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A Review of Matrix SIR Arino Epidemic Models

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Abstract: Many of the models used nowadays in mathematical epidemiology, in particular in COVID-19 research, belong to a certain subclass of compartmental models whose classes may be divided into three “(x, y, z)” groups, which we will call respectively “susceptible/entrance, diseased, and output” (in the classic SIR case, there is only one class of each type). Roughly, the ODE dynamics of these models contains only linear terms, with the exception of products between x and y terms. It has long been noticed that the reproduction number R has a very simple Formula in terms of the matrices which define the model, and an explicit first integral Formula is also available. These results can be traced back at least to Arino, Brauer, van den Driessche, Watmough, and Wu (2007) and to Feng (2007), respectively, and may be viewed as the “basic laws of SIR-type epidemics”. However, many papers continue to reprove them in particular instances. This motivated us to redraw attention to these basic laws and provide a self-contained reference of related formulas for (x, y, z) models. For the case of one susceptible class, we propose to use the name SIR-PH, due to a simple probabilistic interpretation as SIR models where the exponential infection time has been replaced by a PH-type distribution. Note that to each SIR-PH model, one may associate a scalar quantity Y(t) which satisfies “classic SIR relations”, which may be useful to obtain approximate control policies.

Keywords: epidemiological modeling; COVID-19; SIR-PH model; matrix SIR model; reproduction number; first integral

1. Introduction

Motivation. Mathematical epidemiology may be said to have started with the SIR ODE model, which saw its birth in the works of Kermack–McKendrick [1]. This was initially applied to model the Bombay plague of 1905–06, and later to measles [2], smallpox, chickenpox, mumps, typhoid fever and diphtheria, and recently to the COVID-19 pandemic—see, for example in [3–18], to cite just a few representatives of a huge literature.

Note that during the COVID-19 pandemic, researchers have relied mostly on models with quadratic interactions (linear force of infection), which belong furthermore to a parabolic class [19–22] of “(x, y, z)” models. Here, x denotes “entrance/susceptible” classes; y denotes diseased classes, which must converge asymptotically to 0; and z denotes output classes. These models are very useful; to make references to them easier, we propose to call them matrix-SIR (SYR) models, and also SIR-PH [21], when x ∈ R1.

Contents. We begin by recalling in Section 2 several basic explicit formulas for the SIR model. Section 3 presents the corresponding SIR-PH generalizations, and Section 4 offers some applications: the SEIHRD model [23–27], which adds to the classic SEIR (susceptible + exposed + infectious + recovered) a hospitalized (H) class and a dead class (D); the SEICHRD model [28] which adds a critically ill class (C); the SEIARD [29] and SEIAHR/SEIRAH(D) models [30–36], which add an asymptomatic class (A); and the SJQR model [37]. This is just a sample chosen from some of our favorite COVID papers. We note in passing that they seem though all unaware of the existence of the Arino and Feng Formulas (12) and (17).
Finally, Section 5 reviews briefly the case of several classes of susceptibles. This topic requires further development; we include it however due to the recognized importance of heterogeneity factors.

2. The Classic Kermack–McKendrick SIR Epidemic Model

The SIR process \( S(t), I(t), R(t), t \geq 0 \) divides a population of size \( N \) undergoing an epidemic into three classes called “susceptibles, infectives, and removed”. One may also work with the corresponding fractions \( s(t) = \frac{S(t)}{N}, i(t) = \frac{I(t)}{N}, \) and \( r(t) = 1 - s(t) - i(t) \). It is assumed that only susceptible individuals can get infected. After having been infectious for some time, an individual recovers and may not become susceptible again. “Viewed from far away”, this yields the SIR model with demography [1,38]

\[
\begin{align*}
S'(t) &= -\frac{\beta}{N} S(t) I(t) + \xi (N - S(t)), \\
I'(t) &= I(t) \left( \frac{\beta}{N} S(t) - \gamma - \xi \right), \\
R'(t) &= \gamma I(t) - \xi R(t),
\end{align*}
\]

(1)

where

1. \( N \) is the total, constant population size.
2. \( R'(t) \), the number of removed per unit time, is the only quantity which is clearly observable, at least in the easy case when the removed are dead, as was the case of the original study of the Bombay plague [1].
3. \( \xi \) is the population death rate, assumed to equal the birth rate.
4. \( \gamma \) is the removal rate of the infectious, which equals \( 1/duration \) of the infection (under the stochastic model of exponential infection durations, this is the reciprocal of the expected duration).
5. \( \beta \), the infection rate, models the probability that a contact takes place between an infected and a susceptible, and that it results in infection.

