ZERO-DIFFUSION DOMAINS IN REACTION–DIFFUSION MORPHOGENETIC AND EPIDEMIOLOGIC PROCESSES

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Classical models of morphogenesis by Murray and Meinhardt and of epidemics by Ross and McKendrick can be revisited in order to consider the colocalizations favoring interaction between morphogens and cells or between pathogens and hosts. The classical epidemic models suppose, for example, that the populations in interaction have a constant size and are spatially fixed during the epidemic waves, but the presently observed pandemics show that the long duration of their spread during months or years imposes to take into account the pathogens, hosts and vectors migration in epidemics, as well as the morphogens and cells diffusion in morphogenesis. That leads naturally to study the occurrence of complex spatio-temporal behaviors in dynamics of population sizes and also to consider preferential zones of interaction, i.e. the zero-diffusion sets, for respectively building anatomic frontiers and confining contagion domains. Three examples of application will be presented, the first proposing a model of Black Death spread in Europe (1348–1350), and the last ones related to two morphogenetic processes, feather morphogenesis in chicken and gastrulation in Drosophila.

Keywords: Morphogenesis modeling; epidemics modeling; zero-diffusion set; periodic solutions; population size dynamics; gastrulation; feather morphogenesis; Black-Death spread.

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1. Introduction

Recent advances in morphogenesis and epidemics modeling have been obtained by introducing demographic aspects, i.e., by considering morphogen, cell, pathogen, host and vector populations whose global size changes during morphogenetic, epidemic and endemic histories, as well as spatial aspects about their diffusion, spread or genetic changes [Gaudart et al., 2007, 2009, 2010a, 2010b; Glade et al., 2007; Abbas et al., 2009; Horie et al., 2010; Demongeot et al., to appear].

In epidemic studies, for example, the mathematical toolbox allowing these improvements has been introduced making classical models [Bernoulli, 1760; d’Alembert, 1761; Murray, 1763; L’Epine, 1764; de Baux, 1766; May, 1770; Lambert, 1772; Trembley, 1796; Sprengel, 1815; Ross, 1916; McKendrick, 1925; Kermack & McKendrick, 1932, 1933; Mac Donald, 1957; Barry & Gualde, 2006] more realistic, hence more convenient for predicting and anticipating the spread, and also testing scenarios (like vaccination or any health policy limiting the contagion). As applications, infectious disease dynamics, the Black Death spread during the middle age in Europe, and the dynamics of two important processes, feather morphogenesis in chicken and gastrulation in Drosophila, will be studied in the present paper.

Despite their simplicity, the models presented account qualitatively for the global shape of the endemic spatial distributions and of the morphogenetic patterns. Some perspectives will be drawn concerning the present epidemic risks: a model like that used for the Black Death spread retro-prediction would be, “mutatis mutandis”, useful to predict the dynamical behavior of the future epidemics, by considering the population fluxes along the modern aerial routes, responsible of the rapid dissemination of the pathogenic agents and infectives in the present pandemics [Khan et al., 2009].

2. Introduction to Classical Epidemiology: The Ross–McKendrick Model

In the seminal work by Bernoulli [Bernoulli, 1760; Dietz & Heesterbeek, 2000, 2002; Zeeman, 1993] proposed for explaining the small pox dynamics, the population was divided into susceptibles (not yet been infected) and immunes (immunized for the rest of their life after one infection). In [Ross, 1916; McKendrick, 1925; Kermack & McKendrick, 1932, 1933] is proposed a more sophisticated model called Susceptible/Infective/Recovered with immunity (SIR) model, with Eqs. (1):

\[
\begin{align*}
\frac{dS}{dt} &= -\beta SI + (\delta + \gamma)S - \delta S, \\
\frac{dI}{dt} &= \beta SI - (\nu + \delta)I, \\
\frac{dR}{dt} &= \nu I - (\delta + \gamma)R,
\end{align*}
\]

Or equivalently:

\[
\begin{align*}
\frac{dS}{dt} &= -\delta S + \delta(S + I + R) - \delta S + \gamma R \\
\frac{dI}{dt} &= +\beta SI - (\delta + \nu)I \\
\frac{dR}{dt} &= \nu I - (\delta + \gamma)R
\end{align*}
\]

where \( S \) (resp. \( I, R \)) denotes the size of Susceptible (Infective, Recovered) population with \( S + I + R = N \), \( \beta \) (resp. \( \delta, \gamma, \nu \)) being the contagion (resp. death/birth, loss of resistance, immunization) rate.

The epidemic parameter \( R_0 = \beta N/(\nu + \delta) \) is the mean number of secondary infecteds by one primary infective and predicts, if it is greater than 1, the occurrence of an epidemic wave. By defining age classes \( S_i, I_i \) and \( R_i \) \((i = 1, \ldots, n)\) in each subpopulation \( S, I \) and \( R \), we have at any stationary state \((S^*, I^*, R^*)\) the following relationships:

\[
\begin{align*}
u^*(i) &= \frac{S^*}{S^*_1}, \quad \nu^*(i) = \frac{I^*}{I^*_1}, \quad w^*(i) = \frac{R^*}{R^*_1}
\end{align*}
\]

where the probabilities for a newborn being alive and either susceptible \( u^*(i) \), infected \( \nu^*(i) \) or immune \( w^*(i) \) at age \( i \) make the link between the Bernoulli and the Ross-McKendrick models, but the weakness of the latter still resides in many insufficiencies and approximations:

- when the population size of either susceptibles or infectives tends to be very large, the quadratic term \( SI \) has to be replaced by a saturation term (e.g. Michaelian) \( SI/[(k + S)(j + I)] \)
- the immunized infectives or healthy carriers are neglected
- the total population size is supposed to be constant, the fecondity just equalling the natural mortality. The Bernoulli model is taken implicitly
into account the fecundity, and explicitly the natural mortality. The model by d’Alembert [1761] improved that of Bernoulli’s by distinguishing the specific mortality due to the infectious disease from the natural one, being more widely applicable than the model by Bernoulli which was restricted to immunizing infections, d’Alembert’s method requires the knowledge about the survival function after eliminating the specific cause of death due to the infectious disease, but Bernoulli’s approach provides more insight for a mechanistic interpretation of infection.

