

1 *Novel Challenges and Opportunities in the Theory and Practice of*
2 *Matrix Population Modelling*

3
4 An editorial for the Special Feature

5 **“Theory and Practice in Matrix Population Modelling”**
6 *of Ecological Modelling*

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20Abstract

21Demography is at the core of ecology, evolution, and conservation biology. The simple
22recognition that individuals in a given population contribute to its dynamics in different ways
23revolutionised the ways in which demographers approach data collection, analyses, and
24interpretation of their study populations, from bacteria to humans. Matrix population models,
25discrete-time, discrete-state (*i.e.* individuals are categorised into discrete categories based on traits
26such as age or stage), were first introduced to the scientific community by Patrick Leslie 75 years
27ago. Since then, the applications of matrix population models to ecology, evolution, and
28conservation biology have strongly been running strong and in parallel with its robust
29mathematical development. This special feature contains 14 novel contributions that represent
30some the cutting-edge mathematical formulations and applications of this powerful demographic
31tool. In addition to highlighting the key contributions of this manuscripts, we provide suggestions
32to some of the challenges that researchers using matrix population models must overcome in the
33coming decades to truly unlock the potential of this analytical demographic tool.

34

35Keywords

36Comparative biology; Data gaps; Matrix averaging; Population ecology; Potential-growth
37indicators; Population viability; Stochastic growth rate.

381. Introduction

39 The 1945 publication ‘On the use of matrices in certain population mathematics’ by Patrick
40 H. Leslie has revolutionised mathematical demography and population ecology. This manuscript,
41 together with many others produced throughout Leslie’s prolific career (1948; Caswell, 2001, and
42 refs therein), marked the inception of the vastly rich world of *Matrix Population Models* (MPMs,
43 hereafter). Now, at the 75th anniversary of the publication of this seminal contribution, MPMs
44 have moved from being an exclusively useful exercise ‘in certain population mathematics’ to a
45 powerful, frequently used tool across numerous disciplines, including ecology (e.g., Crone et al.,
46 2011), evolution (e.g., van Tienderen, 1995), and conservation biology (e.g., Morris and Doak,
47 2002). MPMs have been applied to a wide range of taxonomic groups, from humans (Nicol-
48 Harper, 2018), to whales (Chiquet et al. 2013), penguins (Jenouvrier et al., 2012), plants
49 (Salguero-Gómez et al., 2016), bacteria (Watwe et al., 2006) and even viruses (Wallace, 2004).
50 Indeed, MPMs have become as equally popular among field ecologists as they continue to be
51 among theoretical ecologists. Even now, Leslie’s pioneering work motivates new developments in
52 the mathematical theory of nonnegative matrices (Li and Schneider, 2002; Tuljapurkar and
53 Haridas, 2006; Protasov and Logofet, 2014; Logofet, 2018, 2019a; Razzhevaikin and
54 Tyrtshnikov, 2020), a single mathematical basis of diverse applications.

55 MPMs represent an elegant, relatively straightforward, powerful tool to describe the dynamics
56 of a population in discrete time and discrete structure (Caswell, 2001). The population structure,
57 whereby individuals are classified following a relevant set of criteria that separates them into
58 discrete states (e.g., ages, size ranges, development, nationality, etc.), together with information
59 regarding the species’ life cycle, allows researchers to draw direct links between transitions and
60 per-capita contributions of different stages along the lifecycle (*i.e.*, the *Life Cycle Graph, LCG*), as
61 well as the likelihood/intensity of those demographic events (*i.e.*, *vital rates*).

62 When calibrated with field or laboratory data, the *Population Projection Matrix* (PPM), the
63 core of any MPM, serves as an indicator of the environment quality and biological properties of
64 the species’ population under study (Caswell, 2001). This PPM provides a rich repertoire of
65 quantitative characteristics that allow not only for the careful characterisation of the current and
66 future state of that population (e.g., Jenouvrier et al., 2009), but also for comparative (e.g., Jones
67 et al., 2013), empirical (e.g. Crone et al., 2011) and theoretical research (e.g., Haridas and
68 Tuljapurkar, 2007).

