THE PARADOX OF VITO VOLTERRA'S
PREDATOR-PREY MODEL

JEAN-MARC GINOUX$^{1,2}$

This article is dedicated to the late Giorgio Israel.

Résumé. The aim of this article is to propose on the one hand a brief history of modeling starting from the works of Fibonacci, Robert Malthus, Pierre Francis Verhulst and then Vito Volterra and, on the other hand, to present the main hypotheses of the very famous but very little known predator-prey model elaborated in the 1920s by Volterra in order to solve a problem posed by his son-in-law, Umberto D’Ancona. It is thus shown that, contrary to a widely-held notion, Volterra’s model is realistic and his seminal work laid the groundwork for modern population dynamics and mathematical ecology, including seasonality, migration, pollution and more.

1. A short history of modeling

1.1. The Malthusian model. If the first scientific view of population growth seems to be that of Leonardo Fibonacci [2], also called Leonardo of Pisa, whose famous sequence of numbers was presented in his Liber abaci (1202) as a solution to a population growth problem, the modern foundations of population dynamics clearly date from Thomas Robert Malthus [20]. Considering an “ideal” population consisting of a single homogeneous animal species, that is, neglecting the variations in age, size and any periodicity for birth or mortality, and which lives alone in an invariable environment or coexists with other species without any direct or indirect influence, he founded in 1798, with his celebrated claim “Population, when unchecked, increases in a geometrical ratio”, the paradigm of exponential growth. This consists in assuming that the increase of the number $N(t)$ of individuals of this population, during a short interval of time, is proportional to $N(t)$. This translates to the following differential equation:

\[ \frac{dN(t)}{dt} = \varepsilon N(t) \]

where $\varepsilon$ is a constant factor of proportionality that represents the growth coefficient or growth rate. By integrating (1) we obtain the law of exponential growth or law of Malthusian growth (see Fig. 1). This law, which does not take into account the limits imposed by the environment on growth and which is in disagreement with the actual facts, had a profound influence on Charles Darwin’s work on natural selection. Indeed, Darwin [1] founded the idea of “survival of the fittest” on the

---

1. According to Frontier and Pichod-Viale [3] the correct terminology should be population “kinetics”, since the interaction between species cannot be represented by forces.

2. A population is defined as the set of individuals of the same species living on the same territory and able to reproduce among themselves.
impossibility of an indefinite population growth. He illustrated this impossibility by a superb parabola describing the descendants of a pair of elephants that, under optimal conditions, would cover the surface of the Earth in a few centuries. However, in laboratory experiments, the predictions of the Malthusian law remain correct on small numbers, while there is divergence for high values of the population. Thus, we are led to conclude that the exponential law remains valid as long as the density of the population does not saturate the environment.

Figure 1. Malthus’s model of exponential growth.

1.2. P.F. Verhulst’s model. It was on the basis of these considerations that the Belgian biologist Pierre-François Verhulst [28] proposed in 1837 a model that took into account the limitation imposed by the increasing population size:

$$\frac{dN(t)}{dt} = \varepsilon N(t) - \lambda N^2(t) = \varepsilon N(t) \left(1 - \frac{1}{K} N(t)\right)$$

where $\varepsilon$ represents the growth rate. The second coefficient $\lambda = \varepsilon / K$ originates in a “mechanistic” interpretation of the phenomenon. Indeed, it is assumed that growth is limited by a kind of interior “friction” within the population, that is, the resources remaining the same, the higher the number of individuals, the more difficult it is for them to feed themselves and then to grow. This is a struggle between individuals for the existence, and so an intra specific competition for food. The factor $K$, called carrying capacity, corresponds to the capacity of the environment to support the population growth and represents the population limit beyond which it can no longer

3. An example of this impossibility is also shown in the film by V.A. Kostitzin and J. Painlevé titled : “Images mathématiques de la lutte pour la vie”, 1937, Médiatèque du Palais de la Découverte, Paris.
grow. This law, which Verhulst called *logistic equation*, is radically different from Malthus’s, since it imposes a limiting value on the population (see Fig. 2 where exponential growth is represented in blue and logistic growth in red). It has been successfully applied to many real life situations, such as population growth in the United States between 1790 and 1950 (see Pearl and Reed [22]) or in experiments conducted by the Russian biologist Georgii Frantsevich Gause [4] on the growth of a protozoan, *Paramecium caudatum*. Independently of these two archetypes, other growth models have been developed: let us mention, for instance, the model by B. Gompertz [9] aimed at evaluating the growth rate of a tumour.