Note that

1. The sum \( S + I + R = N \) is conserved and each value is positive, so the values of \( S, I, R \) remain in the interval \([0, N]\).
2. This system has a unique solution, as (given the boundedness of \( S, I, R \)), the RHS above is Lipschitz.
3. From now on, we will assume that \( \xi = 0 \), and work with the fractions \( s, i, r \), which satisfy

\[
\begin{align*}
s'(t) &= -\beta s(t) i(t), \\
i'(t) &= i(t) [\beta s(t) - \gamma], \\
r'(t) &= \gamma i(t).
\end{align*}
\]

(2)

We will call this the classic SIR model, see Figure 1. Note that

1. \( s(t) \) is monotonically decreasing and \( r(t) \) is monotonically increasing, to, say, \( s_{\infty}, r_{\infty} \); therefore convergence to some fixed stable point \( (s_{\infty}, i_{\infty}, r_{\infty}) \) must hold.
2. the equilibrium set of stable points is \( (s, 0, 1 - s), s \in [0,1] \).
3. solutions starting in the domain

\[
D := \{(s, i, r) : s > 0, i > 0, r \geq 0, s + i + r \leq 1\}
\]
cannot leave it.
4. The second equation of (1) implies the so-called threshold phenomenon: if

\[
\mathcal{R} := \frac{\beta}{\gamma} \leq 1
\]

(3)
then \( i(t) \) decreases always, without any intervention. \( R \) is called reproduction number, and it models the number of susceptibles infected by one infectious (expected number, under more sophisticated stochastic, branching models). A convenient analytical definition, as the Perron–Frobenius eigenvalue of the “next-generation matrix” [39–41], is available as well.

To avoid trivialities, we will assume \( R > 1 \) from now on.

5. When \( R > 1 \), the epidemic grows iff \( s > 1/R \), i.e., until the susceptibles \( s(t) \) reach the immunity threshold

\[
\Theta := \frac{1}{R} = \frac{\gamma}{\beta},
\]

after which infections decline.

An advantage of the classic SIR model is that it is essentially solvable explicitly:

1. We can eliminate \( r \) from the system using the invariant \( s + i + r = 1 \).
2. It can easily be verified that

\[
\mu(s, i) := s + i - \frac{1}{R} \ln(s)
\]

is invariant, (note that we have used different fonts for \((s, i, r)\) when they are functions of \( t \), and standard fonts when they are not), so that \( i \) is explicitly given by

\[
i_R(s) = -s + \frac{1}{R} \ln(s) + \mu_R(s_0, i_0),
\]

and the full system (2) can be reduced to the single ODE

\[
s'(t) = -\beta s(s_0 + i_0 - s) - \gamma s \ln \left( \frac{s}{s_0} \right).
\]

3. The maximal value of the infected \( i \), achieved when \( s = 1/R \), is

\[
i_{\text{max}} = i_{\text{max},R}(s_0, i_0) = i_0 + s_0 - \frac{1 + \ln(s_0 R)}{R} = i_0 + \frac{H(s_0 R)}{R}, \quad H(R) = R - 1 - \ln(R).
\]

4. By differentiating the right-hand side of (7), one finds that the maximal value of the “newly infected” \( s' = \beta i \) \( s \) is achieved when

\[
s = i + \frac{\gamma}{\beta}, \quad s = -\frac{1}{2R} L_{-1} \left[ -2R s_0 e^{-1 - R(s_0 + i_0)} \right],
\]

where the Lambert function \( L_{-1} \) is a real inverse of \( L(z) = ze^z \)– see for example [42–44]. Bounding \( s \) \( i \) is one interesting possibility for accommodating ICU constraints [45] (2.20).

5. The infectious class converges to 0 and the susceptible and recovered converge monotonically to limits which may be expressed in terms of the “Lambert–W(right)” function \( L_0 \).

Let us note that accurate numerical solutions of the evolution of the SIR or other compartmental epidemic may be obtained very quickly.
3. SIR-PH Epidemics with One Susceptible Class (SIR Epidemics with Phase-Type “Disease Time”)

It has been known for a long while that \( R \) and the final size for many compartmental model epidemics may be explicitly expressed in terms of the matrices which define the model, and the works in [19,20,46,47] offer a quite general framework of “xyz” models which ensures this. We believe that these formulas have not received the attention they deserve (they keep being reproved), and decided therefore to review them below; we will call them matrix- SIR models.