69 Depending on the data source and organism under study, MPMs face interesting analytical
70 challenges, but also provide unique research and methodological opportunities. The Special Issue
71 “*Theory and Practice in Matrix Population Modelling*” in *Ecological Modelling* aims to reflect
72 the current state-of-the-art in the *Theory and Practice of Matrix Population Modelling*. Together,

73the special feature includes 14 contributions (marked hereafter with ^{SI}) that address topics at the
74frontiers of MPM's development and applications. Our goal here is to outline some of the next
75frontiers in the theory and praxis of the MPM world, while highlighting how each of these
76publications makes a clear contribution to these areas.

77

782.1. Exploring taxonomic/ecoregion biases

79 Although the usage of MPM across a wide range of species and regions has become a
80reality, important biases remain in the demographic literature (Conde et al., 2019). These biases
81are preventing researchers from examining the validity of general rules (see next Section). For
82instance, even though most of the terrestrial biodiversity worldwide is found in tropical ecoregions
83(Tomašových, 2019; Wang et al., 2019; Loiseau et al., 2020), most of the studies that have used
84MPMs on terrestrial animals and plants have taken place in grasslands and temperate ecoregions
85(Salguero-Gómez et al., 2015a, b). From this angle, the following manuscripts in this special
86feature make contributions to expanding our demographic knowledge across the Tree of Life and
87the planet.

88 Pinto et al. (2020^{SI}) make an important contribution to our understanding of the demography
89of insects. Insects are the most specious animal groups, with an estimated >1 million species
90(Zhang, 2013), yet very little is known about their demography, particularly using MPMs. This
91dearth of insect data contrasts with their rather discrete life cycles, which renders them particularly
92amenable to modelling via MPMs (Caswell, 2001). These authors explore the demographic and
93performance effects of alternative host usage by a Neotropical treehopper (*Alchisme grossa*) to
94examine whether these effects exert a selective pressure sufficient for the evolution of species
95divergence. In the Bolivian Yungas forests, the subsocial treehopper (Hemiptera: Membracidae)
96utilises two sympatric hosts, *Brugmansia suaveolens* and *Solanum ursinum*. Adults use their natal
97host species, and females take care of nymphs (e.g., feeding facilitation, antidepredatory defense)
98during their development on both hosts. However, the duration of pre-imaginal development
99differs markedly between the host species, but two corresponding stage-structured MPMs have not
100revealed any statistical difference in their λ_1 s, hence individuals of *A. grossa*, though hatching to
101different hosts, still belong to a single-species population. Thus, the contribution opens a new
102direction for resolving the eternal dialectics of population dynamics vs. evolution.

103 Another important contribution from the angle of ecoregion representativeness is that by
104Santostasi et al. (2020^{SI}), who developed and introduced a framework to examine the role of
105hybridisation for two case studies, with one of them being in marine species (below). This is
106particularly important, as most of our demographic understanding across the Tree of Lie is

107constrained to terrestrial ecosystems, despite our dependence on the resources and ecosystem
108services that aquatic ecosystems provide (Tallis and Kareiva, 2005).

109

1102.2. Finding generality in ecology, evolution, and conservation biology using big data

111 Ecology, Evolution, and Conservation Biology are ideally positioned to tackle and provide
112solutions to the current biodiversity crisis of the Anthropocene (Proctor, 2013). However, most
113ecological, evolutionary, and conservation biology research that has been carried out to date with
114MPMs has been local in spatial extent and focus on single species. All species, from *Escherichia*
115*coli* to blue whales (*Balaenoptera musculus*), have in common the fact that individuals in their
116populations must invest to a certain degree in four key demographic (vital) rates: survival,
117development, reproduction, and recruitment. Notably, however, there is a great deal of variation in
118how individuals fine-tune the investment of the limited energy, via trade-offs, in each of these
119vital rates (Gaillard et al., 1989; Salguero-Gómez et al., 2016; Healy et al., 2019). However,
120despite the vast amount of life history strategy variation that emerges from this differential
121investment in vital rates, these vital rates ultimately allow species to persist and adapt to their local
122environments (Stearns, 1992).