- variables and parameters do not depend on space (no migration)
- parameters do not depend on time (no genetic adaptation of infectious agent or human population, even very slow compared to the fast dynamics of epidemics).

We will improve in the following the Ross–McKendrick model by trying to partially compensate these defects. We will first introduce the age classes into the host population in order to account for its growth, the space dependence in order to account for the host and vector population migration and their possible coexistence, before presenting an example of application and drawing perspectives. The genetic changes resulting from the adaptation of the concerned populations will be not treated in this paper.

3. Epidemics Modeling with Demography and Diffusion

By exploiting the remarks formulated in the previous section about the classical models of epidemic modeling, we now consider the possibility to merge the demographic dynamics introduced in [Demengeot, 2006] and the reaction-diffusion, by simplifying the von Foerster dynamics: we suppose that the biological age is identical to the chronological age.

For the sake of simplicity, we consider the problem as isotropic and the space coordinate $s$ as unidimensional.

**Proposition 1.** Let us consider the Fisher equation defined by:

$$\frac{\partial n}{\partial t} = \Delta n + n(K - n)$$

with the initial conditions: $n_i(0,0) = \delta_i$, where $\delta_i$ is the Dirac’s distribution in 0. Its asymptotic solution (t tending to infinity) is given by:

$$n(s,t) = K \exp \left( \frac{s^2}{4t} - \frac{\exp(-Kt)}{K} \right) (4\pi t)^{-1/2}$$

**Proof.** Let us suppose that $\frac{\partial n}{\partial t} = \Delta n + n(K - n)$. For $\frac{s}{t} \simeq 1$ (resp. $\frac{s}{t} \simeq 0$), we have:

$$n(K - n) \simeq -nK \ln \left( \frac{n}{K} \right) \quad \text{(resp.} \simeq nK).$$

Then, if we consider the solution $n_1$ of the heat operator, with $n_1(\cdot,0) = \delta_0$, and the solution $n_2$ of the logistic equation

$$\frac{\partial n_2}{\partial t} = -n_2 K \ln \left( \frac{n_2}{K} \right)$$

with $n_2(\cdot,0) = \delta_0$, we have:

1. If $n_1(s,t) = \exp(-\frac{s^2}{4t})(4\pi t)^{-1/2}$, then $\frac{\partial n_1}{\partial t} = -s^2 \frac{n_1}{2t}$, hence

$$\Delta n_1 = \frac{\partial^2 n_1}{\partial s^2} = \frac{n_1}{2t} + s^2 \frac{n_1}{2t}^2$$

and

$$\frac{\partial n_1}{\partial t} = \left( \frac{s^2}{4t} - \frac{1}{2t} \right) n_1 = \Delta n_1$$

2. If $\frac{s}{t} \simeq 1$ (resp. $\frac{s}{t} \simeq 0$), $n_2(s,t) = K \exp(-\exp(-Kt)/K)$ (resp. $n'_2(s,t) = \exp(Kt)$) is a solution of

$$\frac{\partial n_2}{\partial t} = -n_2 K \ln \left( \frac{n_2}{K} \right) \quad \text{(resp.} \frac{\partial n'_2}{\partial t} = n'_2 K).$$

Let us consider now $n = n_1n_2$: we have:

$$\frac{\partial n}{\partial t} = \begin{cases} 
\frac{n_1}{n} \frac{\partial n_1}{\partial t} + \frac{n_2}{n} \frac{\partial n_2}{\partial t} \\
= n_1 \Delta n_1 - n_1 n_2 K \ln \left( \frac{n_2}{K} \right), \quad \text{if} \ \frac{n_2}{K} \simeq 1 \\
n_2^2 \Delta n_1 - n_1 n_2^2 K, \quad \text{if} \ \frac{n_2}{K} \simeq 0
\end{cases}$$
Because \( n_2 \) (resp. \( n_2^* \)) is independent of \( s \), then 
\[ n_2^* \Delta n_1 = \Delta n \] (resp. \( n_2^* \Delta n_1 = \Delta n \)) and we have:

\[
n_1 n_2 K \ln \left( \frac{n_1}{K} \right) = K \exp \left( \frac{-2 \exp(-Kt)}{4} \right) \times \left[ -\frac{s^2}{4t} \frac{\ln(4\pi t)}{2} - \ln(K) \right] (4\pi t)^{-1/2}
\]
tends to 0 when \( t \) tends to infinity, for every \( s \).

Then

\[
n = n_1 n_2 = K \exp \left( \frac{s^2}{4t} \frac{\exp(-Kt)}{K} \right) (4\pi t)^{-1/2}
\]
is the approximate asymptotical solution of the Fisher equation.

It is the same for

\[
n_1 n_2^* K \ln \left( \frac{n_1}{K} \right) = K \exp \left( Kt \frac{s^2}{4t} \right) \times \left[ -\frac{s^2}{4t} \frac{\ln(4\pi t)}{2} - \ln(K) \right] (4\pi t)^{-1/2}
\]
tends to 0, when \( t \) tends to 0. Then

\[
n = n_1 n_2^* = K \exp \left( \frac{s^2}{4t} \frac{\exp(-Kt)}{K} \right) (4\pi t)^{-1/2}
\]
is asymptotically in \( t \) the solution of the Fisher equation \( \frac{dn}{dt} = \Delta n + n(K - n) \), \( n = n_1 n_2^* \) being the approximate solution when \( t \) is small. \( \blacksquare \)

If we consider that the diffusion and the demographic growth are slow compared to the fast epidemic dynamics, then the initial condition of the Fisher equations is the stable steady state of the reaction part of the system of Eqs. (4), defined by:

\[
\begin{align*}
\frac{\partial S}{\partial t} &= \Delta S + S(K_1 - S) - \frac{bSI + k_1 S - k_3 I}{\varepsilon} \\
\frac{\partial I}{\partial t} &= \Delta I + I(K_2 - I) + \frac{bSI - k_3 I}{\varepsilon}
\end{align*}
\]  

(4)