![Figure 2. Verhulst’s model of logistic growth (in red).](image)

1.3. **Volterra’s “predator-prey” model.** In the first half of the twentieth century, the study of the dynamics of several interacting species developed considerably. It was at this time, called “the golden age of theoretical ecology” [26], that the first models based on competition-type behaviours and predator-prey relationships were developed. The paternity of the first model used to transcribe this kind of interactions was the subject of a quarrel between Alfred J. Lotka [18] and Vito Volterra [4], which is described in the work of Giorgio Israel [13, 14, 15, 16]. It has been established that the merit of the development of the first model of predator-prey type applied to a biological situation is Volterra’s. Indeed, Volterra’s interest in problems of equilibria between animal species in ecosystems was prompted by his son-in-law, zoologist Umberto D’Ancona, who for some years had been dealing with statistics on fishing in the northern Adriatic Sea. These data involved the percentage of predatory fish (Selachians) caught in three Italian ports, Trieste, Fiume (now Rijeka) and Venice, during the period 1905-1923. They proved that during the period 1915-1920, when fishing was less intense because of the war, there had been a relative

---

4. For more about the dispute over priority, see [10], pp. 146-147.
increase in the Selachians. According to D’Ancona’s hypothesis, fishing disturbed the natural balance between species. It favoured a relative increase in “prey” species, that is, fish that feed only on plankton, and a decrease in “predatory” species, that is, fish that feed on other fish. The decline in fishing due to World War I had thus restored, at least in part, the natural balance. D’Ancona turned to Volterra, asking him to find a mathematical proof of his hypothesis. In 1926 Volterra published a reply in Italian [29] which took the form of the famous “predator-prey” model, reproduced below. He then presented his results in English and, in more detailed and complete form, in French [30, 31, 32]. In the first chapter of his “Leçons sur la théorie mathématique de la lutte pour la vie” [32], Volterra studies the coexistence of “two species, one of which devours the other”. Considering two species, the first, the prey \( N_1(t) \), would undergo a Malthusian growth if it were alone. The second, the predator \( N_2(t) \), feeds exclusively on the first and in the absence of prey will gradually wear out and disappear. The formulation of the equation representing the predation is based on the méthode des rencontres (“method of encounters”) and on the hypothèse des équivalents (“hypothesis of equivalents”) due to Volterra [32].

The former assumes that for predation to occur between a predatory species and a prey species, it is first necessary to have encounters between these two species and that the number of encounters between them is proportional to the product of the number of individuals composing them, that is, \( N_1(t) N_2(t) \), the coefficient of proportionality being equal to the probability of an encounter. The second hypothesis consists in assuming that “there is a constant ratio between the disappearances and appearances of individuals caused by the encounters”, that is, that predation of the prey is equivalent to increase of the predators. At the beginning, Volterra considers this increase as immediate. This led him to the system:

\[
\begin{align*}
\frac{dN_1}{dt} &= \varepsilon_1 N_1 - \gamma_1 N_1 N_2 = N_1 (\varepsilon_1 - \gamma_1 N_2) \\
\frac{dN_2}{dt} &= -\varepsilon_2 N_2 + \gamma_2 N_1 N_2 = -N_2 (\varepsilon_2 - \gamma_2 N_1)
\end{align*}
\]

where \( \varepsilon_1 \) represents the prey’s growth rate in the absence of the predator; \( \gamma_1 \) the predation rate of the predator on the prey; \( \varepsilon_1 \) the predator’s mortality rate in the absence of prey; and \( \gamma_2 \) the growth rate of the predator due to its predation. From this model, Volterra was able to state the law of the disturbance of the averages:

If an attempt is made to destroy the individuals of the two species uniformly and in proportion to their number, the average of the number of individuals of the species that is eaten increases and that of the individuals of the species feeding upon the other diminishes [31, p. 20]

To establish this result, Volterra assumes that, for a time interval \( dt \), we destroy \( \alpha \lambda N_1 dt \) prey and \( \beta \lambda N_2 dt \) predators. He then proves that the average values of prey and predators, which were previously equal to \( \varepsilon_2/\gamma_2 \) and \( \varepsilon_1/\gamma_1 \), respectively, become, after this destruction, \( (\varepsilon_2 + \beta \lambda)/\gamma_2 \) and \( (\varepsilon_1 - \alpha \lambda)/\gamma_1 \). The phenomenon observed by D’Ancona is thus explained: the increase in the number of predators

---

5. This means that predation is immediately transcribed in terms of growth of the predator species, whereas its effect naturally occurs with some delay. It will be seen below that Volterra also took this delay into account.
and the reduction in the number of prey resulted from the decline in fishing, which before the war had changed the natural balance of this “biological association”. A decrease in fishing favours the more voracious species at the expense of the other. This deterministic model, which constitutes the archetype of the trophic network, aims at transcribing different types of animal behaviour into mathematical functions. Two types of behaviour are represented: those related to increase and those related to decrease. Natality and predation are related to increase, whereas natural mortality and mortality by predation correspond to a decrease in the number of individuals. Each of these behaviours has a mathematical form, called a functional response.

