) - \frac{\sigma}{\gamma} PI - \frac{\alpha}{\beta} \frac{\overline{a}}{\overline{F} \overline{a}P} \right],
\]

\[
\frac{dI}{d\tau} = \frac{1}{\delta} \left( \frac{\overline{\sigma}}{\overline{\alpha}} \frac{\overline{\sigma}}{\overline{\alpha}^2} - \frac{\overline{b}}{\overline{F} - \overline{\mu}} \right),
\]

\[
\frac{dF}{d\tau} = \frac{F}{\delta} \left( \frac{\overline{\sigma}}{\overline{\alpha} + \overline{T} \overline{P}} P + \frac{\overline{\sigma}}{\overline{\gamma} I - \overline{m}} \right).
\]

Let us define the following auxiliary parameters

\[
\alpha = T \sigma, \quad \delta = \frac{1}{2} \overline{\tau}, \quad \beta = \frac{1}{\overline{\alpha}}, \quad \gamma = \frac{1}{\overline{\gamma}}.
\]

where we now make the following choices:

\[
\overline{F} = 2r, \quad \overline{\alpha} = \overline{K}, \quad \overline{\lambda} = \lambda, \quad \overline{\sigma} = \sigma, \quad \overline{a} = a, \quad \overline{F} = T, \quad \overline{b} = \overline{b} = b.
\]

The final form of the system equations reads

\[
\frac{dP}{dt} = \left( 1 - \frac{\mu^2}{\alpha^2} \right) - \frac{1}{\lambda} \frac{a^2 T}{1 + P} F - \frac{1}{2r} PI,
\]

\[
\frac{dI}{dt} = 2I \left( \frac{\lambda}{2ar^2 T^2} \mu^2 - bF - \mu \right),
\]

\[
\frac{dF}{dt} = \frac{F}{T} \left( 1 - \frac{e}{T} \left( 1 + P + ebT - m \right) \right).
\]

Our aim is to study the dynamics of the system depending on the prey carrying capacity \( K \) and the predator mortality rate \( m \).

2.2. Methodology

The main mathematical analysis tool used is bifurcation theory whereby the dependency of the long-term dynamics on parameter variations is studied. We do this by calculation of the equilibria, limit cycles, quasi periodic solutions and chaos. To facilitate the numerical bifurcation analysis we have to choose parameter values. Unless stated otherwise, all the default parameter values used in our analysis are given in Table 1. We used the computer program AUTO [9] to perform the numerical bifurcation analysis of the equilibria and limit cycles.

During a process called continuation the dynamics is followed and changes in the long-term dynamics, for instance from stable to unstable equilibrium, are spotted. These points are called bifurcation points. Table 3 gives a list of the bifurcation points. Also the different line types of the bifurcations used in the diagrams are given. Further information about the basics of bifurcation analysis can be found in e.g. Guckenheimer and Holmes [12], Wiggins [26], Kuznetsov [15], and examples of ecological applications of bifurcation analysis in for instance [2] and [14,22] for the discussion of similar bifurcations as we will find here.

### Table 1

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Value</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( P )</td>
<td>Variable</td>
<td>Healthy or susceptible prey</td>
</tr>
<tr>
<td>( I )</td>
<td>Variable</td>
<td>Diseased or infected prey</td>
</tr>
<tr>
<td>( F )</td>
<td>Variable</td>
<td>Predator</td>
</tr>
<tr>
<td>( a )</td>
<td>0.5</td>
<td>Hunting rate of predator on healthy prey</td>
</tr>
<tr>
<td>( e )</td>
<td>0.5</td>
<td>Conversion factor of prey into predators</td>
</tr>
<tr>
<td>( K )</td>
<td>Variable</td>
<td>Carrying capacity</td>
</tr>
<tr>
<td>( m )</td>
<td>Variable</td>
<td>Natural mortality rate of predators</td>
</tr>
<tr>
<td>( r )</td>
<td>0.7</td>
<td>Intrinsic growth rate of healthy prey</td>
</tr>
<tr>
<td>( T )</td>
<td>0.8</td>
<td>Average time to capture a healthy prey</td>
</tr>
<tr>
<td>( \lambda )</td>
<td>0.7</td>
<td>Contact rate or infection rate for the prey disease</td>
</tr>
<tr>
<td>( b )</td>
<td>0.7</td>
<td>Contact rate of predator and diseased prey</td>
</tr>
<tr>
<td>( \mu )</td>
<td>0.65</td>
<td>Natural+disease-related mortality of infected-prey</td>
</tr>
</tbody>
</table>

### Table 2

<table>
<thead>
<tr>
<th>Description</th>
<th>Attractor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zero-solution equilibrium</td>
<td>( E_0 )</td>
</tr>
<tr>
<td>Disease-free prey-only equilibrium</td>
<td>( E_1 )</td>
</tr>
<tr>
<td>Predator-free predator–prey equilibrium</td>
<td>( E_2 )</td>
</tr>
<tr>
<td>Disease-free predator–prey equilibrium</td>
<td>( E_3 )</td>
</tr>
<tr>
<td>Disease-free predator–prey limit cycle</td>
<td>( L_1 )</td>
</tr>
<tr>
<td>Endemic predator–prey equilibrium</td>
<td>( L_2 )</td>
</tr>
<tr>
<td>Endemic predator–prey quasi-periodic torus dynamics</td>
<td>( L_{23} )</td>
</tr>
</tbody>
</table>

### Table 3

<table>
<thead>
<tr>
<th>Bifurcation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( TC_1 )</td>
<td>Transcritical bifurcation (dashed curve)</td>
</tr>
<tr>
<td>( TC_2 )</td>
<td>Transcritical bifurcation (dashed curve)</td>
</tr>
<tr>
<td>( TC_3 )</td>
<td>Transcritical bifurcation (dashed curve)</td>
</tr>
<tr>
<td>( TC_4 )</td>
<td>Transcritical bifurcation (dashed curve)</td>
</tr>
<tr>
<td>( H_2 )</td>
<td>Hopf bifurcation for disease-free predator–prey system (dotted curve)</td>
</tr>
<tr>
<td>( H_3 )</td>
<td>Hopf bifurcation for disease-free predator–prey system (dotted curve)</td>
</tr>
<tr>
<td>( TR )</td>
<td>Torus bifurcation (long-dashed curve)</td>
</tr>
<tr>
<td>( B )</td>
<td>Bautin bifurcation point</td>
</tr>
<tr>
<td>( C^a )</td>
<td>Global bifurcation for disease-free predator–prey system (solid curve)</td>
</tr>
<tr>
<td>( S )</td>
<td>Destruction of quasi-periodic solution on torus</td>
</tr>
</tbody>
</table>

### Suggestion

Collapsing of system and convergence to equilibrium \( \overline{E}_0 \)
The results are presented in bifurcation diagrams where bifurcation points or curves in the parameter space mark qualitative changes in the long-term dynamics (equilibria, limit cycles and quasi-periodic solutions). In bifurcation diagrams where the state variables are plotted as functions of time or of a parameter, solid (dashed) curves denote stable (unstable) equilibrium values. For limit cycle solutions they denote local maximum and minimum values. We used the computer program Maple [17] to do a symbolic analysis of equilibria. The classical ode45 ode-solver of Matlab [18] was used to perform simulations by integration in time for specific parameter values and initial conditions.

Before we analyse the three-population system (2), called the PIF-model, we start with the analysis of the two two-dimensional subsystems: the SI-model for the prey population, called the PI-model where \( F = 0 \) in (2), and then the demographic predator-prey model, called the PIF-model where \( I = 0 \) in (2), with only healthy prey individuals.

3. The epidemic prey population model

Let us begin by studying the non-dimensional prey (healthy and diseased) only system PI where no predator is present. Strictly speaking the conversion from \( R \) to \( P = \sqrt{r} \) is not necessary since the term with the square root in the predation term in (1a) and (2a) is missing. However, to be able to easily interpret the results obtained with those for the full PIF-model we adhere to model (2) with \( F = 0 \).