**Proposition 2.** If we denote the fast endemic steady state of (4), supposed to be stable, by \( (S^*, I^*) \), then we have as asymptotic (in \( t \)) solution of (4) the Gaussian functions:

\[
\begin{align*}
S(s, t) &= S_0^* \exp \left( -\frac{s^2}{4t} \frac{\exp(-K_1 t)}{K_1} \right) (4\pi t)^{-1/2} \\
I(s, t) &= I_0^* \exp \left( -\frac{s^2}{4t} \frac{\exp(-K_2 t)}{K_2} \right) (4\pi t)^{-1/2}
\end{align*}
\]

for \( t \geq t^* \), where \( t^* \) denotes the first time where \( \langle S(s, t^*), I(s, t^*) \rangle \) approximates \( (S^*, I^*) \) with a precision equal to \( \varepsilon \) in Euclidean norm:

\[
((S(s, t^*) - S^*)^2 + (I(s, t^*) - I^*)^2) = \varepsilon
\]

**Proof.** The result is the direct consequence of Proposition 1 and of the fastness of the epidemic dynamics. \( \blacksquare \)

There is an asymptotic identity between the zero-diffusion lines of the susceptibles and of the infecteds, if the asymptotic mode and variance of their Gaussian diffusion functions are the same. When the contagion coefficient \( b \) is considered as depending on space, we choose \( b(s) \) as maximal, equal to \( b^* \), on zero-diffusion lines, i.e. where susceptibles and infecteds have the maximum chance to coexist and we denote the steady state values of the fast epidemic dynamics (for \( b^* \)) as \( S^{**} \) and \( I^{**} \).

If there is no asymptotic identity, we can define \( b(s) \) as inversely proportional to the distance between \( s \) and the zero-diffusion set located around the concentration peaks.

Let us suppose now that the fast dynamics are the demographic and epidemic ones and that they are driven by the following differential equations:

\[
\begin{align*}
\frac{dx}{dt} &= kx(N - x) - C_1 \frac{xy}{(K + y)} - k_1 x \\
\frac{dy}{dt} &= Cy + C_2 \frac{xy}{(K + y)} - k_2 y
\end{align*}
\]

(5)

where the host population growth (its size being represented by the variable \( x \)) is logistic, the fecundity being limited by a Malthusian term depending on the maximal population size \( N \). The contagion
interaction is supposed to have a Michaelian saturation term [Crauste et al., 2008] for controlling a possible excess of infective vectors (whose population size is $y$), and the mortality is assumed to be different between host and vector. By denoting $a = k(N - k_1)$ and $b = -f + k_2$, the system (5) becomes:

$$\begin{align*}
\frac{dx}{dt} &= -kx^2 - C \frac{xy}{(K + y)} + ax \\
\frac{dy}{dt} &= C \frac{xy}{(K + y)} - by
\end{align*}$$

(6)

**Proposition 3.** The steady states of the system (6) are of three types:

- a stable node (resp. focus) $(x^*, y^*)$, in case of small mortality of infecteds ($b \ll 1$), if
  $$a > 1 \quad \text{and} \quad (C - a)(a - 1) > 2kb$$
  (resp. $a < 1, 4(1 - a)C > ab$)
- a stable node, in case of fast epidemics ($C \gg K$ and $a = b = C$), if $k \leq 1$.
- a neutral steady state $(x^{**}, y^{**}) = (0,0)$.

**Proof.** The Jacobian matrix $B^*$ of the system (6) at the steady state $(x^*, y^*)$ is equal to:

$$B^* = \begin{pmatrix}
-2kx^* - b \frac{y^*}{x^*} + a & -b \frac{C}{(K + y^*)} \\
0 & b \frac{K}{(K + y^*)} - b
\end{pmatrix}$$

The nonzero stationary is defined by:

$$x^* = \frac{b}{C}(K + y^*) \quad \text{and} \quad kx^2 + (C - a)x^* - bK = 0,$$

therefore the only positive solution, distinct from the saddle, is given by:

$$x^* = \frac{a - C + \sqrt{(a - C)^2 + 4bkK}}{2k}.$$

$$y^* = \frac{C}{b}x^* - K.$$

Therefore, the characteristic polynomial of the matrix $B^* - \lambda I$, denoted $P_{B^*}$, is equal to:

$$P_{B^*}(\lambda) = \lambda^2 + \left[2kx^* + b \frac{y^*}{x^*} + b - a - b \frac{K}{(K + y^*)}\right] \lambda + ab \frac{K}{(K + y^*)} - ab - 2kx^*$$

$$\times \left[\frac{K}{(K + y^*)} - b\right] + b^2 \frac{y^*}{x^*} = 0$$

and, because

$$bK \left(\frac{b}{C} - 1\right) - (a + b - C)x^*,$$

we have:

$$2x^*\lambda = bK \left(\frac{b}{C} - 1\right) - (a + b - C)x^*$$

$$\pm \left(\frac{bK}{b} - 1\right) - (a + b - C)x^*\right)^2$$

$$+ 8kx^2 \left(\frac{b^2K}{C} - bx^*\right) - 4b^2y^*x^* - 4aba^* \left(bK \frac{K}{C} - x^*\right)^{1/2}.$$

We also have

$$8kx^2 \left(\frac{b^2K}{C} - bx^*\right) - 4b^2y^*x^* - 4aba^* \left(bK \frac{K}{C} - x^*\right)$$

$$= 4bx^* - 2kx^2 + x^* \left(2bkK \frac{K}{C} - C + a\right)$$

$$- bK \left(\frac{a}{C} - 1\right)$$

and

$$2C\left(b - C\right) - C(a + b - C) = \left[bK(b - C) - C(a + b - C)\right]^2$$

$$+ 4bC\left[2Ck^2x^2 + 2bkK \right]$$

$$+ C(a - C)x^* - bK(a - C)]^{1/2}.$$

Hence, we have:

(1) if $b \ll 1$ such as $bK/(C - a) \ll 1$, then we have:
$x^* \simeq b \frac{K}{C - a} > 0$, \quad $y^* \simeq a \frac{K}{C - a} > 0$, \quad $6K \left( \frac{b}{C - 1} \right) - (a + b - C)x^* \simeq -ab^2 \frac{K}{C} (C - a) < 0$ \quad and

$$8kz^2 \left( \frac{K}{C} - kx^* \right) - 4k^2 y^* x^* - 4axz^2 \left( \frac{K}{C} - x^* \right) \simeq -4ab \frac{K^2}{C(C - a)^2} \left( \frac{bK}{C} - a \right) + 1 - a$$

$(x^*, y^*)$ is a stable node ($a > 1$ and $(C - a)(a - 1) > 2kb$) or focus ($a < 1$ and $4a(1-a)b^2 K^2 > a kb^4 K^2 / C$, i.e. $4(1-a)C > ab$). For example, if we choose $a = b = C/3 = 4K = 1/3$ and $k = 1$, then: $x^* = 0.236/6$, $y^* = 0.035$ and the $B^*$ eigenvalues are given by:

$$\lambda = -0.0054 \pm [0.0054^2 + 0.052 - 0.0031 - 0.0238 + 0.0093]^{1/2}$$

and $(x^*, y^*)$ is a stable focus.

(2) if $a = b = C \gg K$ and $k \leq 1$, then $x^* = (CK/k)^{1/2}$, $y^* = (CK/k)^{1/2} - K$ and the $B^*$ eigenvalues are:

$$\lambda = \left[ C^2 \pm C \left( 1 - 8 \left( \frac{Kk}{C} \right)^{1/2} \right) \right]^{1/2} \times \left( 1 - \left( \frac{Kk}{C} \right)^{1/2} \right)^{1/2}.$$

Then $(x^*, y^*)$ is a neutral steady state.

Suppose that the contagion interaction has a double Michaelian saturation term for controlling both a possible excess of susceptibles (whose population size is $x$) and of infective vectors (whose population size is $y$), then system (6) becomes:

$$\frac{dx}{dt} = -kx^2 - C \frac{x y}{(J + x)(K + y)} + ax \quad (7)$$

$$\frac{dy}{dt} = C \frac{x y}{(J + x)(K + y)} - by$$

**Proposition 4.** The steady states of the system (7) are of three types:

- a stable node (resp. focus) $(x^*, y^*)$, in case of large saturation of susceptibles ($J \gg 1$) and of small mortality of infecteds ($b \ll 1$), if

\[
\begin{align*}
\alpha &> 1 \quad \text{and} \quad \left( \frac{C}{J} - a \right)(a - 1) > 2kb \\
\text{(resp.} \quad \alpha < 1 \quad \text{and} \quad 4(1-a)b^2 K^2 > a kb^4 K^2 / C, \quad \text{i.e.} \quad 4(1-a)C > ab) \\
\quad \text{a neutral steady state} \quad (x^*, y^*) = (0,0)
\end{align*}
\]

**Proof.** Let us prove the second assertion. If $k \ll 1$, the $x^2$ term is negligible, and $(x^*, y^*)$ verifies:

$$\frac{C x^* y^*}{a(J + x^*)(K + y^*)} \simeq x^* \simeq y^*$$

hence

$$C x^* \simeq a(J + x^*)(K + C x^*) \quad = aJC + a(JC + K)x^* + aC x^*$$

and

$$aJC x^* + (aJC + aK - C)x^* + aJC = 0.$$

Then, if we denote $D = C - aJC - aK > 0$, we have:

$$D^2 - 4a^2 CJK = (C + aJC - aK)^2 - 4aC J$$

$$> (C - aK)^2 - 4aC K$$

$$> (C - 0.5)^2 - 2C$$

$$> 0$$

and

$$2aJK > x^* = \frac{D - (D^2 - 4a^2 CJK)^{1/2}}{2aC} > 0.$$

Then the eigenvalues $\lambda$ of the Jacobian matrix $B^*$ of system (7) at the stationary point $(x^*, y^*)$
are given by \( \text{det}(B^* - \lambda I) = P_{22} - (\lambda) = 0 \), where:

\[
B^* - \lambda I = \begin{pmatrix}
-CJ (J + x^*)^2 (K + y^*) & \frac{y^* (J + x^*)^2 (K + y^*)}{a - \lambda} \\
-CJ (J + x^*)^2 (K + y^*) & \frac{-CK (K + y^*)^2 (J + x^*)}{b - \lambda}
\end{pmatrix}
\]

\[
= \begin{pmatrix}
\frac{a x^*}{J + x^*} - \lambda & \frac{-bK}{K + y^*} \\
\frac{bK}{K + y^*} & \frac{-CK x^* (J + x^*)}{b - \lambda}
\end{pmatrix}
\]

Fig. 1. Attractor of the system with double saturation (7) showing (top) a limit cycle and attractor of the system with a unique saturation term (6) showing (bottom) a stable focus for different initial conditions and for the same set of values of parameters in cases (6) and (7) except for the affinity coefficient \( J \) (cf. for the numerical simulations the web site: http://www.zweigmedia.com/RealWorld/deSystemGrapher/func.html).
Hence, we have for the characteristic polynomial $P_B$ of $B^*$:

$$P_B(\lambda) = \lambda^2 - \left[\frac{a\lambda^*}{J + \lambda^*} - \frac{b}{K + \gamma}\right] \lambda + ab \frac{KJ - \lambda^* \gamma}{(J + \lambda^*)^2(K + \gamma^*)}.$$  

Then the eigenvalues $\lambda$ are complex with a positive real part and give birth to a limit cycle after a Hopf bifurcation (cf. Fig. 1), because: $K > J$ implies

$$\frac{a\lambda^*}{J + \lambda^*} - \frac{b}{K + \gamma^*} > 0$$

and since $\lambda^* < 2aJK$ with $a < 1/(2K) < 1/(2J)$, we obtain:

$$\frac{4abKJ}{(J + \lambda^*)(K + \gamma^*)} > \frac{4a^2JK}{C^2}\frac{aK + aJ + 2a\lambda^*}{C^2}$$

$$= \left(\frac{a\lambda^*}{J + \lambda^*} + \frac{b}{K + \gamma^*}\right)^2.$$  

The other results of Proposition 4 are proved as in Proposition 3.