123 The last decades have witnessed the utilisation of ever-growing volumes of published MPMs
124from open-access databases (Salguero-Gómez et al., 2015a, b) to tackle questions regarding the
125universality of the evolution of senescence (Jones et al., 2014; Salguero-Gómez, 2017), the link
126between functional traits and demographic performance (Adler et al., 2014), the predictive power
127of demographic schedules for species presence and absence (Csörgő et al., 2017), the examination
128of main axes of variation in life history strategies (Salguero-Gomez et al., 2016; Healy et al.,
1292019), or responses of organisms to climate change (Herrando-Perez, S., 2013).

130 Decomposition analyses in demographic studies, such as Life Table Response Experiments
131(LTRE, Caswell, 2000) have become a common, powerful way to examine ways in which
132experimental treatments affect the overall dynamics of a population via its underlying vital rates.
133These solutions have now been developed both for deterministic (Caswell, 2000) as well as
134stochastic environments (Caswell, 2010, 2019; Davidson et al., 2010). Davison et al. (2019^{SI})
135apply stochastic LTRE (sLTRE) tools to a subset of 220 populations from 62 species archived in
136COMADRE and COMPADRE. They find that stochasticity drives 28% of fitness effects,
137exceeding the mean effects by 7.78%, and thus further evidencing the need to describe
138populations and their abiotic effects in stochastic rather than deterministic terms.

139 Together, these novel contributions accentuate the promise of using large volumes of
140demographic data to find generality in ecology, evolution, and conservation biology.

141

1422.3. Eco-evolutionary dynamics

143 Classically, *evolution* (i.e., the change in phenotypes in populations linked to their heritability
144via natural selection) and *ecology* (i.e., the interactions of individuals with their own kind, other
145species, and their local environments) have been treated as self-contained disciplines acting in
146their separate silos. The main argument for their separation was a temporal one, under the
147assumption that evolution occurs at much longer time scales than ecology. However, research in
148the last decade has revealed that evolution can and oftentimes does shape individual performance
149and community dynamics at relatively short time scales (e.g., via rapid evolution; Hart et al.,
1502019). Similarly, ecological interactions can have long-lasting repercussions that place them right
151at the same temporal scale as classical evolutionary thinking (Hendry, 2017; Power et al., 2018).
152Demographic tools allow for the reconciliation of eco-evolutionary dynamics in a single, robust
153framework (Govaert et al., 2018; Takada and Shefferson, 2018; Shefferson and Salguero-Gómez,
1542015). Santostasi et al. (2020^{SI}) use a novel approach via matrix population models to examine the
155extinction of specific genomes in hybridizing populations. This approach is extremely relevant to
156our understanding of conservation biology, as hybridization is a main mechanism of biodiversity
157loss. In their paper, the authors model the breeding of two parental groups as separate
158components. They apply this model to two case studies regarding terrestrial (the wolf *Canis lupus*
159and its domestic counterpart *Canis lupus familiaris*) and marine species (the striped dolphin
160*Stenella coeruleoalba* and the common dolphin *Delphinus delphis*) facing hybridization. This
161article presents original utilisations of the matrix-based demographic machinery to advance our
162understanding of eco-evolutionary dynamics.

163

1642.4. Testing and overcoming classical assumptions in demographic modelling

165 Since the rigid postulates of Leslie's demographic model, testing and overcoming classical
166assumptions have been representing the main course of the mathematical developments of MPMs
167(Lefkovitch, 1965; Cohen, 1979; Tuljapurkar, 1990, 2013; Logofet, 1993, 2018; Cushing and
168Yicang, 1994; Caswell, 2001, and refs therein; Li and Schneider, 2002; Logofet, 2013, 2018). The
169following contributions make important progress along this course: de Vries et al. (2020^{SI}),
170Barraquand and Gimenez (2019^{SI}), and Takada and Kawai (2020^{SI}).