The dimensionless model where \( P \) represents healthy or susceptible prey and \( I \) is the diseased or infected prey population size reads

\[
\frac{d\tilde{P}}{dt} = P \left( 1 - \frac{\mu}{\sqrt{r} K T^2} \right) - \frac{1}{2T} \frac{\lambda}{\sqrt{r}} \tilde{P} I, \tag{3a} \\
\frac{dI}{dt} = 2I \left( \frac{\lambda}{2\sqrt{r} T^2} \mu - \frac{\sqrt{r}}{2} \right). \tag{3b}
\]

In the next two sections we will study feasibility and stability of the equilibria of this PIF-system (3).

3.1. Equilibria: PI-system

In the PI phase space, the equilibria \( \tilde{E}_k = (\tilde{P}_k, \tilde{I}_k) \) of the system (3) are found as follows. We have the origin \( \tilde{E}_0 = (0, 0) \), the disease-free \( \tilde{E}_1 = (aT\sqrt{r}, 0) \), and possibly the endemic prey population equilibrium \( \tilde{E}_2 = (\tilde{P}_2, \tilde{I}_2) \), with

\[
\tilde{P}_2 = aT \sqrt{2\mu r}, \quad \tilde{I}_2 = \frac{2r(\lambda K - 2\mu r)}{\lambda K}, \tag{4}
\]

with feasibility condition

\[
\lambda \geq \frac{2\mu r}{K}. \tag{5}
\]

Observe that the carrying capacity \( K \) is involved in such a way that when the system is enriched (by increasing its carrying capacity \( K \)) the prey population becomes more vulnerable to the disease.

In Fig. 1 the vector field for \( K = 20 \) in combination with \( m = 0.6 \) is shown. The equilibrium \( \tilde{E}_1 \) is the intersection of the two null-clines, the vertical curve is the \( I \)-null-cline where the time-derivative of \( I \) vanishes, i.e. the curve \( P = aT \sqrt{2\mu r} \).

3.2. Stability: PI-system

The PI system (3) has the following Jacobian matrix

\[
\tilde{J} = \begin{pmatrix}
1 - \frac{3\mu^2}{KT^2 a^2} - \frac{1}{2T} & -\frac{P}{2T} \\
\frac{2\lambda I P}{r a^2 T^2} & \frac{\lambda P^2}{ra^2 T^2} - 2\mu
\end{pmatrix}. \tag{6}
\]

At the origin the eigenvalues of the Jacobian matrix are \( \tilde{\omega}_{11} = 1 \), \( \tilde{\omega}_{12} = -2\mu \), showing that \( \tilde{E}_0 \) is always an unstable equilibrium. Their respective eigenvectors are \((1, 0)^T\) (unstable manifold) and \((0, 1)^T\) (stable manifold). This will be an important fact when we study the full three-dimensional model below.

At equilibrium \( \tilde{E}_1 \) where \( P = aT\sqrt{r} > 0, I = 0 \) we find the following Jacobian matrix \( \tilde{J}_1 = J(\tilde{E}_1) \)

\[
\tilde{J}_1 = \begin{pmatrix}
-2 - \frac{aT\sqrt{K}}{2T} \\
0 & \lambda K - 2\mu
\end{pmatrix}. \tag{7}
\]

The eigenvalues of the Jacobian matrix at \( \tilde{E}_1 = (aT\sqrt{r} \cdot 0) \) read \( \tilde{\omega}_{11} = -2 \) and \( \tilde{\omega}_{12} = (\lambda K - 2\mu r)/r \). Thus equilibrium \( \tilde{E}_1 \) is stable for

\[
K < \frac{2\mu r}{\lambda} \frac{aT \sqrt{r}}{K}. \tag{8}
\]

and unstable conversely. The parameter value \( K^* = 1.3 \) for the default parameter values, marks a so called transcritical bifurcation parameter \( T C_{12} \) in general identified by the criterion \( \det \tilde{J}_1 = J(\tilde{E}_1) = 0 \). Invasion of the disease is possible when the boundary, disease-free equilibrium \( \tilde{E}_1 \) becomes unstable. To which kind of interior long-term dynamics (whether be it an equilibrium, a limit cycle or chaotic attractor) this invasion leads to is, however, not predicted. On the other hand we know that there is an interior equilibrium given in (4) when the feasibility condition (5) is fulfilled.

The eigenvalues of the Jacobian matrix evaluated at the endemic equilibrium \( \tilde{E}_{12} \) are explicitly evaluated as

\[
\tilde{\omega}_{21} = 2 - r r + \sqrt{r^2 \mu^2 - (\lambda K - 2r\mu)\mu K}, \quad \lambda K \]

\[
\tilde{\omega}_{22} = 2 - r r + \sqrt{r^2 \mu^2 - (\lambda K - 2r\mu)\mu K}. \tag{8}
\]

The expression in the square root term is always smaller than \( r^2 \mu^2 \), in view of the feasibility condition (5). Hence, when the discriminant is positive both eigenvalues are real and negative and the equilibrium is a stable node. On the other hand when the discriminant is negative the real part of the conjugate eigenvalues is negative and the equilibrium is a stable focus. Hence, stability of the equilibrium \( \tilde{E}_{12} \) given by (4) of the endemic prey system is ensured when condition (5) is satisfied.
There is no Hopf bifurcation since the real part of the conjugate eigenvalues differs from zero. In general the Hopf bifurcation of a two-dimensional system occurs when $\text{tr}(J)_{12} = 0$, that is the trace of the Jacobian matrix is zero. Here we have $J_{12} = \hat{\omega}_{21} + \hat{\omega}_{22} = -4\mu/(\lambda K) \neq 0$.

4. The demographic predator–prey model

We begin by recalling and extending some interesting results already obtained in [5,8] for the disease-free or purely demographic model PF. Here we give the extended model formulation, assess its the equilibria and their stability. But furthermore and most importantly for the following analysis of this paper, we perform the full bifurcation analysis and state its results.

In non-dimensionalized form, the model derived in [8], reads as follows, where $P$ denotes prey population size and $F$ again denotes the predator size:

\[
\begin{align*}
\frac{dP}{dt} &= P \left( 1 - \frac{P^2}{a^2 + FK} - \frac{1}{\lambda} \frac{a^2 T}{1 + F^{r}} \right), \\
\frac{dF}{dt} &= F \left( \frac{1}{T} \frac{e}{1 + P} - m \right).
\end{align*}
\]

(9a)

(9b)

This model is obtained by taking $I = 0$ in (2).

In the next sections we perform an existence and stability analysis of the equilibria of this PF-system (9) completed by a bifurcation analysis where we also study the existence and stability analysis of limit cycles.

4.1. Equilibria: PF-system

In the PF phase space the equilibria $E_{k} = (\hat{P}_{k}, \hat{F}_{k})$ of the system (9) are: the origin $E_{0} = (0, 0)$, the predator-free point $E_{1} = (aT\sqrt{K}, 0)$, and possibly coexistence $E_{13} = (\hat{P}_{1}, \hat{F}_{1})$, with

\[
\hat{P}_{1} = \frac{mT}{e - mT}, \quad \hat{F}_{1} = \frac{m}{aT}\sqrt{K}(e - mT)^{2} - m^{2}
\]

(10)

It is feasible for $0 \leq \hat{P}_{1} \leq aT\sqrt{K}$, i.e. explicitly for

\[
eq \frac{ae\sqrt{K}}{1 + aT\sqrt{K}} m \leq \frac{a\sqrt{K}}{1 + a\sqrt{K}} m.
\]

(11)

4.2. Stability: PF-system

The PF system (9) has the following Jacobian matrix

\[
\hat{J} = \begin{pmatrix}
\frac{1}{T} \left( 1 - \frac{3P^2}{KTT^2a^2} + \frac{a^2 T}{\lambda(1 + P^2)} - \frac{a^2 T}{\lambda(1 + P)} \right) \\
\frac{rT}{rT(1 + P)} - \frac{eF}{eFP} - \frac{m}{r} \frac{m}{rT(1 + P)}
\end{pmatrix}
\]

(12)

The eigenvalues of the Jacobian matrix of (9) evaluated at the origin are $\hat{\omega}_{01} = 1$, $\hat{\omega}_{02} = -m/r$; their respective eigenvectors are $(1, 0)^{T}$ and $(1, \lambda(r + m)/(a^2T))^{-1}T$. Thus $E_{0}$ is an unstable saddle.