4. Definition of the Biological Age

By introducing a biological age different from the chronological age $t$ of the demographic dynamics [Demongeot, 2009], we replace the logistic term in Eqs. (5)–(7) by a von Foerster-like partial differential equation, where we denote by $\sigma_x$ the biological age shift of an individual susceptible with respect to its chronological age $t$:

$$\sigma_x x_i(a,t) + x_i(a,t) = -\mu_x x_i(a,t),$$  

where $x(a,t)$ is the number of susceptibles in biological age $a$ at time $t$.

If an ageing acceleration $\gamma_S$ of an individual with respect to its chronological age $t$ is allowed and if the individuals can diffuse, a generalized von Foerster’s equation can be used in the continuous case of size structured populations. In the discrete case of age-class structured populations, the diffusion has to be replaced by a discrete progression in age (stepwise ageing) [Demongeot, 2009],

$$\sigma_x x_i(a,t) + \Box x + x_i(a,t) = -\mu_x x_i(a,t),$$  

where the demographic d’Alembertian operator is equal to $\Box x = \gamma_S \Box x/\partial a^2 - \Delta x$ and where $\mu_x$ is the natural mortality coefficient of the susceptibles. The values of parameters like $\sigma_x$, $\gamma_S$ and $\mu_x$ can depend both on space, biological age and time.

5. Introduction of a Spatial Dynamics

The introduction of the space in Ross-McKendrick models can be done through stochastic spatial Markovian or renewal models [Demongeot, 1985; Demongeot & Fricot, 1986; Ivanoff & Merzbach, 2006; Davies & Braun, 2010, to appear] or through deterministic Partial Differential Equations (PDE’s) in which the diffusion of hosts or vectors is modeled by the Laplacian operator $\Delta$ or possibly by the d’Alembertian $\Box$, when some subpopulations can present an accelerated ageing [Demongeot, 2009]. These models are called SIR with Diffusion (SIRD) [Gaudart et al., 2010a]. During the susceptible and infective vector spread, the maximum of contagion is observed on the common zones of least diffusion, which can be asymptotically identical, the common zero-diffusion domains allowing a maximum of contacts between interacting species (cf. Fig. 2), i.e. reducing the effect of the thermic fluctuations which give birth to large values of the diffusion coefficients.

Taking into account the diffusion of all vector subpopulations (vector susceptibles, infecteds/noninfectives and infectives), it is possible to simulate a model and compare its numerical results to the data recorded on the ground. For improving the fit, we take into account the diffusion of the human subpopulations $S$, $G$ and $R$ (susceptibles, infecteds/infectives and recovereds).

The contagion parameters are chosen depending on space, e.g. being maximum in zones where diffusion of infective vectors ($A_i$) and susceptible hosts ($S$) is minimum and in zones where sizes $S$ of susceptible hosts and $A_i$ of infective vectors are maximum, where for example $D_S/S$ and $D_{A_i}/A_i$ are minimum, ensuring locally a large coexistence time, i.e. a high contagion rate between interacting subpopulations. In case of isotropic diffusion, the zero-diffusion or zero Laplacian (or zero curvature or maximal gradient) lines of the concentration surfaces of the concerned populations are, if they are identical (cf. Fig. 2), such a contagion domain,
where hosts, vectors and infectious agents interact. These lines correspond to regions where the mean Gaussian curvature of the concentration surfaces $S$ and $A_i$ vanishes, these line regions being defined respectively by

\[
\frac{\partial^2 S}{\partial x^2} \frac{\partial^2 S}{\partial y^2} \left( \frac{\partial^2 S}{\partial x \partial y} \right)^2 = 0, \quad \text{and by}
\]

\[
\frac{\partial^2 A_i}{\partial x^2} \frac{\partial^2 A_i}{\partial y^2} \left( \frac{\partial^2 A_i}{\partial x \partial y} \right)^2 = 0.
\]

We can show the possibility of intersection of these lines on one tangency point or on two points (cf. Fig. 2) or on whole zero-diffusion sets if they are asymptotically identical (cf. Fig. 6, bottom) for some values of the ratio between diffusion coefficients $D_S/D_{A_i}$ [Michon et al., 2008].

6. An Example of Application: The Black Death in Europe Between 1348 and 1350

Plague was considered as endemic in the steppes of Southern-Russia where Mongols originated [Zhang et al., 2008]. Born in the Caspian sea area (probably triggered by contacts between Mongolian and Genoa sailors and warriors during wars around 1346), epidemic wave went through the Mediterranean routes (cf. Fig. 3). It reached ports like Marseilles in France and Genoa in Italy at the end of the year 1347. In five years, it spread widely in Europe from two large commercial cities and returned to the Caspian reservoir. A simple Susceptible-Infective-Recovered model with Diffusion (SIRD) explains the essential of the observed front wave dynamics during the years between 1348 and 1350 [Gaudart et al., 2010a]. The model uses only three coefficients: (i) a local viscosity proportional to the altitude, (ii) a contagion parameter and (iii) a death/recovering parameter (representing the future of infecteds/infectives as dead or immunized after being cured of the plague).

The Fisher equation [Fisher, 1937; Murray, 2003] was first used for representing the evolution of the host and vector subpopulations during the spread of the Black Death.

The model used for modeling the Black Death spread is a SIRD model as in the Bankoumana study [Gaudart et al., 2007, 2009, 2010a], but without vector terms and has for its reaction term the form of a Lotka–Volterra Ordinary Differential Equation (ODE) of dimension 3, plus a diffusion term:

\[
\frac{dS}{dt} = \varepsilon \Delta S - \beta SI,
\]

\[
\frac{dI}{dt} = \varepsilon \Delta I + \beta SI - \gamma R,
\]

\[
\frac{dR}{dt} = \varepsilon \Delta R + \gamma R,
\]