A particular but revealing case is that when there is only one susceptible class, which we will call SIR-PH, following Riano [21], who emphasized its probabilistic interpretation—see also in [48].

Definition 1. A “SIR-PH \((\vec{\alpha}, V, \beta, W)\) epidemic” contains a homogeneous susceptible class, but vector “diseased” state \(\vec{i} (t)\) (which may contain latent/exposed, infective, asymptomatic, etc.) and vector removed states (healthy, dead, vaccinated, etc.). It is defined by an ODE system:

\[
\begin{align*}
    s'(t) &= -s(t) \vec{i}(t) \beta \\
    \vec{i}'(t) &= s(t) \vec{i}(t) \beta \vec{\alpha} + \vec{i}(t) A \\
    \vec{r}'(t) &= \vec{i}(t) W
\end{align*}
\]

where

1. \( \vec{i}(t) \in \mathbb{R}^n \) is a row vector whose components are fractions of diseased individuals of various types, which must satisfy \( \vec{i}(\infty) = 0 \).
2. \( \beta \in \mathbb{R}^n \) is a column vector whose components represent the relative transmission ability of the various disease classes.
3. \( \vec{\alpha} \in \mathbb{R}^n \) is a probability row vector with the components representing the fractions of susceptibles entering into the corresponding disease compartments, when infection occurs.
4. \( A \) is a \( n \times n \) Markovian sub-generator matrix describing rates of transition between the diseased classes \( \vec{i} \) (i.e., a Markovian generator matrix for which the sum of at least one row is strictly negative). Alternatively, \( V := -A \) is a non-singular \( M \)-matrix ( \( M \)-matrix is a real matrix \( V \) with \( v_{ij} \leq 0, \forall i \neq j \), and having eigenvalues whose real parts are nonnegative [49]).
5. \( \vec{r}(t) \in \mathbb{R}^p \) is a row vector which must satisfy \( \vec{r}(\infty) > 0 \), whose components represent (fractions of) various classes which survive at the end of an infection.
6. \( W \) is a \( n \times p \), matrix whose components represent the rates at which classes of diseased individuals become recovered. We assume that the \( n \times (n + p) \) matrix \( \tilde{V} = (A \ W) \) has row sums 0 (which implies that the total population \( N(t) = s(t) + \vec{i}(t)1 + \vec{r}(t)1 \) is constant).
We turn now to a deceivingly simple particular example of the SIR-PH model, which explains its name.

**Remark 1. Probabilistic interpretation of SIR-PH epidemics.** For simplicity, let us group all the output classes of (10) into one \( r = \vec{r}_1 \), where \( \vec{1} \) is a column vector of ones of appropriate dimension (here \( p \)), yielding

\[
\begin{align*}
& s'(t) = -s(t)\vec{i}(t)\beta \\
& \vec{i}'(t) = s(t)\vec{i}(t)\beta \vec{\alpha} + \vec{i}(t)A \\
& r'(t) = \vec{i}(t)a,
\end{align*}
\]

where we put \( a := W\vec{1} = (-A)\vec{1}. \)

(11) emphasizes the fact that SIR-PH models are in one to one correspondence with laws of phase-type \((\vec{\alpha}, A)\) [21] (21).

Let us recall now, as known essentially since the work in [50]—see also [51] (Theorem 2.2.7)—that under proper scaling, the expected fractions \( s(t), \vec{i}(t), r(t) \) of stochastic SIR (One such model stipulates that each infective \( j = 1, \ldots, J \) infects a randomly chosen susceptible, at encounter times which belong to independent Poisson processes \( P_j(t), j = 1, \ldots, J \), of rate \( \beta_j \), and that infection durations are i.i.d. r.v.'s which are exponentially distributed at the end of which the individual recovers (or dies.) and more general compartmental models obey a “law of large numbers/fluid limit” which recovers the deterministic epidemic.

As an example, the SIR-PH model (10) may be derived as limit of a stochastic SIR model in which the exponential infection time has been replaced by a phase-type \((\vec{\alpha}, A)\) “dwell period” [48].

**Proposition 1.** For processes defined by (10), with \( V = -A \) a non-singular M-matrix, the basic reproduction number is given by [19] (Theorem 2.1) (This can be also derived as the expected number of susceptibles infected during a dwell period, for the stochastic model (the so-called “survival method”)—see the work in [52] for an excellent review).

\[
R = \vec{\alpha} V^{-1} \beta. \tag{12}
\]

A disease-free equilibrium \((s_0, \vec{0}, \vec{r}_0)\) is asymptotically stable iff \( s_0 < \frac{1}{R} \).