At equilibrium $E_{1}$ the eigenvectors of $J$ read

\[
\hat{\omega}_{11} = \hat{\omega}_{12} = \frac{1}{r} \left( \frac{e\sqrt{K}}{1 + a\sqrt{K}} - m \right).
\]

Thus equilibrium $E_{1}$ is stable for $m \neq m$ and unstable conversely, (see (11)). The parameter value $m > m$ marks a so called transcritical bifurcation parameter $TC_{1}$. Invasion of the prey is possible when the boundary equilibrium $E_{1}$ becomes unstable. To which kind of interior long-term dynamics (equilibrium, limit cycle or chaotic attractor) this invasion leads to is not predictable from this information.

There is an interior equilibrium $E_{13}$, see (10), when the feasibility condition (11) is fulfilled.

The matrix $\hat{J}$ of the PF-system given in (12) is now evaluated for $P = \hat{P}_{1}$ and $F = \hat{F}_{1}$ given in (10). The eigenvalues are

\[
\omega_{21,22} = \frac{1}{2} \left( \text{tr}(\hat{J}) \pm \sqrt{\text{tr}(\hat{J})^2 - 4\text{det}(\hat{J})} \right).
\]

with

\[
\text{tr}(\hat{J}) = 2 \left( 1 - \frac{3\hat{P}_{1}^2}{\lambda^{2}K} \right) + \frac{a^2 K}{T} \frac{1}{1 + \hat{P}_{1}^2} \hat{F}_{1}.
\]

(13)

\[
\text{det}(\hat{J}) = \frac{2e}{r^2} \frac{a^2 K}{T^2(1 + \hat{P}_{1})^2} \hat{F}_{1}.
\]

(14)

The interior equilibrium $E_{13}$ is stable when $\text{tr}(\hat{J}) < 0$.

The region in the parameter space $(K, m)$ where the interior equilibrium $E_{13}$ is stable is bounded by the codimension-one transcritical $TC_{1}$ and Hopf $H_{2}$ bifurcation curves. The codimension-one transcritical bifurcation curve $TC_{1}$ specified by $\text{det}(J) = 0$ is described by

\[
K_{TC_{1}} = \left( \frac{m}{a(e - mT)} \right)^{2}.
\]

(15)

It has a horizontal asymptote for $m = e/T = 0.625$ for the default parameter values given in Table 1. The Hopf bifurcation curve for this two-dimensional system is given by the trace $\text{tr}(J)_{13} = 0$

\[
K_{H_{1}} = \frac{m^2(e^2T + 3e)}{a^2(m^2T^2 - em^2T + e^3)}.
\]

(16)

In the next subsection these bifurcation curves are calculated using the program auto [9] for the specific parameter set given in Table 1.

4.3. Bifurcation analysis: PF-system

In order to study equilibrium $E_{13}$ and the limit cycle $L_{13}$ originating at the Hopf bifurcation, we use a numerical bifurcation analysis where $K$ and $m$ are taken as the variable parameters while all other ones are fixed at the default values given in Table 1.

![Fig. 2. One-parameter bifurcation diagram for K = 20 and free parameter m, for the demographic PF system (9) for the healthy prey population, P and the predator population, F. The solid curve between the transcritical bifurcation point TC1 and the Hopf bifurcation point H1 denotes stable equilibrium E13 values. Below H2 the equilibrium E13 is unstable and shown as a dashed curve. Between H2 and the global bifurcation point C the maximum and minimum peak values of the stable limit cycle L are shown as solid curves. Table 2 gives a list of the asymptotic dynamics and Table 3 gives a list of the bifurcations.](image)
The one-parameter bifurcation diagram is shown in Fig. 2 where \( m \) is the bifurcation parameter and \( K = 20 \). With large mortality rates (say \( m = 0.7 \)) only the prey population persists. Decreasing the parameter \( m \) the predator population invades at the transcritical bifurcation \( T_{C1} \) leading to the existence of the interior solution of the predator–prey PF system (9). Decreasing the mortality \( m \) further the predator population size increases and at the Hopf bifurcation, \( H_2 \), a limit cycle \( L_{13} \) occurs. Lowering \( m \) the amplitude of these limit cycles grows fast. It is broken by a heteroclinic connection between two saddle equilibrium points where \( F = 0 \) at the global bifurcation point \( G^o \) at \( m = 0.335475 \). (see also [8,22]). Related to this phenomenon is the fact that equilibrium \( E_0 \) is reached in a finite time as it will be shown below.

The pictures in Fig. 3 show the vector field for \( K = 10 \) in combination with \( m = 0.3 \) and 0.278745 respectively. The trajectories shown terminate asymptotically in the stable equilibrium \( E_{13} \) and stable limit cycle \( L_{13} \), respectively. From the origin three curves originate. The dotted line is a numerical approximation of the stable manifold \( W^s_0 \) passing through \( E_0 \) calculated by time backwards simulations. The long-dashed curve is the linear tangent manifold \( T^s_0 \) passing through \( E_0 \) which is a local approximation of the tangent to the stable manifold \( W^s_0 \) corresponding to the negative eigenvalue \(-m/r\) with eigenvector \((1, \lambda \sqrt{r \sigma T} - 1)^T\).

The vector fields in Fig. 3 reveal that for points starting above the curve \( W^s_0 \) leads to crossing the vertical axis where \( P = 0 \). In order to explain this fact, we revisit the dynamics analysis. The Jacobian matrix evaluated at the origin \( E_0 \) yields the eigenvalues 1 and \(-m/r\) and the eigenvectors \((1, 0)^T\) and \((1, \lambda \sqrt{r \sigma T} - 1)^T\). This means that for the dynamics restricted to the line \( P = 0 \) the single eigenvalue is negative \(-m/r\) and that equilibrium \( E_0 \) is stable once \( P(t) \) vanishes in finite time. There can be bistability when there is an interior limit set, for instance the stable equilibrium \( E_{13} \) or the stable limit cycle \( L_{13} \). In order to substantiate this statement we simulated backward in time the system starting from points where \( P(0) = 0 \) and \( F(0) > 0 \) but close to zero. These calculated trajectories are the dotted lines in Fig. 3. They form approximately the stable manifolds \( W^s_0 \) passing through \( E_0 \) which act as separatrices between \( E_0 \) and \( E_{13} \) or \( L_{13} \). In each figure the linear tangent manifold \( T^s_0 \) is the straight long-dashed curve passing through \( E_0 \) which is tangent to and is a local approximation of stable manifold \( W^s_0 \) corresponding to the negative eigenvalue \(-m/r\) with eigenvector \((1, \lambda \sqrt{r \sigma T} - 1)^T\).

In Fig. 4 the population solutions for the parameter combinations \( K = 1 \) and \( m = 0.05 \) are shown where the zero equilibrium is globally attracting, that is there is no stable equilibrium \( E_{13} \) or stable limit cycle \( L_{13} \). These results show that the prey population \( P \) goes extinct in finite time. The time of extinction depends on the initial conditions. Thereafter the predator population \( P \) goes extinct asymptotically, despite the fact that the origin equilibrium \( E_0 \) was unstable. This is a result of the non-uniqueness of the solution due to the square root singularity of the \( \sqrt{ } \) that describes the dynamics of the prey population \( P \).