where $\beta SI$ term comes from the “law of mass action”, assuming homogeneous mixing between susceptibles and infecteds, $\beta$ being the rate of transition from susceptible to infected state calculated per infected and per susceptible, $\gamma$ is the rate of transition from infected to post-infected state (e.g. death or immunity) per infected person and $\varepsilon$ is the diffusion coefficient. By taking the viscosity (inverse of $\varepsilon$) proportional to the altitude, the simulated front waves are more similar to the observed.
ones (cf. Fig. 3) than in the previous simulations [Murray, 2003]. The initial population size of susceptibles in the main middle age cities has been fixed following the demographic data. The results of simulations (cf. Fig. 3, bottom) are in agreement with the data observed in the 370 hospitals of the order of St. Anthony (cf. Fig. 3, top right). Improvements could come from considering multiple entrance points (ports like Barcelona reached in June 1348 or La Rochelle, Rouen and Dover reached later in 1348), and taking into account all the commercial sea (Mediterranean and Atlantic) and overland routes (cf. Fig. 3, top left) as well as the demography (fecundity and natural mortality). The present endemic state (cf. Fig. 4) could be explained by a new model taking into account the air routes [WHO, 1999]. An efficient prediction from simulations of a realistic model considering new aerial routes with a minimal viscosity [Khan et al., 2009] could serve this cause. Another improvement could come from introducing saturation effects like those made explicit in system (7). The contagion parameter $\beta$ could also be chosen depending on space, e.g. maximum in zones which constitute overlaps between domains where diffusion of infective vectors and hosts is minimum and domains where concentration of susceptibles is maximum, ensuring locally a large coexistence time, hence a high contagion rate between these large interacting subpopulations.
7. The Feather Primordia Morphogenesis

The feather primordia morphogenesis is an embryonic process, which allows to well position adult feathers permitting, for example, the peacock to do the wheel in order to attract the female (cf. Fig. 5).

The reaction–diffusion system corresponding to the feather primordia morphogenesis [Michon et al., 2008] rules three variables, the density \( n \) of migrant primordial cells and the concentration \( u \) (resp. \( v \)) of an activator (resp. inhibitor), the BMP-7 (resp. BMP-2), following the equations:

\[
\begin{align*}
\frac{\partial n}{\partial t} &= -\Delta n - b n + \nabla(\chi n \nabla u), \\
\frac{\partial u}{\partial t} &= D_u \Delta u + f_0(u, v) - k_u u, \\
\frac{\partial v}{\partial t} &= D_v \Delta v + g_0(u, v) - k_v v,
\end{align*}
\]

with \( f_0(u, v) = c_1 u v^2/(1 + v) \), \( g_0(u, v) = c_2 u v^2 \), \( n(x, 1, t) = \int_0^t 2Q(x, a, t)\, \text{d}a \), and also with Neumann boundary conditions. For the sake of simplicity, we will use in the following a simplified equation for \( n \):

\[
\frac{\partial n}{\partial t} = D_n \Delta n - \nabla(\chi n \nabla u) + bn(1 - n). \tag{12}
\]

Then it is possible to derive explicitly Turing’s instability necessary conditions [Turing, 1952], where \( u \) (resp. \( v \)) denotes the stationary concentration of \( u \) (resp. \( v \)) and \( f_{ub} \) (resp. \( g_{ub} \)) the first derivative of \( f_0 \) (resp. \( g_0 \)) with respect to \( u \) at \((u, v)\),

\[
\begin{align*}
(1) \quad f_{ub} + g_{ub} < 0 & \Rightarrow 2c_1 k_u u_0/(k_u + c_2 u_0^2) - k_u - k_v < 0, \\
(2) \quad f_{ub}g_{ub} - f_{ub}g_{vb} > 0 & \Rightarrow -2c_1 k_u^2 u_0/(k_u + c_2 u_0^2) + k_u k_v + 2c_1 c_2 u_0^2/(k_u + c_2 u_0^2)^2 > 0 \Rightarrow -2c_1 k_u^2 u_0/(k_u + c_2 u_0^2)^2 + k_u k_v > 0, \\
(3) \quad D_{ub}g_{ub} + D_{vb}f_{ub} < 0 & \Rightarrow 2D_u c_1 k_u u_0/(k_u + c_2 u_0^2) - D_u k_u - D_u k_v < 0, \\
(4) \quad (D_{ub}g_{ub} + D_{vb}f_{ub})^2 & > 4D_u D_v (f_{ub}g_{ub} - f_{ub}g_{vb}) \Rightarrow 2D_u c_1 k_u u_0/(k_u + c_2 u_0^2) - (D_u k_u + D_u k_v) > (4D_u D_v c_1 k_u u_0/(k_u + c_2 u_0^2))^2/4.
\end{align*}
\]

If \( v \ll 1 \) and \( n \) are near their stationary value, e.g. if \( D_u, \chi \) and \( b \) are large, such that the system reaches rapidly its slow \((u, v)\) manifold, we can decompose
the two last equations of (11) in order to get a potential-Hamiltonian system, with:

\[
\begin{align*}
\frac{\partial u}{\partial t} &= -\frac{\partial P}{\partial u} + \frac{\partial H}{\partial v}, \\
\frac{\partial v}{\partial t} &= -\frac{\partial P}{\partial v} - \frac{\partial H}{\partial u},
\end{align*}
\]

\[
P = \frac{k_u u^2 + k_v v^2}{2},
\]

\[
H = c_1 n u^3 \ln(1 + v) - c_2 n^3 v^3.
\]

Then \(c_1\) and \(c_2\) (resp. \(k_u\) and \(k_v\)) can be considered as frequency (resp. amplitude) modulating parameters [Demongeot et al., 2007a, 2007b; Forest et al., 2007; Glade et al., 2007] and the synchronizability can be estimated by considering the isochrons landscape of the simplified system [Demongeot & Francoise, 2006; Ben Amor et al., 2010].