To illustrate the power of the SIR-PH formalism, consider now the case with two diseased states—latent and infectious—with phase-type dwell times, parametrized by \((\vec{\alpha}_e, A_e)\) and \((\vec{\alpha}_i, A_i)\), respectively. Using the well-known convolution formula—see, for example, [53] (Theorem 3.1.26) we find that formulas like (12) (see other examples of such formulas below) still apply, with \((\vec{\alpha}, A, \beta)\) given by

\[
\vec{\alpha} = (\vec{\alpha}_e, 0), A = \begin{pmatrix} A_e & a_e \vec{\alpha}_i \\ 0 & A_i \end{pmatrix}, \beta = \begin{pmatrix} 0 \\ 0 \\ \vdots \\ \beta_{1,1} \\ \beta_{1,2} \\ \vdots \end{pmatrix}. \tag{13}
\]

The “epidemic dwell structure” \((\vec{\alpha}, A, \beta)\) of examples with more complicated network structures for the diseased may be constructed using Kronecker sums of the matrices defining each component.

Let us give now an example which does not in general belong to the SIR-PH class.
Example 1. The SIRV model—see, for example, in [54]—is defined by
\[
\begin{align*}
    s'(t) &= -s(t)(\beta i(t) + \gamma), \\
    i'(t) &= i(t)(\beta s(t) - \gamma), \\
    r'(t) &= \gamma_i i(t), \\
    \varphi'(t) &= \gamma s(t).
\end{align*}
\]
This is of the form (10) with \( \gamma = (i) \), \( \tau = (r, v) \) iff \( \gamma_s = 0 \).
In the opposite case \( \gamma_s \neq 0 \), one may still compute an invariant (Note that \( \gamma_s > 0 \) has been used for modeling vaccination, and \( \gamma_s < 0 \) has been used for modeling loss of immunity)
\[
\mu(s, i) := \beta(s + i) - \gamma \ln(s) + v \ln(i),
\]
and for fixed \( s \), putting \( \tilde{\gamma} = \frac{\gamma}{\tilde{s}} \), \( \tilde{\beta} = \frac{\beta}{\tilde{s}} \), \( \tilde{\mu} = \frac{\mu_0}{\tilde{s}} \), it holds that \( i \) is explicitly given by
\[
i(s) = \frac{1}{\beta}L_0 \left[ \tilde{\beta} \tilde{s}^{\tilde{\gamma}} e^{\tilde{\mu}_0 - \tilde{\beta} s} \right].
\]
When \( \gamma_s > 0 \), the final size is \( s_\infty = 0 \).

We provide now a list of several formulas, obtained by replacing \( i \) in SIR by a scalar linear combination (17) [47]. They are all easily proved; however the formula for the maximal value of the newly infected involves also a second linear combination \( y \) (21).

Proposition 2. For processes defined by (10), with \( V = -A \) a non-singular M-matrix, it holds that:
1. The following weighted sum of the diseased variables [47] (24)
\[
Y(t) = \frac{\tilde{i}(t)}{\tilde{\alpha}} V^{-1} \tilde{\beta} = \frac{1}{\mathcal{R}} \tilde{i}(t) V^{-1} \tilde{\beta}
\]
has the property that
\[
\frac{dY}{ds} = \frac{1}{\mathcal{R}} \tilde{i}(t) \left( s(t) \tilde{\beta} \tilde{\alpha} - V \right) V^{-1} \tilde{\beta} = \frac{\tilde{i}(t) \beta \left( s(t) \mathcal{R} - 1 \right)}{-\mathcal{R} s(t) \tilde{i}(t) \tilde{\beta}} = -1 + \frac{1}{\mathcal{R} s},
\]
and that
\[
\begin{align*}
    z(t) := \mu( s(t), Y(t)) := Y(t) + s(t) - \frac{1}{\mathcal{R}} \ln[s(t)], \\
    Z(t) := e^{-\mathcal{R} z(t)} = s(t) e^{-\mathcal{R}(s(t) + Y(t))}
\end{align*}
\]
are constant along the paths of the dynamical system (10).
The solution of \( Z(s) = Z(0) \) with respect to \( s \) may be expressed in terms of
\[
s(t) = -\frac{1}{\mathcal{R}} L_0 \left[ -\mathcal{R} Z_0 e^{\mathcal{R} Y(t)} \right],
\]
where \( [-e^{-1}, \infty) \ni z \rightarrow L_0(z) \in [-1, \infty) \) is the principal branch of the Lambert-W function.
2. The derivative with respect to time is
\[
\frac{dY}{dt} = \left( s - \frac{1}{\mathcal{R}} \right) \tilde{i} \beta := \left( s - \frac{1}{\mathcal{R}} \right) y.
\]
Therefore, \( \frac{dY}{ds} = 0 = \frac{dY}{dt} \) iff \( s = \mathcal{R}^{-1} \).
3. The maximum value of $Y$ occurs for $s = \min[1/R, 1]$. In the case $R > 1$, this yields [47] (Section 2.1):