The vector field close to the origin is shown in Fig. 5. The dashed curves are the null-clines, the vertical curve is the \( E \)-null-cline where the time-derivative of \( P \) vanishes, i.e. the curve \( P = mT(e^{-{mT} - 1} \). This curve goes through the unstable internal equilibrium \( E_{13} \) where it intersects the \( P \)-null-cline where the time-derivative of \( P \) vanishes. The \( P \)-null-cline intersects the horizontal curve at the origin \( E_0 \) and at \( E_1 \) where \( P = aT \sqrt{ } R = 0.4 \) and \( F = 0 \). Note that the arrows cross the vertical axis, the line \( P = 0, \) with rate \( dP/ dt = -a(\sqrt{ } \lambda) F \) and it is negative for \( F > 0 \). The solid line...
is the trajectory for the same initial values, labeled by a ‘O’, as in Fig. 4. These plots show that for all initial values the solution converges to the origin, whereby first the prey goes extinct in finite time (terminating on the vertical axis) and then the predator disappears asymptotically.

In order to substantiate this statement we simulated the system backward in time starting from points where \(P(0) = 0\) and \(F(0) > 0\). Two of these trajectories are shown in Fig. 5 starting from points labeled by the ‘□’ symbol. Both terminate at the interior equilibrium point \(E_{13}\). The curve with \(F(0) = 0.2925\) is special in the sense that starting below this point where \(P(0) = 0\) again, the trajectory goes directly toward the equilibrium without intersecting the \(P\)-nullcline above the equilibrium. On the other hand starting above this point the intersection occurs, as for the upper curve with \(F(0) = 0.9358\) where the time-backward curve intersects the ‘O’ point before it lands on the interior equilibrium point \(E_{13}\). There is also an initial condition where the trajectory lands on equilibrium \(E_1\) (not shown). Starting above this point leads to unbounded solutions for \(t = -\infty\).

For \(m = 0.278745\) the stable manifold \(W^0_0\) is the separatrix between two attractors namely the interior limit cycle \(L_{13}\) and the equilibrium \(E_0\). The separatrix is the common boundary of the basins of attraction of the two attractors \(E_0\) and \(L_{13}\). The stable manifold is also invariant and therefore no trajectory can cross this manifold. For \(m = 0.05\), however, equilibrium \(E_0\) is globally attracting. Therefore, there is a switching point when \(m\) is continued from \(m = 0.278745\) to 0.05 where the separatrix disappears. This happens at the global bifurcation where the limit cycle is broken into two non-smooth connected parts: from \(E_0\) to \(E_1\) and from \(E_1\) to \(E_0\). The part on the boundary is the straight line between \(E_0\) and \(E_1\) where \(F = 0\) and the interior part where \(F > 0\) is the heteroclinic connection where the trajectory starting from \(E_1\) lands exactly in the origin \(E_0\). In that situation the stable manifold \(W^0_1\) is precisely also the nullcline connecting \(E_1\) with \(E_0\) where the stable manifold terminates. At that critical parameter value the basin of attraction of the limit cycle finishes and the separatrix between the attractor \(E_0\) and the interior limit cycle \(L_{13}\) disappears.

In Fig. 6 the two-parameter diagram is shown for the predator–prey system \(PF\), where both \(m\) and \(K\) are varied simultaneously. The transcritical bifurcation \(TC_1\) separates regions where we have \(E_1 \Rightarrow E_{13}\), the Hopf bifurcation \(H_2\) where \(E_{13} \Rightarrow L_{13}\), and the global bifurcation \(G^c\) where \(L_{13} \Rightarrow E_0\). In Figs. 3 and 5 the phase plane plot are given for three points in this two-parameter diagram. In Fig. 3a the parameter combination \(K = 10, m = 0.3\) is a point in Fig. 6 between the curves \(TC_1\) and \(H_2\) where equilibrium \(E_{13}\) is stable. In Fig. 3b the parameter combination \(K = 10, m = 0.278745\) is a point between the curves \(H_2\) and \(G^c\) where limit cycle \(L_{13}\) is stable. In Fig. 4 where \(K = 1, m = 0.05\) is a point in Fig. 6 that lies below the curve \(G^c\) where \(E_0\) is globally stable.

In summary: for low mortality rates the system collapses always completely. For intermediate mortality rates there is bistability between the equilibrium \(E_0\) and equilibrium \(E_{13}\), limit cycle \(L_{13}\), and for higher mortality rates equilibrium \(E_1\). This is indicated in the phase plane plots of Figs. 3 and 5. The boundaries of the basins of attractions (separatrix points) form the switching points in the phase space \((P, F > 0)\) to which attractor the system will converge: interior attractor or a total collapse. Note that for \(F = 0\) there is no positive \(P\) where such a switch occurs like in models with an Allee effect.

In the next section we use these results for comparison with the results of the model where the prey is also suffering from the disease.

5. The predator–prey model with abandoned diseased prey

We study the main three-dimensional system formulated in (2) which is recalled here for the convenience of the reader.

\[
\begin{align*}
\frac{dP}{dt} &= P \left(1 - \frac{P^2}{a^2KT^2}\right) - \frac{1}{\lambda} \frac{a^2T}{T + P} F - \frac{1}{2\lambda} Pl, \\
\frac{dI}{dt} &= 2I \left(\frac{\lambda}{2a^2T^2} F^2 - bF - \mu\right), \\
\frac{dF}{dt} &= \frac{F}{I} \left(1 - \frac{e}{\mu(T + P)} P + ebl - m\right).
\end{align*}
\]

We recall that when \(F = 0\) we have system \(PL\) (3) and when \(I = 0\) system \(PF\) (9).

In the next subsections we discuss the equilibria and their stability. Table 4 summarizes the equilibria and Table 5 their stability for the subsystems and the full system.
Table 4

<table>
<thead>
<tr>
<th>Cases</th>
<th>Equilibria</th>
<th>System</th>
</tr>
</thead>
<tbody>
<tr>
<td>(0, 0, 0)</td>
<td>$E_0 = (0, 0, 0)$</td>
<td>( P )</td>
</tr>
<tr>
<td>(0, 0, 0)</td>
<td>$E_1 = (aT\sqrt{K}, 0, 0)$</td>
<td>( P )</td>
</tr>
<tr>
<td>(0, 0, 0)</td>
<td>$E_{12} = \left( \tilde{P}_1, aT\sqrt{2\mu r}, \tilde{I}_1 = \frac{2r(\lambda K - 2\mu r)}{\lambda K} \right)$</td>
<td>( P )</td>
</tr>
<tr>
<td>(+, +, 0)</td>
<td>$E_{13} = \left( \tilde{P}_3, \tilde{P}_1, \tilde{P}_2, m\lambda e^{\frac{mT}{1+a^nT}} \right)$</td>
<td>( P )</td>
</tr>
<tr>
<td>(+, +, 0)</td>
<td>$E_{123} = (P_0, I_0, 0)$</td>
<td>( P )</td>
</tr>
</tbody>
</table>

5.1. Equilibria: PIF-system

The equilibria $E_3 = (P_0, I_0, F_0)$ are found as follows. We have: the origin $E_0 = (0, 0, 0)$, $E_1 = (aT\sqrt{K}, 0, 0)$ the healthy-prey-only equilibrium, the disease-free predator–prey case equilibrium $E_{12} = (P_0, 0, F_0)$, the endemic predator-free equilibrium $E_{123} = (P_0, I_0, F_0)$ and possibly the endemic predator–prey case $E_{123} = (P_0, I_0, F_0)$. The endemic predator-free prey and the disease-free equilibrium of the two-dimensional subsystems were already discussed in the previous section. The introduction of the extra state variable to obtain the three-dimensional system does not change the previous equilibrium results; only, the extra state variable is zero.