171 The contribution by de Vries et al. (2020^{SI}) provides methods to overcome two classical
172assumptions in MPMs. First, motivated by application to pesticide resistance in the flour beetle
173(*Tribolium castaneum*), the authors add genotype as the second basis to classify individuals (a
174rarely considered, yet important trait; but see Coulson et al. 2011), thus expanding the classical
175formalism to a general framework for the *stage*- and *genotype*-structured population. Second,
176when the classical linearity of demography meets the classical nonlinearity of population genetics,

177the PPM becomes a nonlinear density-dependent operator governing selection in this kind of
178population. The Jacobian matrix of this operator enables obtaining the stability conditions of
179homozygous equilibria, in particular, those of pesticide resistance domination. These results are
180useful for the pest management theory and they open a novel dimension in studying eco-
181evolutionary dynamic.

182 Barraquand and Gimenez (2019^{SI}) have overcome the classical MPM paradigm of single-
183species population dynamics by introducing a stage-structured model for a predator-prey
184community in which the populations are represented by juveniles and adults, predation is stage-
185specific, and the juvenile prey density positively affects predator fecundity. Estimating the
186parameters of species interaction in this framework requires a combination of traditional and novel
187ways to integrate data for single-species demography, thus leading to Integrated Community
188Models (ICMs). In a sense, this returns to the second birth of population dynamic models, i.e., the
189classical Lotka-Volterra prey-predator equations (Lotka, 1925; Volterra, 1931). Our authors
190assessed the value of different data sources using simulations of ICMs under different scenarios
191contrasting data availability and the presence/absence of intraspecific density-dependent effects.
192Combining all the data types (capture-recapture, counts, and reproduction surveys) allows the
193estimation of both demographic and interaction parameters, unlike count-only data which
194typically generate high bias and low precision in the interaction parameter estimates for short time
195series. As an accurate representation of stage structure in community dynamics, the ICMs open a
196wide spectrum of research perspectives, from the development of efficient observational study
197designs for monitoring communities in the field to the analysis of more sophisticated stage-
198structured MPMs.

199 Takada and Kawai (2020^{SI}) avail from the large volume of MPMs stored in the COMPADRE
200Plant Matrix Database (Salguero-Gómez, 2015a) to examine the most optimal way to analyse and
201depict elasticities of population growth rate with respect to vital rates in a comparative framework.
202*Elasticities*, i.e., the relative impacts of different demographic processes on the overall dynamics
203of a population (De Kroon et al., 1986, 2000) remain one of the most widely ways to quantify
204demographic processes' impacts on the viability of species (Morris and Doak, 2002). However,
205the ways in which they have been quantified and visualised (e.g., Enright et al., 1995; Franco and
206Silvertown, 2004) has not been free of criticism (e.g., Shea et al., 1994; Franco and Silvertown,
2071994). Specifically, Takada and Kawai use 68 MPMs from semelparous plant species in
208COMPADRE to examine the biologically meaningful patterns that should emerged from their
209elasticity vectors when presented in a ternary plot. Then they contrast these results to those of
210randomly generated MPMs with the same life history strategy. Their analyses help explain why,
211when the elasticities to survival, growth, and reproduction of plant species are plotted on a ternary

212plot, the overall pattern shows a strong curvature, with a “gap” in the ternary plot that is not
213“invadable”.

2142.5. Novel contributions to stochastic demography

215 Examining stochastic demographic properties by means of MPMs is typically done when
216more than one (a time series of) PPMs (“annual” PPMs) exist for a single population. This
217approach has become popular in ecology as most ecological systems are characterised by
218oscillatory environmental conditions (Salguero-Gómez and de Kroon, 2010), and climate change
219projects are likely to make natural systems suffer from a high frequency of climatic anomalies
220(IPCC, 2020). In this approach, the *stochastic growth rate*, λ_s , is typically defined via an infinite
221sequence of annual PPMs chosen at random from the given (finite) set of annual PPMs, and it is
222generally used as a proxy to population viability in the stochastic environment. In the special case
223of permanent non-stochastic environment, λ_s coincides with λ_1 , the asymptotic growth rate. On
224the one hand, theoretical approximations of λ_s exist that require certain additional assumptions
225regarding the given set of PPMs, such as the deviations from an average PPM with known
226temporal auto-covariances of the vital rates (Tuljapurkar, 1990; Caswell, 2001, Section 14.3.6.3).
227On the other hand, once a rule for the random choice of matrices, such as the *i.i.d.* (*independent*,
228*identically distributed* matrices, *ibid.*), the simplest and most popular rule (e.g. Cohen, 1979;
229Caswell, 2001; Buckley et al., 2010) has been accepted (but see Paniw et al., 2018), the λ_s can be
230estimated from a series of Monte Carlo simulations of the long enough random sequence.
231Unfortunately, the assumptions of both theoretical estimates of λ_s and Monte Carlo simulations
232may often not be accepted in practice. The challenge is therefore to attenuate those assumptions in
233response to theoretical and practical needs.