$$Y_0 + s_0 - Y^* - \frac{1}{R} + \frac{1}{R} \ln(s_0 R),$$

(22)

by the conservation of $Y(t) + s(t) - \frac{1}{R} \ln(s(t))$ between the time 0 and the time $t_1/R$ of reaching the immunity threshold.

4. The final size of the susceptibles satisfies [19] (Theorem 5.1):

$$\ln(s_0/\bar{s}_\infty) = R(s_0 - s_\infty) + \int_0^1 V^{-1} \beta = R(s_0 - s_\infty + Y_0),$$

(23)

by the conservation of $Y(t) + s(t) - \frac{1}{R} \ln(s(t))$ between the times 0 and $\infty$; explicitly,

$$s_\infty = -\frac{1}{R} L_0[-R Z_0] = -\frac{1}{R} L_0[-R s_0 e^{-R(s_0 + Y_0)}].$$

(24)

5. The integrated infectives $\tilde{I}^{(a,b)} = \int_a^b \tilde{i}(s) ds$ satisfies

$$\begin{cases}
\tilde{I}^{(a,b)} V = \tilde{I}_a - \tilde{I}_b, \\
(\tilde{I}_a - \tilde{I}_b) V^{-1} \beta = \log \left( \frac{S_a}{S_b} \right),
\end{cases}$$

(25)

and the total integrated infectives $\tilde{I}^{(\infty)} = \int_0^\infty \tilde{i}(s) ds$ satisfies [19] (6)

$$\tilde{I}^{(\infty)} V = \tilde{i}_0 + (s_0 - s_\infty) \tilde{x}.$$ 

(26)

6. The final size of the removed satisfies:

$$\bar{r}_\infty - \bar{r}_0 = I^{(\infty)} W = \left( \tilde{i}_0 + (s_0 - s_\infty) \tilde{x} \right) V^{-1} W,$$

(27)

7. The value of the infected combination $Y$ when $s = 1/R$ is

$$Y_{max} = i_{max,R}(s_0, \tilde{i}_0) = Y_0 + s_0 - \frac{1 + \ln(s_0 R)}{R} = i_0 + \frac{H(s_0 R)}{R}, H(R) = R - 1 - \ln(R).$$

(28)

8. The maximum size of the newly infected is achieved when

$$s(t) = \frac{Y^2 + \beta Y(t)}{\tilde{x} \beta y}.$$ 

(29)

Remark 2. In particular, for the SIR model (2),

$$\log \left( \frac{s(a)}{s(b)} \right) = \beta I^{(a,b)} = R(J_a - J_b), J = s + i.$$ 

(30)

Note that this has been used to model the total cost of an epidemic [55].

Remark 3. Let us note that for control problems involving optimization objectives which only depend on $Y(t)$, we are effectively optimizing a SIR model; this SIR approximation may be used to offer practical solutions for optimizing more complicated compartmental models.

Example 2. For SEIR, putting $\tilde{i} = (e, i)$, we may write

$$\begin{cases}
\tilde{s}' = -\beta s i \\
\tilde{i}' = (\beta s i - \gamma_e e, \gamma_e e - \gamma i) = \beta s i (1, 0) - (e, i) \begin{pmatrix} \gamma_e & -\gamma e \\ 0 & \gamma \end{pmatrix} \\
r' = \gamma i
\end{cases}$$
so that
\[
\begin{aligned}
\bar{a} &= (1, 0), \\
b &= \begin{pmatrix} 0 \\ \beta \end{pmatrix}, \\
V &= \begin{pmatrix} \gamma & -\gamma e \\ 0 & \gamma \end{pmatrix} \quad \Rightarrow \quad V^{-1} = \frac{1}{\gamma e} \begin{pmatrix} \gamma & \gamma e \\ 0 & \gamma \end{pmatrix}, \\
Y &= \begin{pmatrix} e_i, i \\ 0, \gamma e \end{pmatrix}.
\end{aligned}
\]

4. Examples of SIR-PH Models Used in COVID-19 Modelling

We derive now $R$ and $Y$ from (12), (17), for some popular compartmental models. Note that we will be reformulating the original results (which, unfortunately, have already appeared several times with different notations), using a unifying notation.