For the predators–free endemic prey equilibrium $E_{12}$ we have, see (4)

$$P_2 = \frac{aT\sqrt{2\mu r}}{e^{-mT}}, \quad I_2 = \frac{2r(\lambda K - 2\mu r)}{\lambda K}, \quad F_2 = 0,$$

with feasibility condition (see (5))

$$\lambda \geq \frac{2\mu r}{K}.$$

For the disease-free predator–prey system we have equilibrium $E_{13}$, see (10)

$$P_3 = \frac{mT}{e^{-mT}}, \quad I_3 = 0, \quad F_3 = \frac{m\lambda e^{-\frac{mT}{1+a^nT}}}{a^4 K(e-mT)^4}.$$

with feasibility condition (see (11))

$$e^{-mT}, \quad m \leq \frac{ae\sqrt{R}}{1 + a^nT} = m^4.$$

For coexistence, solving the second and third equilibrium equation of the system (17), we have

$$F_4 = \frac{\lambda P_2^2 - 2r\mu a^2 T^2}{2ba^2 T^2}, \quad I_4 = \frac{mT(1 + P_2) - eP_2}{ebT(1 + P_2)},$$

and for determining $P_4$ we have the fourth degree polynomial equation

$$\frac{1}{a^2 KT^2}(P_4^3 + P_4^2) + \left( \frac{m}{2reb} - 1 \right)(P_4^2 + P_4) - \frac{a^2 T^2}{\lambda b} = 0.$$

Substitution of this expression into (22) gives the expression for $I_4$ and $F_4$ just in terms of the parameters.

Descartes’ rule of signs tells us the number of roots with positive and negative real parts. For $m > 2reb$, $m = 2reb$ and $m < 2reb$

Table 5

<table>
<thead>
<tr>
<th>Equilibria</th>
<th>Eigenvalues</th>
<th>Stability conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\tilde{E}_0$</td>
<td>$\lambda_{E_0} = 1$</td>
<td>Always unstable</td>
</tr>
<tr>
<td>$\tilde{E}_1$</td>
<td>$\lambda_{E_1} = 2\mu$</td>
<td>Stable $K &lt; \frac{2\mu}{\lambda}$</td>
</tr>
<tr>
<td>$\tilde{E}_{12}$</td>
<td>$\lambda_{E_{12}} = \left( \lambda K - 2\mu r \right)$</td>
<td>( \lambda K &gt; 2\mu r )</td>
</tr>
<tr>
<td>$\tilde{E}_{13}$</td>
<td>$\lambda_{E_{13}} = \frac{mT}{1+a^nT}$</td>
<td>Unstable saddle</td>
</tr>
<tr>
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</tr>
<tr>
<td>$\tilde{E}_1$</td>
<td>$\lambda_{E_1} = 2\mu$</td>
<td>Unstable saddle</td>
</tr>
<tr>
<td>$\tilde{E}_{12}$</td>
<td>$\lambda_{E_{12}} = \left( \lambda K - 2\mu r \right)$</td>
<td>( \lambda K &gt; 2\mu r )</td>
</tr>
<tr>
<td>$\tilde{E}_{13}$</td>
<td>$\lambda_{E_{13}} = \frac{mT}{1+a^nT}$</td>
<td>Unstable saddle</td>
</tr>
<tr>
<td>$\tilde{E}_{123}$</td>
<td>$\lambda_{E_{123}} = \frac{mT}{1+a^nT}$</td>
<td>Unstable saddle</td>
</tr>
</tbody>
</table>

Section 4.3

Section 5.3
there is only one sign change + + + − , + + + and + + + − respectively; hence, there is just one positive solution and therefore it must be real and it gives the value of $P_4$.

With Maple [17] (or other computer programs for symbolic computations) it is possible to derive symbolic expressions for the solutions of the equilibria. However, these expressions are very long and therefore are omitted here.

5.2. Stability: PIF-system

In contrast to the equilibria values of $E_0$, $E_{12}$ and $E_{13}$ their stability for the three-dimensional full PIF-system has to be determined now anew since the results can differ from those of the two-dimensional $P$ and PF-systems derived above.

System PIF [17] has the following Jacobian matrix

$$
J = \begin{pmatrix}
\frac{1}{T} & \frac{3P^2}{aT^2K} + \frac{aT^2F}{K(1 + P)^2} - \frac{P}{2T} & -\frac{aT^2}{K(1 + P)^2} \\
\frac{eF}{rT(1 + P)^2} & \frac{ef}{rT(1 + P)^2} - \frac{ebT}{r} & \frac{eb}{r} - \frac{m}{T} \\
\frac{erP}{rT(1 + P)^2} & \frac{ebP}{rT(1 + P)^2} - \frac{efT}{r} & \frac{ef}{rT(1 + P)^2} - \frac{ebT}{r} - \frac{m}{T}
\end{pmatrix}
$$

(23)

The origin, equilibrium $E_0$ of the PIF-system [17], is again unstable as it was for the subsystems $P$ and $F$. The eigenvalues of the Jacobian matrix (23) evaluated at the origin $E_0$

$$
J_0 = \begin{pmatrix}
1 & 0 & -\frac{aT^2}{K} \\
0 & -2\mu & 0 \\
0 & 0 & -m/T
\end{pmatrix}
$$

(24)

are $\omega_{01} = 1$, $\omega_{02} = -2\mu$ and $\omega_{03} = -m/r$, showing that $E_0$ is always an unstable equilibrium. Their respective eigenvectors $v_{0i}$, $i = 1, 2, 3$ are

$$
v_{01} = \begin{pmatrix} 1 \\ 0 \\ 0 \end{pmatrix},
\quad v_{02} = \begin{pmatrix} 0 \\ 1 \\ 0 \end{pmatrix},
\quad v_{03} = \begin{pmatrix} \frac{\lambda(m + r)}{\lambda aT^2} \\ 0 \\ 0 \end{pmatrix}
$$

(25)

Observe that when starting close to the plane $P = 0$ in $\mathbb{R}^3$, the trajectory converges finally to this equilibrium $E_0$. The situation is now more complex that in the demographic predator–prey system discussed in Section 4. The stable manifold of $E_0$ between $E_0$ and the interior attractors $E_{123}$, $L_{123}$, $T_{123}$ is now two dimensional instead of one dimensional. In the sequel we will not calculate this separatrix explicitly.

At equilibrium $E_1$ we find the following eigenvalues of the Jacobian matrix $\omega_{1i} = -2$, $\omega_{12} = \lambda K/r - 2\mu$, and $\omega_{13} = (eaK - m\sqrt{K} - \alpha T^K)/(r(\sqrt{K} + \alpha T^K))^{-1}$. Thus it is stable for

$$
m > \frac{ae\sqrt{K}}{1 + \alpha T^{\sqrt{K}}} = m^1, \quad \lambda < \frac{2\mu}{K}.
$$

(26)

At equilibrium $E_{12}$ where $F = 0$ the Jacobian matrix reads

$$
J_{12} = \begin{pmatrix}
\frac{1}{T} & \frac{3P^2}{aT^2\alpha} - \frac{P_2}{2T} & -\frac{aT^2}{\lambda(1 + P_2)} \\
\frac{2\lambda aP_3}{r^2T^2} & \frac{2\lambda a^2P}{r^2T^2} - \frac{2\mu}{2T} & -2\beta T \\
0 & 0 & \frac{eT}{rT(1 + P_2)} + \frac{ebT}{r} - \frac{m}{T}
\end{pmatrix}
$$

(27)

and the eigenvalues are explicitly evaluated as

$$
\begin{align*}
\omega_{21} &= \frac{eP_2}{rT(1 + P_2)} + \frac{ebT}{r} - \frac{m}{T}, \\
\omega_{22,23} &= 2\frac{\lambda a^2}{2\alpha T} + \frac{\sqrt{K}}{\lambda} - \frac{\lambda K}{2\alpha} - \frac{2\mu}{\lambda}(\lambda K - 2\mu)T^{\sqrt{K}}.
\end{align*}
$$

Observe that in view of feasibility (5) both eigenvalues $\omega_{22}$ and $\omega_{23}$ have negative real parts. Stability is thus ensured only by

$$
m > \frac{eP_2}{rT(1 + P_2)^{\sqrt{K}}} + \frac{ebT}{r} - \frac{m}{T}.
$$

(28)

where $P_2$ and $I_2$ are given by (18).