234 A key paper in this regard is by Sanz (2019^{SI}). In it, the author considers minimal and
235maximal vital rates among the given PPMs and obtains four new lower and upper bounds for the
236stochastic growth rate. These bounds enable calculating the sufficient conditions for population
237growth or population extinction. The author illustrates those conditions with a hypothetical
238random environment toggling between “poor” and “rich” situations.

239 Practical needs dictate rejecting the assumption of modelling stochastic dynamics of a natural
240population according to i.i.d. Instead, a more realistic approximation is to arrange the simulations
241in a way that reflect ways in which the natural environment oscillates through time (IPCC, 2020).
242Since the i.i.d. assumption represents a particular degenerate case of the time-homogeneous
243Markov chain (Logofet et al., 2020c), an important challenge for future research is to construct a
244governing Markov chain whose states would correspond to different types of environmental
245conditions and to calibrate its transition matrix from real data on those conditions.

246 The metapopulation environments may change randomly as well as that of a single
 247 population. In this special issue, Sanz and Bravo de la Parra (2019^{SI}) analyse the situation where
 248 the migrations between patches and demographic processes proceed at different time scales, the
 249 former being k times faster than the latter. Common formalisms for discrete-structured populations
 250 suggest that these two kinds of event occur sequentially one after the other, so that the combined
 251 effect of both processes can be described by the product of the slow-process matrix times the k^{th}
 252 power of the fast-process matrix. In nature, however, the processes may act simultaneously. The
 253 authors alternatively included survival into the fast process and approximated its effect during the
 254 fast time unit, referring to such an approach as the *re-scaling* of survival to the fast time scale.
 255 They adapt their original method of re-scaling to the proposed stochastic models and compared
 256 the asymptotic behaviour of the model populations with and without re-scaled survival. The
 257 outcome of principal importance is that the corresponding stochastic growth rates may appear
 258 qualitatively different for the same vital rate values: while the former predicts exponential growth,
 259 the latter indicates extinction. So, the order of events does matter in the MPM formalism for
 260 metapopulation dynamics in the stochastic environment; the both versions are ready in theory, and
 261 now it is the turn of real migrating populations to “migrate” right here.

262 Van Daalen and Caswell (2020^{SI}) introduce a framework to simultaneously quantify the
 263 variance components in demographic vs. environmental stochasticity in life history outcomes
 264 using matrix population models. The introduced mathematical machinery allows for the
 265 calculation of such effects in a general manner. The authors further illustrate the power of their
 266 approach using frailty analyses of fruit flies and explicitly quantifying survival–reproduction
 267 trade-offs in a plant study. This approach has the great potential to understand the extent to which
 268 demographic stochasticity vs. environmental stochasticity shape population outcomes worldwide.

269 As regards climate change, Romanov and Masterov (2020^{SI}) compare the effects of adult
 270 survival and productivity on the persistence of the Steller’s sea eagle (*Haliaeetus pelagicus*)
 271 populations. The authors simulated the corresponding MPM under several scenarios in a
 272 stochastic environment and found that the fecundity alone, varying within broad limits, can lead to
 273 population decline in this charismatic species. Although the population growth is less sensitive to
 274 the fecundity than to adult survival, the authors argue that the former is more appropriate as a
 275 target of conservation management.

276 2.6. Averaging the PPMs.