Example 3. The SEIHRD model \cite{23–27} has disease states $\vec{i} = (e, i, h)$. We use here the version in \cite{25} (we would rather call this SI$^2$HRD model) defined by,

\[
\begin{aligned}
\bar{a} &= (1, 0, 0), \\
\beta &= \begin{pmatrix} \beta_e \\ \beta_i \\ 0 \end{pmatrix}, \\
V &= \begin{pmatrix} \gamma_e & -\gamma_e \\ 0 & \gamma_e \end{pmatrix} \\
W &= \begin{pmatrix} e_r & 0 \\ i_r & 0 \\ h_r & 0 \\ h_d & 0 \end{pmatrix},
\end{aligned}
\]

where we denoted by $\gamma_e, \gamma_i, \gamma_h$ the sum of the constant rates out of $e, i, h$, and by $i_h$ the rate out of $i$ and reaching $h$, etc. See Figure 2.

Then, $R = \frac{\beta_e}{\gamma_e} + \frac{i_h}{\gamma_i} \gamma_i \gamma_h \beta_i \beta'_i$ \cite{25} (2), and $Y = e + i \gamma_e \beta_i \beta'_i$. When $\beta_e = 0 = e_r \Rightarrow \frac{\beta_e}{\gamma_e} = 1$, we recover $R = \frac{\beta_i}{\gamma_i}$ \cite{24} and $Y = e + i$.

![Figure 2](image)

Figure 2. Flowchart of the SEIHRD model. The forces of infection are $F_e = \beta_e e$, $F_i = \beta_i i$, $F = F_e + F_i$. The red edge corresponds to the entrance of susceptibles into the diseased classes; $\bar{a}$, the dashed green edges correspond to contacts between diseased to susceptibles, the brown edges are the rates of the transition matrix $V$, and the remaining yellow dashed flows correspond to the rates of $W$.

Example 4. The SEIHCRD model of the work in \cite{28}, has disease states $\vec{i} = (e, i, h, c)$ and is defined by

\[
\begin{aligned}
\beta &= \begin{pmatrix} \beta_i \\ \beta_e \\ 0 \\ 0 \end{pmatrix}, \\
\bar{a} &= (1, 0, 0, 0), \\
V &= \begin{pmatrix} \gamma_e & -\gamma_i & 0 & 0 \\ \gamma_i & -i_h & 0 \\ 0 & 0 & \gamma_h & -h_c \\ 0 & -c_h & \gamma_c & \gamma_e \end{pmatrix}, \\
W &= \begin{pmatrix} 0 & 0 \\ 0, i_r \\ h_r & 0 \\ 0 & 0 & e_d \end{pmatrix},
\end{aligned}
\]
see Figure 3. Then,

\[ \mathcal{R} = \frac{\beta_i}{\gamma_i}, Y = e + i. \]

Then,

\[ R = \beta_i \gamma_i, Y = e + i. \]

\[ \mathcal{R} = \frac{\beta_i}{\gamma_i}, Y = e + i. \]

Figure 3. Flowchart of SEIHCRD model. The force of infection is \( F = \beta_i i \).

**Example 5.** The SEIRAH(SEIAHR) model \([30,36]\) has disease states \( \vec{i} = (e, i, a, h) \) and is defined by

\[
\beta = \begin{pmatrix} 0 \\ \beta_i \\ \beta_a \\ 0 \end{pmatrix}, \bar{\alpha} = (1, 0, 0, 0), V = \begin{pmatrix} \gamma_e & -e_i & -e_a & 0 \\ 0 & \gamma_i & 0 & -i_h \\ 0 & 0 & \gamma_a & -a_h \\ 0 & 0 & 0 & \gamma_h \end{pmatrix}, W = \begin{pmatrix} e_r \\ i_r \\ a_r \\ \gamma_h \end{pmatrix}.
\]