At equilibrium $E_{13}$ where $I_3 = 0$ the Jacobian matrix reads

$$
J_{13} = \begin{pmatrix}
\frac{1}{T} & \frac{3P^2}{aT^2\alpha} - \frac{P_3}{2T} & -\frac{aT^2}{\lambda(1 + P_3)} \\
\frac{2\lambda aP_2}{r^2T^2} & \frac{2\lambda a^2P}{r^2T^2} - \frac{2\mu}{2T} & 0 \\
0 & 0 & \frac{eT}{rT(1 + P_3)} + \frac{ebT}{r} - \frac{m}{T}
\end{pmatrix}
$$

(29)

One eigenvalue factors out, namely

$$
\omega_{31} = \frac{\lambda a^2}{2\alpha T} - \frac{2\mu}{2T} - 2\mu.
$$

The remaining $2 \times 2$ minor $\tilde{J}$ is the matrix of the PF-system given in (12) evaluated now for $P = P_3$ and $F = F_3$. The eigenvalues are

$$
\omega_{32,33} = \frac{1}{2} \left( \text{tr}(\tilde{J}) \pm \sqrt{\text{tr}(\tilde{J})^2 - 4 \det(\tilde{J})} \right).
$$

The Routh–Hurwitz sufficient condition for asymptotic stability becomes

$$
\text{tr}(\tilde{J}) = 2\frac{\lambda a^2}{2\alpha T} + \frac{aT^2}{T} (1 + P_3)^2 F_3 < 0.
$$

(30)

The second condition obviously holds, so that stability of $E_{13}$ is guaranteed if

$$
\frac{\lambda a^2}{2\alpha T} < \frac{K}{2} + \mu, \quad \frac{aT^2}{(1 + P_3)^2} + r < \frac{3rP^2}{\alpha T^2}.
$$

(31)

For the other equilibria, with symbolic manipulators it is possible to compute the expressions for the eigenvalues of the Jacobian matrix evaluated at $E_{123}$ in terms of parameters and to derive stability criteria. However, these expressions are intractable. Furthermore it is not possible to perform a symbolic analysis in the case of limit cycles (and in general for chaos) or phenomena related to global bifurcations. This holds also for the situations when extinction of one of the populations occurs in finite time. In order to complete the study we perform a numerical bifurcation analysis using the parameter values given in Table 1.

5.3. Bifurcation analysis: PIF-system

In Fig. 7 the two-parameter diagram is shown for the endemic predator–prey system PIF, where both $m$ and $K$ are varied simultaneously. The three bifurcation curves for the disease-free system, $T_{C_1}$, $H_2$ and $G^*$, were already discussed among the results presented in Fig. 6. Before we start the analysis we point out that everywhere in this diagram $E_0$ will also be an attractor when starting close to the plane $P = 0$ where $F > 0$ and $I > 0$. However, in the diagrams we indicate only the additional interior attractors. Firstly,
the healthy prey population goes extinct in finite time and thereafter the infected prey and predator populations asymptotically.

We split up the analysis in three different ranges for the parameter $K$, the low range $0 < K < 4$, the intermediate one $0 < K < 20$, and the high range $K \geq 20$.

5.3.1. Low carrying capacity $0 \leq K \leq 4$ analysis: PIF-system

We start our description of the long-term dynamics of the system for small values of $K$. A blow up of the two-parameter diagram presented in Fig. 7 is shown for the range $0 \leq K \leq 4$ in Fig. 8.

The transcritical bifurcation $TC_1$ between the origin $E_0$ and a codimension-two point $N_1$ separates the parameter space between the two equilibria $E_1$ and $E_{13}$. This is clear from Fig. 9 where the long-term state variable values are plotted for fixed $K = 1$ and varying $m$. Above the transcritical bifurcation $TC_1$ only the healthy prey population exists at equilibrium $E_1$. Between $TC_1$ and the Hopf bifurcation $H_2$ the predator and disease-free prey exist at equilibrium $E_{13}$. Below the $H_2$ the maximum and minimum values for the stable limit cycle $L_{13}$ are plotted till the collapse of the complete system at the global bifurcation point $G^\#$.

With increasing $K$ and $m$ from point $N_1$ in Fig. 8, three new transcritical bifurcations $TC_2$, $TC_3$ and $TC_4$ emerge. At $TC_2$ the prey population becomes infected, namely a predator-free system $PI$ arises: the equilibrium $E_{12}$ appears consisting of susceptible and infected sub-populations. Furthermore at both $TC_3$ and $TC_4$ the infected prey invades forming the system $PIF$ with interior positive equilibrium $E_{123}$ from the subsystems $PI$ and $PF$ respectively. Fig. 10, where $K = 4$ (instead of $K = 1$ in Fig. 9), illustrates the latter case with the transition from $PF$ to $PIF$ at $TC_4$.

5.3.2. Intermediate carrying capacity $4 \leq K \leq 20$ analysis: PIF-system

Fig. 11 is also an enlargement of a part of the two-parameter diagram of Fig. 7 for $0 \leq K \leq 20$. The interior equilibrium $E_{123}$ of the PIF-system becomes unstable at the Hopf bifurcation curve $H_3$. This Hopf bifurcation can be supercritical, denoted by $H^*_3$, giving rise

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Fig. 7. Two-parameter diagram in terms of the parameters $K$, the carrying capacity, and $m$, the natural mortality, of the PIF ecoepidemic system (17) with $I = 0$. All parameter values are given in Table 1. Table 2 gives a list of the asymptotic dynamics and Table 3 gives a list of the bifurcations.

Fig. 8. Two-parameter diagram for parameters carrying capacity, $K$, and natural mortality, $m$, of the ecoepidemic PIF system (17). This is a blow up of the diagram presented in Fig. 7 for the range $0 \leq K \leq 4$.

Fig. 9. One-parameter bifurcation diagram for the ecoepidemic PIF system (17) for the susceptible prey population, $P$, the infected prey population $I$, and the predator population, $F$ with free parameter $m$ where $K = 1$. See Fig. 2 for a description of the symbols.

Fig. 10. One-parameter bifurcation diagram for the ecoepidemic PIF system (17) showing the susceptible prey population, $P$, the infected prey population $I$, and the predator population, $F$ with free parameter $m$ where $K = 4$. See Fig. 2 for a description of the symbols.
to a stable limit cycle $L_{123}$, or subcritical, then denoted by $H^+_3$, in which case the originating limit cycle $L_{123}$ is unstable. At three so-called Bautin (or Generalised-Hopf) bifurcation points, denoted by $B_i$, ($i = 1, 2, 3$) where the switch from super- to subcritical or vice versa, takes place continuing the Hopf bifurcation, tangent bifurcation curves for limit cycles, $T_i$, ($i = 1, 2$) emerge (Fig. 11). At $K = 20$ the Hopf is supercritical (a stable limit cycle originates). Following the Hopf curve $H^+_1$ by lowering $K$, at the first Bautin point $B_1$ it becomes subcritical. The originating tangent bifurcation curve for limit cycles, $T_1$, goes via a cusp to the global bifurcation curve $G^\phi$ curve. It terminates there at a point $N_2$ where there are no infected prey, $I = 0$, and where the system reduces to the P- system and merges there with the curve $G^\phi$ of PF-system shown in Fig. 6.

Continuing on the Hopf curve $H^+_2$ in Fig. 11 again from point $B_1$, first the Hopf curve $H^+_2$ becomes supercritical at the Bautin point $B_2$, then changes again feature and becomes once more subcritical at $B_1$. Between the two Bautin points $B_2$ and $B_3$ there is a tangent bifurcation $T_2$ above the supercritical Hopf bifurcation curve $H^+_2$ (this supercritical Hopf bifurcation curve is not labeled in Fig. 11). Continuing from $B_3$, the Hopf curve terminates also at the point $N_2$ where there are no infected prey, $I = 0$, and where the system reduces to the PF-system. At that point it merges with the Hopf bifurcation $H^+_2$ of the PF-system shown in Fig. 6.