The very last important parameter is the ratio between the diffusion coefficients \(D_u/D_v\), which is less than 1 as usually in lateral inhibition [Demongeot et al., 2009]: if the ratio is equal to the critical value 0.06, we observe both in experiments (cf. Fig. 5) and in simulations (cf. Fig. 6) a temporal-spatial synchrony between the effectors \(u\) and \(v\). Both experiments and simulations show a coincidence of their remarkable Gaussian lines, i.e. the projections of the null-curvature lines on the \(u\) and \(v\) concentration surfaces, defined by the following equations expressing the vanishing of the mean Gaussian curvature:

\[
C_u(x, y, t) = \frac{\partial^2 u}{\partial x^2} \frac{\partial^2 u}{\partial y^2} - \left(\frac{\partial^2 u}{\partial x \partial y}\right)^2 = 0,
\]

\[
C_v(x, y, t) = \frac{\partial^2 v}{\partial x^2} \frac{\partial^2 v}{\partial y^2} - \left(\frac{\partial^2 v}{\partial x \partial y}\right)^2 = 0.
\]

These two remarkable lines for the effectors \(u\) and \(v\) coincide for the critical value of \(D_u/D_v = 0.06\) (cf. Fig. 6). The 2D projections of these lines form front waves moving in the same direction as the fronts of the concentration contour lines, and where they coincide, the diffusion term vanishes and \(u\) and \(v\) are susceptible to form at this location a self-assembly like the phospho-lipo-proteic plasmic membrane or the inner mitochondrial membrane [Demongeot et al., 2007c]. The coexistence at this common least diffusion location of migrant cells \(n\) as well as

![Fig. 5. Feather morphogenesis with identification of an activator \(u\) (BMP-7), an inhibitor \(v\) (BMP-2) and a mediator (Follistatine) as morphogens (left) interacting at the genetic level, where \(G_u\) (resp. \(G_v\)) and \(O_u\) (resp. \(O_v\)) denote the gene coding for \(u\) (resp. \(v\)) and its operator (top middle and bottom right) for giving first feathers primordia and after, adult feathers allowing the wheel of feathers in the peacock (top right).](image-url)
Fig. 6. Coincidence of the null-curvature lines of $u$ (in red) and $v$ (in blue) concentration surfaces, when $D_u/D_v$ varies from $0.05$ (left) to $0.07$ (right). For $D_u/D_v = 0.06$ (middle), the coincidence is perfect on the central part of the picture, which corresponds roughly to the experimental value of the diffusion coefficients ratio.
Fig. 7. On the four thumbnails (left) diminishing $k$ (from 35 to 0 with step of 5) causes the decrease of the feather number and amplitude. On the four thumbnails (right) diminishing $c_2$ (from 4500 to 1200 with step 1100) causes rather motif disparition and diffusion wins over reaction.
morphogens \( u \) and \( v \) permits indeed to build an anatomic boundary for the future feathers, avoiding chemical reactions between these components, which change their physical nature and involve thermal fluctuations (hence no zero diffusion). These phenomena are summarized in Fig. 6 which shows the coincidence (or the spatial synchrony) between the remarkable lines in 2D, suggesting that this mechanism can be permitted in many circumstances of formation of an anatomic boundary: for example, in [Demongeot et al., 2007c], a lateral inhibition mechanism is also used to show a spatial synchrony between transmembrane proteins (the ATPase and the Translocase) allowing the realization of a variational principle, which maximizes the mitochondrial ATP production and minimizes the mean free path of adenylates inside the mitochondrial membrane, by favoring the spatial vicinity between the ATPase and Translocase sites inside the inner mitochondrial membrane. Many other parameters like \( c_2 \) and \( k_v \) are critical for the occurrence of feathers (cf. Fig. 7).

8. The Gastrulation Process

The gastrulation process is critical for a living organism, because it initiates the construction of the digestive tube, just before the neural chord (cf. Figs. 8 and 9). Many experimental observations show that invagination preceding tube cylinderization starts on the two embryo extremities and propagates until its middle part (cf. Fig. 10), where occurs a high concentration of myosin in bottle cells (yellow in Fig. 10). In these cells, apical constriction occurs when actomyosin contractility folds the cell membrane to reduce the apical surface area. By considering a 3D mesh representing the terminal region where curvature changes, we can simulate “in silico” the phenomenon by only taking into account the contractility in the central cells of the mesh due to a local excess of myosin diffusing from a random fluctuation in the central embryonic part.

A gastrulation model needs to account for four mechanisms, allowing realistic simulations:

1. change (due to random fluctuations) in concentration of metabolites critical for cell shape, like myosin, actin, tubulin or of the substrates (notably ATP, GTP) and enzymes ruling adenylate and guanylate pools (mainly ADPribosyl Kinase or ADK, Guanylate Kinase or GK and Nucleoside-Diphosphate Kinase or NDK) required for their polymerization. The cell shape change into a truncated pyramid (or bottle, or flask) shape is achieved in the apical portion of the cell which constricts

2. diffusion of critical metabolites provoking locally the bottle cell differentiation (in region 1 of Fig. 11)

3. cell contraction from the apical cell surface (cf. 2 of Fig. 11) and centrosome displacement in the cell depth at the cell extremities due to the elastic forces balance during the first invagination stage of the gastrulation (3 of

Fig. 8. In gray, eyes, gills and gastrula cavity anatomic frontiers in Zebra fish embryo (a) [Zanella et al., 2010], corresponding to negative curvature, maximal proliferation and minimal morphogen diffusion domains. Adult animal (b).
Fig. 9. Building gastrulation cavity with invagination first phase showing four types of cells: (1) the exothelial, (2) the bottle (BC on middle left, arch “keystone”), (3) the trapezoidal and (4) the endothelial (top). Genetic network ruling the gastrulation, with only two fixed point attractors, if genes $b$ and $c$ vanish (middle right and bottom).