See Figure 4 below. Then,

\[ \mathcal{R} = \frac{\epsilon_a}{\gamma_e} \mathcal{R}_a + \frac{\epsilon_i}{\gamma_e} \mathcal{R}_i, \quad \mathcal{R}_i = \frac{\beta_i}{\gamma_i} \mathcal{R}_a = \frac{\beta_a}{\gamma_a}, \]

and

\[ Y = e + \frac{\mathcal{R}_i}{\mathcal{R}} + \frac{\mathcal{R}_a}{\mathcal{R}}. \]

Figure 4. Flowchart of the SEIAHR model. The forces of infection are \( F_{si} = \beta_i i, F_{sa} = \beta_a a, F = F_{si} + F_{sa}. \)
Example 6. The S\textsuperscript{aps}QR model (with asymptomatic, pre-symptomatic, and symptomatic infectious) [37,56] (3.2). See Figure 5;

![Flowchart of S\textsuperscript{aps}QR model](image)

Figure 5. Flowchart of S\textsuperscript{aps}QR model. The forces of infection (rates) are \( F_{sa} = \beta_a i_a \), \( F_{sp} = \beta_p i_p \), \( F_{ss} = \beta_s i_s \), \( F = F_{sa} + F_{sp} + F_{ss} \).

The disease states are \( \mathbf{i} = (i_a, i_p, i_s, q) \), and the model is defined by

\[
\beta = \begin{pmatrix} \beta_a \\ \beta_p \\ \beta_s \\ 0 \end{pmatrix}, \quad \mathbf{\alpha} = (\phi, 1 - \phi, 0, 0), \quad V = \begin{pmatrix} \gamma_a & 0 & 0 & 0 \\ 0 & \gamma_p & -\gamma_p & 0 \\ 0 & 0 & \gamma_s & -\gamma_q \\ 0 & 0 & 0 & \gamma_q \end{pmatrix}, \quad W = \begin{pmatrix} \gamma_a \\ 0 \\ i_{sr} \\ \gamma_q \end{pmatrix}.
\]

Then, from the work in [37],

\[
R = \phi \frac{\beta_a}{\gamma_a} + (1 - \phi) \left( \frac{\beta_p}{\gamma_p} + \frac{\beta_s}{\gamma_s} \right),
\]

and,

\[
Y = \frac{1}{R} \left[ i_a R_a + i_p (R_p + R_s) + i_s R_s \right],
\]

where \( R_a := \frac{\beta_a}{\gamma_a} \), \( R_p := \frac{\beta_p}{\gamma_p} \), and \( R_s := \frac{\beta_s}{\gamma_s} \).

5. S\textsuperscript{(m)}YR Models with \( m \) Groups of Susceptibles

The SIR-compartment model makes the unrealistic assumption that the population through which the disease is spreading is well-mixed. However, differences in susceptibility and rates of contact between individuals strongly affect their likelihood of catching COVID-19. A model which attempts to capture this aspect is:

\[
\begin{align*}
    s_k'(t) &= -s_k(t) i_k(t) \beta_k, \quad k = 1, \ldots, m \\
    i_k'(t) &= \sum_{k=1}^{m} s_k(t) i_k(t) \beta_k \mathbf{\alpha} - i_k(t) V \\
    r'(t) &= i(t) W
\end{align*}
\]

Lemma 1. A disease free equilibrium \((s_1, s_2, \ldots, s_m, 0, 0)\) of (31) is asymptotically stable iff \( sR < 1 \), where \( s = \sum_k s_k \) and

\[
\mathcal{R} = \sum_k \frac{s_k}{s} \bar{\alpha} V^{-1} \beta_k = \sum_k \frac{s_k}{s} \mathcal{R}_k, \quad \mathcal{R}_k := \bar{\alpha} V^{-1} \beta_k
\]

is the spectral radius of the next generation matrix.
While the final size may also be obtained under this model [20] (Theorem 2.1), for transient behavior it is convenient to turn to a simpler model.