277 An alternative way to assess population viability in stochastic environments from a given set
 278 of annual PPMs is to calculate $\lambda_1(\mathbf{G})$, the dominant eigenvalue of the average matrix $\mathbf{G}$. However,
 279 in contrast to nonnegative numbers, it is not trivial to average several given nonnegative matrices
 280 unless the average is arithmetic or weighted arithmetic (due to the linearity of matrices as

operators in the vector space). Arithmetic average might make sense when the matrices to be averaged represent a *metapopulation* distributed among several habitats at the same moment of time, but it does not make sense when we have a time series of $m \geq 2$ PPMs, from the initial moment of observation (L_1) to the final one (L_m). If we want the average matrix, G , to project population vectors in the exactly same manner as the annual PPMs do, then it has to be their *multiplicative* (or *geometric* by the analogy with numbers) average (Logofet, 2018; Logofet et al., 2018a,b), which suggests extracting the m -th root of the matrix product.

To find a unique nonnegative matrix root is a matter of nontrivial calculations with a serious theory behind (Polity, Popolizio, 2014, 2015). A practical sufficient condition for the unique *principal root*, $A^{1/m}$, to exist for a given matrix A is the lack of negative/zero eigenvalues in the spectrum of A (*ibid.*), and the routine can really succeed in this case. However, what the routine returns will hardly have the same matrix pattern as the PPMs to be averaged – it will rather be positive-valued at best or complex-valued in general, the both cases making no sense as an average PPM that should correspond to the same LCG as the given PPMs to be averaged.

This requirement motivated a new concept of matrix average, namely, the *pattern-geometric* average (Logofet, 2018), or *pattern-multiplicative* average to be pedantic, meaning the same matrix pattern of the average matrix G as of those to be averaged. It is also logical that each element of G lies within the bounds of this element known from the set of given PPMs. Mathematically, finding the pattern-multiplicative average of a given set of annual PPMs reduces to a kind of constrained optimization problem, and this problem was efficiently solved for some practical cases (Logofet, 2018; Logofet et al., 2018a,b), but theoretical issues are still to be investigated in what concerns the uniqueness and accuracy of the optimal solution.

The λ_s and $\lambda_1(G)$, these two measures of viability assessment, generate the question of how they correlate quantitatively when applied in real cases, and Dmitrii Logofet has tried to give an answer (2019b^{SI}). Two former case studies of alpine Red-Book perennials (*Eritrichium caucasicum* and *Androsace albana*) resulted in respectively two sets of 8 annual PPMs, based on which both λ_s estimates and $\lambda_1(G)$ values are reported. These both measures of population viability turn out fairly close each other in both case studies, but *E. caucasicum* λ_s estimates are unambiguously less than $\lambda_1(G)$, whereas those for *A. albana* are certainly greater than $\lambda_1(G)$. The reason for this difference is not yet known, which gives one more incentive to further developing both the λ_s and $\lambda_1(G)$ methods.

2.7. Uses and abuses

314 As with any research tool that is both highly technical and popular, matrix population models
315 are not free of accidental mis-usage. We are not the first ones to call for caution in the
316 construction, analysis, and interpretation of MPMs (e.g., Caswell 2001). Our hope here is to be the
317 last ones. In this special feature, Nguyen et al. (2019^{SI}) and Kendall et al. (2019^{SI})-provide
318 invaluable advices on how to use this tool as well as the vast amount of demographic data
319 available in the format of an MPM.

320 Nguyen et al. (2019^{SI}) highlight once again (Doak et al., 2002) the importance of getting the
321 basic life cycle right before parameterising the pertinent matrix population model. There is a
322 wealth of MPMs for species across the Tree of Life (Werner and Caswell, 1977; Caswell 2001;
323 Salguero-Gómez, 2015a,b). However, some of the models, particularly in plants, may not contain
324 all, or the most, key life cycle stages. In the case of some plant species, permanent seed banks do
325 occur, but are often not included due to logistical practicalities of the fieldwork. Nguyen et al.'s
326 work show that their inclusion/exclusion can have very important repercussions for the ways we
327 assess the deterministic and stochastic dynamics of the populations under study. Importantly, the
328 emerging message therein is that the ways in which ignoring cryptic life stages affects our
329 estimates of population performance are not systematically biased in a specific way, rendering
330 generalisations with these key data rather limiting.