Fig. 11), which is purely mechanical without proliferation (4) cell cycle arrest for bottle cells [Kurth, 2005] and after, proliferation at the end of gastrulation, consolidating the tube formation (4 of Fig. 11). For example, the onset of gastrulation in rodents is associated with the start of bottle cell differentiation within the embryo proper and after, with a dramatic increase in the rate of growth and proliferation, the cell cycle time being 7 h to 7.5 h in ectoderm and mesoderm, but 3 h to 3.5 h in the cells of the primitive
Fig. 10. Progressive invagination due to the first bottle cell differentiation in gastrulation process: experimental data (top left, from http://www.molbio1.princeton.edu/wieschaus/); tentative mechanism of propagation of the random myosin fluctuation (in blue, top right); model with myosin diffusion (in red) and cell contraction, yellow color indicating the zones of minimal diffusion (middle left) with explanation of the inward movement (middle right) and central mesh contraction showing the terminal invagination in axonometric and profile views, with calculation of forces exerted by the elastic constraints propagation on a central and distal cell, located respectively on the cylindric (A) and curved terminal (B) parts of the embryo (bottom).
Fig. 11. (a) First steps of the gastrulation with (1) diffusion of constriction metabolites from the first bottle cells (in blue), (2) occurrence of the invagination from the initial bottle cell and (3) proliferation of bottle cells, before (4) closing and enlarging the tube. (b) Curves showing the cell surface $S$ over cell volume $V$ ratio ($S/V$) depending on the mode of attachment of the myosin fibers inside the cells $A$ and $B$ of Fig. 10.
streak, whose total cell cycle time is reduced by shortening $S$ and $G_2$, as well as $G_1$ in contrast to cells later in development, where the cell cycle duration is modulated by varying the $G_1$ length [Mac Auley et al., 1993].

The gastrulation model formalizes the mechanisms causing the mechanical perturbations due to the first bottle cell differentiation (cell 2 in blue of Fig. 11); after the apical constriction of the upper cells (cf. Figs. 10 and 11) and myosin diffusion, each cell evolves with its walls following the Newton law: the sum of exerted forces is equal to the acceleration of the wall in the resultant direction (the mass of a wall being equal to 1), and each cell is submitted to forces related to internal and external pressures (created by elastic forces applied from the centrosome to the cells’ extremities in Fig. 10, bottom), plus contact forces imposed by neighbor cells. Each force is orthogonally applied to the concerned cell wall and is proportional to its length, coefficients being either the pressure or the cadherins concentration. The updating of each cell force balance is sequential: when a cell moves, it takes its neighbors with itself. These movements cause variations of cells areas: we suppose that growth occurs where the forces are high and cells are incompressible. After a radial division due to a small nutritive surface/volume ratio of the bottle cells (following the Thom’s law, described in [Forest & Demongeot, 2004, 2008; Forest et al., 2004, 2006]), we suppose that the growth in $G_1$ following the mitosis increases this ratio, ensuring a convenient nutrition. Cells are often shaped by requirements of cell surface $S$ over volume $V$ ratio ($S/V$) and namely intestinal cells have tendency to increase the area through which nutrients are absorbed [Stanek, 1983; Miller & Levine, 2002]. Ratio $S/V$ decreases when invagination occurs (cf. Fig. 9, bottom), especially if actomyosin fibers are orthogonal to microtubules [Silverman-Gavrila et al., 2008], depolymerization reducing the extent of the apical constriction [Lee & Harland, 2010]. Forces exerted on walls push centrosome and nucleus to move in the cell depth at the level of the first curved distal parts of the embryo (cf. Figs. 9 and 10) in agreement with experiments [Leptin & Grünwald, 1990], due to the location of the elastic forces application points supposed to be the same on neighbor walls, located at the cadherin and myosin-membrane-attached sites [Inoue, 1995;
The domains of minimal diffusion of myosin are shown in yellow in Fig. 10 and are located on the frontiers of the invagination (zones of zero-curvature). The link between these domains and the anatomic boundaries have to be confirmed in further 3D microscopic studies by comparing the null-curvature maps of the embryo, the zero-diffusion domains of the critical metabolites and the maximal proliferation zones. We conjecture in concluding these two short studies about feather morphogenesis and gastrulation that the zero-diffusion sets could be good candidates for ensuring locally the coexistence, and after the self-assembling of the components (carrier, receptor and attachment proteins as well as phospho-lipids) of cell interfaces between two tissues needed for separating organs functionally specified by differentiated cells. In the zero-diffusion zones indeed the effect of the temperature on the diffusion is minimum, because the viscosity (inverse of the diffusion coefficient \(D\)) is proportional to \(\exp(E/kT)\), where \(E\) is an activation energy, \(T\) the absolute temperature and \(k\) the Boltzmann’s constant. We also notice that if the concentration front wave is Gaussian, the zero-diffusion zone corresponds to the domain where the partial de Donder affinity is \(\ln(u)\), \(u\) being the concentration of the diffusing substance [Dutt, 2000], that does not vary, after reaching the reaction equilibrium, in the case where the reaction has a fast dynamics with respect to a slow diffusion:

\[
\frac{\partial \ln(u)}{\partial t} = \frac{1}{u} \frac{\partial u}{\partial t} = \frac{D \Delta u}{u}
\]

If we authorize the value of \(D \Delta u/u\) to be sufficiently small, that corresponds also to the domain where the diffusion of \(u\) is minimum and its concentration maximum. If the corresponding value of \(u\) minimizes a chemical potential from which the reaction velocity derives, like in \(\pi\)-switches involved in morphogenetic processes [Cinquin & Demyanov, 2002a; 2002b], then the zero-diffusion domains, in the case where they coincide for several constituents (e.g. of a membrane or aponeurosis), correspond to a local constancy of their concentrations favoring their interactions in order to build the self-assemble the least sensitive to the thermal fluctuations.

9. Conclusion
We have considered firstly in this paper some natural extensions of the classical Ross–McKendrick–Mac Donald approaches, in order to account for demographic and spatial dependencies of the variables involved in an infection process. One example has been presented, concerning the Black Death spread in Europe during the middle-age, which shows the interest of introducing space and biological age into the classical equations. In the future, other infectious diseases (like Sexually Transmitted Diseases) could be treated with the same approach showing the importance of the demography (the sexual relationships depending on the age of the partners) and of the socio-geography (conditioning the sexual behavior). Based on the knowledge of the new aerial routes [Khan et al., 2009], epidemics modeling will also be revisited in the near future for predicting new pandemics, with a viscosity minimal on the aerial routes. A second type of spatial dependency in a reaction–diffusion process occurs in the morphogenesis modeling: like for the epidemics, both age and spatial diffusion can explain the occurrence of spatial patterns, e.g. in the cases of feather morphogenesis and gastrulation.

In both epidemics and morphogenesis models, further studies have to be done in order to definitely emphasize and make more precise the functional role of the zero-diffusion domains, in which chemical or infectious agents coexist and interact.

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