A Generalization of Heterogeneous SEIR

Assume now that \( \beta_k = \beta_1 \beta_i \) where \( \beta_i, \beta_1 \in \mathbb{R}^+ \) and \( W = \beta \bar{w} \), where \( \bar{w} \) is a row vector. Putting \( \gamma = \gamma \beta \), the system \((31)\) becomes (Such dynamics was first considered in [57]).

\[
\frac{d \log s_k}{dt} = -\beta_k y, \quad \frac{d i}{dt} = \left( \sum_{k=1}^{m} \beta_k s_k \right) y \bar{a} - \gamma V, \quad \frac{d r}{dt} = \gamma W = y \bar{w},
\]

and

\[
\bar{r}(t) = \int_0^t y(u) du \, \bar{w} := I(t) \, \bar{w}.
\]

It is convenient to reparametrize the model taking \( I \) as parameter, or, equivalently, by taking

\[
r = \bar{r} \mathbf{1}, \gamma = \bar{w} \mathbf{1}.
\]

Solving

\[
\frac{d}{dt} (\log s_k) = -\beta_k y = -\beta_k \frac{d}{dt} r,
\]

we find that the system has a family of first integrals which includes

\[
\begin{align*}
\frac{1}{m} \log(s_1/s_1(0)) &= \cdots = \frac{1}{m} \log(s_k/s_k(0)) = \cdots = \frac{1}{m} \log(s_m/s_m(0)) = r(0) - r, \\
\sum_{k=1}^{m} s_k + \sum_{k=1}^{n} y_k + \sum_{k=1}^{n} r_k &= 1
\end{align*}
\]

Also

\[
s_k(t) = s_k(0) e^{-\beta_k I(t)} = s_k(0) e^{-\frac{\gamma}{\gamma} r(t)},
\]

where \( \gamma \) is defined in \((35)\).

We conclude with some preliminary results on this model.

**Lemma 2.** (a) The authors of [58] put \( s(t) = \sum_k s_k(t), p_k = s_k(0)/s(0) \). The solution of \((33)\) satisfies the time-dependent SYR system:

\[
\frac{d s}{dt} = -a(t) s(t) y(t), \quad \frac{d i}{dt} = \left( \sum_k \beta_k s_k(t) \right) y(t) \bar{a} - \gamma(t) V, \quad \frac{d r}{dt} = \gamma y(t),
\]

where

\[
a(t) = \frac{\sum_k \beta_k s_k(t)}{s(t)}
\]

is a positive non-increasing function with \( a(0) = \sum_k p_k \beta_k := \bar{\beta} \).

(b) \( Y(t) \) defined in \((17)\) with \( R = \sum_k p_k R_k, R_k = \beta_k V^{-1} \beta \), satisfies

\[
\frac{1}{y(t)} \frac{d Y}{dt} = \frac{1}{R} \left( \sum_k s_k(t) R_k - 1 \right) = \frac{R e(t) - 1}{R} = \frac{\gamma}{\gamma} \frac{d Y}{d r}
\]

and is unimodal, with a maximum on the immunity/recovery line

\[
\sum_k s_k R_k = 1.
\]

(c) The stable stationary solution \( (s_1^*, s_2^*, \ldots, 0, 0, \ldots, \bar{r}^*) \) is determined by the unique solution with \( r^* = r > r(0) \) of

\[
1 = \sum_{k=1}^{m} s_k(0) e^{-\frac{\beta_k}{\gamma} r}, \quad s_k^* = s_k(0) e^{-\frac{\beta_k}{\gamma} r}, \quad k = 1, 2, \ldots, m.
\]
Proof. (a) The only thing which requires a proof is that \( a(t) \) is a positive non-increasing function. However, the derivative of \( a \) satisfies
\[
\frac{da}{dt} = -\frac{\gamma(t)\bar{\beta}}{s(t)^2} \left( \sum_k \beta_k(t) \sum_k \beta_k^2 s_k(t) - \left( \sum_k \beta_k(t) s_k(t) \right)^2 \right) \leq 0,
\]
by the Cauchy–Schwarz inequality.

(b) \( Y(t) = \frac{1}{\pi} \bar{\gamma}(t) V^{-1} \beta \implies Y'(t) = \frac{1}{\pi} \bar{\gamma}'(t) V^{-1} \beta = \frac{1}{\pi} \left[ \left( \sum_k \beta_k(t) \beta_k \right) Y(t) \bar{\alpha} - \bar{\gamma}(t) \bar{V} \right] V^{-1} \beta \)
\[
= \frac{\bar{\gamma}(t)}{\pi} \left( \sum_k \beta_k(t) \bar{R}_k - 1 \right)
\]
(c) This follows from the conservation of mass and \( \bar{\gamma}(\infty) = 0 \) [58]. □

6. Conclusions
We believe that the matrix SIR Arino epidemic models could provide a guiding thread to a great part of the epidemics literature. We are currently investigating this family of epidemic models in the presence of extra births and deaths.

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