331 Kendall et al. (2019^{SI}) present a summary of a subset of published MPMs in animal species.
332 The authors highlight three errors commonly encountered in published MPMs: (1) failing to
333 include survival in the fertility coefficient; (2) introducing a one-year delay in the age at first
334 reproduction; and (3) incorrectly calculating the growth rate out of a stage class. The first two
335 errors are found in 34% and 62%, respectively, of the published studies collected in COMADRE;
336 of the studies where stages may last longer than one time step, 53% constructed the growth rate
337 using inappropriate formulas for estimating the asymptotic population growth rate or its sensitivity
338 to demographic parameters. Kendall et al. review the sources of such errors and reveal their
339 impact on model predictions, using lionfish and American alligator models as examples. These
340 results suggest that further efforts are required to educate biologists on the construction of MPMs,
341 and many studies that are based on MPMs may need to be re-examined; synthetic studies using
342 the COMADRE Database need to be accompanied by careful examination of the underlying
343 knowledge and data. Luckily, the researchers curating large demographic datasets are well aware
344 of these issues, and are expediently working together with the demographic community to help the
345 users of big data overcome these challenges. The solutions are already discussed in the response to
346 Kendall et al.'s paper in a complementary response by Che-Castaldo et al. (2020).

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3492.8. How many bases to classify individuals?

350 What is mentioned above concerns mostly what to do with ready MPMs. But to get them
351ready, given some knowledge of/data on a particular population is not a self-evident, nor a
352simple task. Historically, the ‘certain population mathematics’ adopted highly rigid postulates
353about the population *age* structure and proposed a highly specific pattern of the *Leslie* matrix
354(Leslie, 1945). Then, the postulates were attenuated and the pattern expanded to the *stage*
355structure (Lefkovitch, 1965) and to the *generalised stage* structure that suggested a
356classification of individuals by any observable/measurable trait such as size, weight, etc.
357(Caswell, 2001). Soon, a single trait became insufficient to practical demands where data on
358two traits, such as *age* and developmental *stage* (Logofet et al., 2006, 2014, 2016; Caswell,
359Salguero-Gómez, 2013), *stage* and *space* (Hunter, Caswell, 2005), or *stage* and *genotype* (de
360Vries et al., 2019^{SI}), were available, and the model formalism was expanded to block-
361structured PPMs (*ibid.*; Caswell, 2012; more samples of pairs in Roth and Caswell, 2016).
362The block-structured PPMs appear also in the single-trait metapopulation model with
363migrations among the local habitats (Sanz, Bravo de la Parra, 2019^{SI}); here the issue of time
364scale difference between demographic and migration processes needs further investigation.
365 Today, we witness an attempt to formalise an MPM for multi-trait population data, a 3-
366trait case in Coste and Pavard (2020^{SI}). The PPM transforms now to a 3D array of vital rates,
367which is called a *tensor* in mathematics. Authors have not yet dared to launch this term to the
368field of MPMs, but it will perhaps become common for the future generations of matrix
369modellers. Note however that a *hyperstate* matrix model, a competing term, has already been
370launched by Roth and Caswell (2016), with a general formalism for many bases.

3713. Concluding remarks

372 The diversity of topics presented above are sufficient to recognize the MPM as a powerful
373“multi-tool” for modeling the population and community dynamics. However, the range of topics
374covered in this editorial does not represent the wide spectrum of MPM applications in ecology,
375evolution, and conservation science. Below we discuss two (of many potential) areas of further
376enquiry where MPMs have much to offer to researchers.

3773.1. Potential-growth indicators

378 Among the current challenges of MPMs, studying their mathematical foundations is a key
379one. In particular, the most commonly used outcome of an MPM is $\lambda_1(\mathbf{L})$, its dominant
380eigenvalue, i.e., the asymptotic population growth rate, which is interpreted as a measure of
381population adaptation that is local in space and time. Matrix modellers are so focused on λ_1
382that they frequently forget its subscript 1 in the notation and what the subscript means itself.