In the region bounded by the Hopf bifurcation curve in Fig. 11 there is a torus bifurcation TR where the stable limit cycle $L_{123}$ becomes unstable. This torus bifurcation terminates at point $N_4$ on the tangent bifurcation curve $T_1$. In the very small region bounded by the tangent $T_1$ between $B_1$ and $N_4$, the torus bifurcation TR starting at $N_4$ and the Hopf bifurcation $H^+_3$ between $B_1$ and $B_2$ there is tri-stability of a stable equilibrium $E_{123}$ and a stable limit cycle $L_{123}$ and zero equilibrium $E_0$.

This is explained by showing results of continuation studies presented in Fig. 12 where $K = 10$ and varying $m$ and Fig. 13 where $m = 0.42$ and varying $K$.

Comparison of the diagrams of Figs. 12 and 10 where $K = 4$ instead of $K = 10$ shows the new dynamics related to the periodic solutions, where $I > 0$ in the region $0.4 < m < 0.5$. We follow the maximum values of $I$ curve starting from the heteroclinic connection point $G^\phi$ at $m = 0.2862$. Decreasing $m$ gives a tangent bifurcation curve $T_1$ at $m = 0.2757$ and then increasing at $m = 0.43884$ a new point on the tangent bifurcation curve $T_1$ is found. Except close the second $T_1$ point, the limit cycle is unstable. The small region where it is stable is bounded by $T_1$ at $m = 0.43884$ and a torus bifurcation TR at $m = 0.42938$. The origin of the limit cycles and what happens below the TR will be explained now by studying the results in Fig. 13.

Fig. 13 shows the bifurcation sequence for increasing $K$ till $K = 20$ where $m = 0.42$ (see also Fig. 11 again). Starting from say $K = 0$ firstly part of the healthy prey populations in equilibrium becomes infected at the transcritical $TC_2$ bifurcation. Increasing $K$ at $TC_3$ the
The dynamic on the torus, the dynamics remains on the torus while during time intervals the trajectory is close to the saddle limit cycle depicted in green.

In Fig. 16 where $k = 9.91596$ on the other hand, starting on the torus shown in Fig. 15 where $k = 9.91496$, firstly the dynamics in red follows closely the torus dynamics but after passing the saddle cycle, depicted in green, it converges in blue toward the stable equilibrium $E_{123}$. Together with the torus dynamics also its basin of attraction disappears at the transition point. In other words the stable manifold associated with the saddle limit cycle loses its function as a separatrix between $T_{123}$ and $E_{123}$. Hence, beyond this bifurcation point $S$ there is only coexistence between $E_0$ and $E_{123}$.

Here we considered the case where the stable equilibrium $E_{123}$ exists, that is for $k$ values below $H_1^+$ in Fig. 11. Then, in the interval between the tangent $T_1$ and the torus bifurcation $TR$ there can occur several possibilities: tri-stability of the stable equilibrium $E_{123}$ and the stable limit cycle $L_{123}$ and a collapse of the whole system to equilibrium $E_0$ after the prey population becomes extinct in a finite time. Between $TR$ and $S$ the stable limit cycle $L_{123}$ is replaced by the quasi-periodic torus attractor $T_{123}$. Beyond $S$ there is bistability of $E_{123}$ and $E_0$.

However, in the region above the Hopf bifurcation $H_1^+$ the interior equilibrium $E_{123}$ is unstable, see Fig. 11. In the next section

predator population enters into the system and starts to feed on the two prey sub-population. Here, there is coexistence between two attractors $E_0, E_{123}$. This stable interior equilibrium $E_{123}$ becomes unstable at a subcritical Hopf bifurcation $H_1^+$ at $k = 10.6978$. The originating limit cycle is unstable and coexists with the stable equilibrium. This unstable limit cycle becomes stable at a tangent bifurcation $T_1$. In this region there is coexistence between three attractors $E_0, L_{123}$ and $E_{123}$. Following this stable branch the limit cycle becomes unstable at the torus bifurcation $TR$.

This torus bifurcation $TR$ occurs with $m = 0.42$ at $k = 9.68296$. The dynamics on the torus emerging from this torus bifurcation with increasing $k$ is quasi-periodic (characterized by two Lyapunov exponents equal zero). This can be shown by plotting the points where the trajectory intersects the Poincaré plane for $dP/dt = 0$ (see Fig. 14). The dots in the graph form closed curves when simulations continue for longer times and hence show quasi-periodic dynamics on the torus (which we will call for short torus dynamics, denoted by $T_{123}$). Fig. 15 shows the attractor $T_{123}$ in the phase-space for the three state variables on the torus for $k = 9.91496$. In Fig. 13 also the saddle limit cycle which originated from subcritical Hopf bifurcation $H_1^+$ at $k = 10.6978$ is shown. Hence in this region there is coexistence between three attractors $E_0, T_{123}$ and $E_{123}$.

When we continue following the quasi-periodic torus attractor $T_{123}$ with increasing $k$ this torus dynamics is destructed at a point labeled $S$ by the saddle limit cycle that emerges at the subcritical Hopf bifurcation $H_2^+$ at $k = 10.6978$. This phenomenon occurs between $k = 9.91496$ and 9.91596 (see Figs. 15 and 16). This bifurcation point is denoted by $S$. In Fig. 15 where $k = 9.91496$ starting...
we will study the dynamics of the system in this region in Fig. 7. These results will be important for higher carrying capacities.

5.3.3. High carrying capacity $K \geq 20$ analysis: PIF-system

For a high carrying capacity keeping $K = 20$ fixed in Fig. 17 (instead of $K = 10$ in Fig. 12) we give the one-parameter diagram by varying $m$.

Starting with a high mortality rate, say $m = 0.7$ above the transcritical bifurcation $T_{C3}$, the prey-only diseased system exists stably.

Decreasing $m$ at $T_{C3}$ the predator invades the system and the interior equilibrium $E_{123}$ is stable. It becomes unstable crossing the supercritical Hopf bifurcation $H^2_{2}$ and an interior stable limit cycle $L_{123}$ emerges. Lowering $m$ further this limit cycle becomes unstable at the torus bifurcation $TR$. It appears that the dynamics on the emerging torus is quasi-periodic. This dynamics is shown in Fig. 18 for $m = 0.628$, a value slightly below the torus bifurcation $TR$ for $m = 0.629$.

Decreasing $m$ further shows a very sudden change of the dynamics. The amplitude of the oscillations grow and also the final shape of the trajectory changes drastically. This is shown in Fig. 19 where $m = 0.57109$. We started simulations in time with initial values on the torus shown in Fig. 19 for $m = 0.5711$ and then first slowly in time the trajectory changes drastically. The amplitude especially in the $F$ direction increases and during the oscillatory dynamics the trajectory passes the origin closely and finally the system collapses, first $P$ in finite time and thereafter $F$ and $I$ exponentially. Observe that the collapse of the system is unrelated to a destruction by a saddle limit cycle as we saw in Fig. 16.

What occurs is more similar to what happened in the case of the two-dimensional $PF$ system when the heteroclinic orbit was approached, see Fig. 3 where the vector field for $K = 10$ with $m = 0.278745$ was shown. That trajectory ended in the stable limit cycle $L_{13}$ but for lower values this limit cycle was broken and the system collapses as in Fig. 5. At the switching point the trajectory lands exactly in the origin. In that situation the stable manifold $W^{s}_{0}$ terminates being a separatrix so that $E_{0}$ becomes a global attractor.

However, the situation for the three-dimensional $PIF$ system differs much from that in the two-dimensional $PF$ system due to the fact that the interaction between the infected and the predators with the healthy prey differs, see Sections 3 and 4. Now when the trajectory approaches the zero equilibrium, not only $F$ becomes small but $I$ also. This was shown in Fig. 1 for the $PI$-system and for the $PF$-system Figs. 3 and 5. The vector field in the three-dimensional state space close to the origin is shown in Fig. 20 for $m = 0.57109$ and $K = 20$.