In fact, the subscript originated from the standard of nonnegative matrix theory to order all the eigenvalues of a given $\mathbf{L}$ by their decreasing absolute values, so that the dominant eigenvalue (which is always positive by the Perron–Frobenius theorem) becomes λ_1 . However, recent findings have revealed $\lambda_2(\mathbf{L})$ to be important, too. Specifically, if λ_2 is positive and less than 1, then a simple function of matrix elements, $R_1(\mathbf{L}) = 1 - \det(\mathbf{I} - \mathbf{L})$, can serve as a *potential-growth indicator* (hereafter PGI). In other words, under these conditions, the PGI always locates at the same side of $\lambda = 1$ as $\lambda_1(\mathbf{L})$ does (Protasov, Logofet, 2014; Logofet, 2019a). There are certain merits of *indication* prior to the calculation of λ_1 ; in particular, $R_1(\mathbf{L})$ has turned out useful in solving calibration problems under data uncertainty (Logofet, 2013).

Another, more senior, PGI instance is the well-known *net reproductive rate* $R_0(\mathbf{L}) = \rho(\mathbf{F}[\mathbf{I} - \mathbf{T}]^{-1})$, where $\rho(\dots)$ denotes the *spectral radius* (the largest absolute value of matrix eigenvalues) and $\mathbf{L} = \mathbf{T} + \mathbf{F}$, the sum of its transition and fertility parts (Cushing, Yicang, 1994; Caswell, 2001). However, calculating the spectral radius is, in general, much more difficult than the determinant in $R_1(\mathbf{L})$. The problem with $R_1(\mathbf{L})$ is that its PGI property is proved for a less general class of PPM patterns than $R_0(\mathbf{L})$ is valid for.

A recent expansion of the classical Perron–Frobenius theory, namely, the theory of *rank-1 corrections* of nonnegative matrices (Protasov, Logofet, 2014), has enabled proving the PGI property of $R_1(\mathbf{L})$ when the rank of fertility part $\mathbf{F}$ equals 1. For instance, when $\mathbf{F}$ has only one nonzero row or column of fertility rates meaning a single recruiting or a single generative stage in the LCG, respectively. Can this wide, yet not universal, class of PPMs be expanded? Some particular examples give a positive answer, but theoretical justification is still a target for future efforts.

3.2. Integer-valued formalism

The ‘identified individuals’, a standard for stage-structured demographic models (Caswell, 2001, p. 134), can often provide the calibration of PPMs in a unique way. Since all the events of transitions along the LCG links are inherent in the data, the transition rates are just the frequencies of the corresponding events, hence the rates are certain *rational* numbers according to the arithmetic way of calculation. Naturally, the number of recruited individuals is always integer. However, the tradition is to treat the basic calibration equation of the MPM as an equation in *real* numbers in spite the rational-valued PPM does project $\mathbf{x}(t)$, an integer-valued population vector, into the integer-valued $\mathbf{x}(t + 1)$.

The old tradition to deal with real numbers did make certain sense in the last century, when computers had only been dealing with binary representations of all numbers including the machine approximations of the irrational roots of the characteristic polynomials. However,

417modern systems of mathematical software, such as R or MatLab, with its Symbolic Algebra
418Toolbox (Mathworks, 2020), enable algebraic manipulations with the rational-/integer-valued
419numbers as symbols, thus avoiding the machine round-off errors and providing for the
420absolute (logical) accuracy of calculations. Also, the existence of strong computational
421support enables us to validate each step of the calibration or manipulation in other related
422problems, e.g., backward prediction (Clutton-Brock and Sheldon, 2010; Ehrlén et al., 2016;
423Nater et al., 2018; Logofet et al., 2020a) or uncertainties in the dynamics of cryptic stages
424(Paniw et al., 2017; Nguen et al., 2019^{SI}; Logofet et al., 2020b^{SI}). The calibration equation or
425other pertinent expressions hold true, with the absolute accuracy, for the integer- or rational-
426valued parameters, unlike the approximate real values to be found from the classical least-
427squares procedures of curve fitting. This approach helps solve at least some of the ‘persistent
428problems in the construction of matrix population models’ and we encourage matrix
429modellers to incorporate them in their demographic enterprises.

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