When both $F$ and $I$ get small (blue and green arrows) the trajectory remains close to the $F = 0$ and $I = 0$ plane where $P$ increases (red arrows) toward the saddle equilibrium $E_{5}$. Thereafter by increasing $I$ and to a much lesser extent $F$ it approaches as a spiral close to the $P-I$ plane the unstable prey-only equilibrium $E_{12}$. Indeed condition (28) is not satisfied at this point while the real
parts of the eigenvalues $\omega_{22}$ and $\omega_{23}$ are negative. However, close to $E_{12}$ the predator population $F$ increases sharply and the trajectory spirals with relatively small amplitude in the $P-I$ plane toward the unstable interior equilibrium $E_{12}$. Now the amplitude of the oscillatory dynamics in the $P-I$ plane grow while the predator size $F$ decreases again. So, there is a kind of torus dynamics. But for all $m$ values where the quasi-periodic solution was not stable, we found a total collapses of the system where first the healthy prey become extinct in finite time and thereafter the infected prey and predator.

The projection of the trajectory of Fig. 20 is shown in Fig. 21. This plot shows that just before collapsing the trajectory crosses the linear tangent manifold $T_j^0$ and by assumption also the stable manifold $W^s_j$ which does not act as a separatrix anymore. Since we found this for all values below the point where the quasi-periodic torus dynamics became unstable (the extra zero Lyapunov exponent became positive) this point is catastrophic in the sense that below this curve in the two-parameter diagram the system always goes extinct.

6. Discussion and conclusions

The analysis of the prey-only system shows that the prey can be infected when the carrying capacity is above a certain threshold given in (8) independently of the mortality rate of the predator. It depends, however, on the contact rate for the transmissible and unrecoverable disease of the prey.

The demographic predator–prey model shows that the predators invade the system when the prey-only equilibrium becomes unstable, i.e. for a small enough predator mortality rate, see (11). Varying the mortality rate of the predator the predator–prey equilibrium exists whenever the predators’ mortality rate falls below a threshold (see Fig. 6). Through the prey carrying capacity, $K$, the environment always influences that threshold level. The predators too contribute to this phenomenon, since their efficiency in hunting appears in the threshold expression, i.e. through the hunting rate of predator on healthy prey, parameter $a$. Note that even in the purely demographic model besides the hunting rate the prey capture time $T$ and the conversion factor of prey into predator $e$ appear explicitly in the expression for the threshold (11).

The demographic predator–prey equilibrium becomes unstable with lowering $m$ at a Hopf bifurcation. The amplitude of the originating limit cycle grows very fast and the limit cycle disappears suddenly at a heteroclinic connection in a global bifurcation $G^e$ from a saddle disease-free prey-only equilibrium point $E_1$ to the zero-solution $E_0$ where the total system collapsed. This has already been discussed in [5,8] but this analysis is extended here by a thorough phase portrait analysis in this paper.

The linear stability analysis of the origin shows it to be always unstable, a fact that would guarantee the survival of at least some part of the ecosystem. This would occur also for the disease-free predator–prey model, thereby showing that this ecosystem behavior is due essentially to demographic reasons. The healthy prey reproduction rate provides the positive eigenvalue responsible for the origin instability. However, the study of the phase plane analysis shown in Figs. 3 and 5 reveals the existence of a sector in the phase plane, for which trajectories are doomed to end up into the origin, see Fig. 4 (see also [25]). This indicates that the actual ecosystem behavior is prone to become extinct in the region labeled $E_0$ in the bifurcation diagram (Fig. 8). In unfavorable circumstances, the prey in fact becomes extinct in finite time, followed by an exponential decay of the predators. This phenomenon is related to the presence of the square root terms in the Holling type II functional response in accordance with the findings of Braza in [5]. Here, however, we stress that our findings further indicate that the phenomenon occurs after the prey population gets extinct in a finite time.

We studied using a phase plane analysis the functioning of the separatrix of two coexisting attractors, the demographic predator–prey equilibrium $E_{13}$ or limit cycle $L_{13}$ and the zero equilibrium $E_0$ where a total collapse of the system occurred. The relationship with the heteroclinic connection was illustrated. The fact that a stable manifold is invariant disallows that a trajectory crosses this manifold. Hence, the stable manifold loses its separatrix property at the heteroclinic connection where the zero equilibrium becomes a global attractor.

There is resemblance with the dynamics of predator–prey systems with a strong Allee effect (see for instance [22]). There the quadratic logistic is replaced by a cubic growth function of the prey giving three prey-only equilibria instead of two. Also in that
predator–prey systems an unexpected collapse can occur. There this event is referred to as over-exploitation.

The bifurcation analysis reveals an organizing center point N1 in the two-dimensional parameter space in which K and m are taken as bifurcation parameters [Figs. 7 and 8]. Transcritical bifurcations, TCs, show the change in the composition of the system induced by a change in a parameter: these occur when the predator invades a prey equilibrium at TC1 (P → PF) or the prey population becomes infected at TC2 (P → PI). At the transcritical bifurcations TC3 or TC4 the prey population becomes infected (PF → PIF).

Non-equilibrium, oscillatory dynamic behaviors occur in the two parameter bifurcation diagram for the two parameters K and m. The non-equilibrium solutions emerge at Hopf bifurcations. The pattern of the originating limit cycles shows a cusp bifurcation and this gives rise to more complex dynamics together with the fact that these limit cycles become unstable at a torus bifurcation. Generally this can be the onset of chaotic dynamics. We found however only quasi-periodic dynamics originating from the torus bifurcation.

The way the torus dynamics is destructed by the saddle limit cycle (for instance shown in Figs. 15 and 16), resembles the way a limit cycle is broken by a saddle point giving a homoclinic connection of this saddle to itself (see also [4]). Here it happens in one dimension higher: the point is replaced by a limit cycle and the limit cycle by a quasi-periodic dynamics on a torus (see also [6] for more details). In [3] the destruction by a saddle-cycle, is called a homoclinic bifurcation.

We stress that these results were obtained for the parameter values given in Table 1. Because of the smoothness of the model these results are robust for small perturbations of the other parameters, that are now fixed. For larger deviations, however, the region where complex dynamics occurs can grow or even disappear and furthermore even other complex dynamics may show up. This is inherent in applying a numerical bifurcation analysis. On the other hand, the results obtained for the standard transcritical and Hopf bifurcations remain valid.

A model with diseased predators instead of infected prey was described and analysed in [8, Eq. (14)], with the three state variables: the prey, healthy predators and infected predators. Their topological structure was that of a food chain, namely infected predators have a negative effect on healthy predators and healthy predators on the prey whereby the self-regenerating prey have a negative effect on themselves. The topological structure of the ecopath studied here, system (17), is the one of an ecosystem with omnivory, namely here predators have a negative effect on both infected and healthy prey while infected prey have a negative effect on healthy prey. Hence, the infected prey and the predator are also competitors. The healthy prey population has a negative effect on itself expressed by the logistic growth because they are self-replicating. The resulting bifurcation diagrams respect these topologies (see also [13,14]). There is an organizing center N2 where the population at the lowest level, the healthy prey, can be invaded by both the other populations, infected prey and predator, either separately or together (see [13, Fig. 11 (left-bottom panel)]). This is the typical invasion of the prey by two predator populations that compete for the prey.

From the two-parameter diagram (Fig. 7) due to the weakening of the prey population by infection we conclude that the predator feeding on the prey population can exist for larger natural mortality rates of predators when prey carrying capacities are sufficient. However, with larger carrying capacities the system starts first to show oscillatory dynamics, a phenomenon related to the “paradox of enrichment” [20], and for higher values a collapse of the system occurs for a wide range of natural mortality rates where the prey population goes extinct in finite time.

References