1.4. Different types of functional responses. Since the mid-1920s, these functional responses have been the subject of numerous studies and developments aimed at making the representation of animal behaviour by a mathematical function more realistic. Natural growth, that is by natality, of the prey, represented by a functional response of Malthus type \[20\] was later modified by Verhulst \[28\] to account for its being bounded. The decrease by natural mortality was initially considered in a way symmetrical to that of natural growth, that is, by substituting in the equations \[11\] and \[12\] the growth rate \(\varepsilon\) with a natural mortality rate \(-\varepsilon\). It is important to emphasise that this functional response, sometimes called closure relation, was later the subject of special studies aimed at transcribing the specific behaviour of certain species, for instance cannibalism. All these functional responses are summarized in Table 1.

|            | natural increase | natural mortality |
|------------|------------------|-------------------|
| Malthus    | \(\varepsilon N(t)\) | \(-\varepsilon N(t)\) |
| Verhulst   | \(\varepsilon N(t) \left(1 - \frac{N(t)}{K}\right)\) | \(-\varepsilon N(t) \left(1 + \frac{N(t)}{K}\right)\) |

Table 1. Functional responses for natural increase and decrease.

The functional response proposed by Volterra \[32\] to describe predation and which was based on the principle of encounters was proportional to the product of the number of individuals of each species: \(N_1(t)N_2(t)\). In other words, the predation rate was a linear function of the prey, that is, \(N_1(t)\). A few years later, Gause \[4, 5\], who was one of the first to make “experimental verifications of the mathematical theory of the struggle for life”, proposed another type of functional response to describe predation, a “nonlinear” one, aimed at transcribing a certain “satiety” of the predator with respect to its prey: \(N_1^g(t)N_2(t)\) with \(0 < g \leq 1\). In this case, the predation rate becomes indeed a “nonlinear function” of the prey, that is, \(N_1^g(t)\) (see Fig. 3).

In the late 1950s, entomologist Crawford Stanley Holling \[11, 12\] developed, from the celebrated “disc equation” two new functional responses for predation, also intended to describe a certain satiety of the predator with respect to its prey: Holling function of type II and Holling function of type III. This formulation assumes that the predator divides its time between two kinds of activities: the search for its prey,
and its capture, which includes the time spent hunting, killing, devouring and digesting it. Holling’s type II (see 4) is a functional response in which the predator’s “attack” rate increases when the prey number is low and then becomes constant when the predator reaches satiety. In other words, the predator causes maximum mortality at low prey densities. Thus, type II functional responses are typical of predators specializing in attacking one or few prey. In this case the mortality of the prey decreases with their density. Holling’s type II is represented by:

(4)  \[
\frac{N_1(t)}{h + N_1(t)} N_2(t)
\]

where \(h\) represents half-saturation, that is, the value of the prey density \(N_1(t) = h\) for which the predation level reaches a value equal to half its maximum.

Holling’s type III (see Fig. 5 where Holling’s type II is represented in blue and Holling’s type III in red) is a functional response in which the attack rate of the predator first increases when the prey number is low and then slows down when the predator reaches satiety. In other words, the predator increases its research activity when prey density increases. Thus, type III functional responses are typical of generalist predators moving from one species of prey to another and concentrating their activities in areas where resources are abundant. In this case, the mortality of the prey initially increases with their density and then decreases. Holling’s type III is represented by:

(5)  \[
\frac{N_1^2(t)}{h^2 + N_1^2(t)} N_2(t)
\]

where \(h\) represents half-saturation, as above.
Contrary to what one might think, Holling did not deduce his two functional responses from the observation of a natural environment, but, as he himself wrote in his article, by concocting an artificial predatory-prey situation:
In the first artificial situation devised the “prey” were sandpaper discs four centimetres in diameter thumb-tacked to a three-foot square table. A blindfolded subject, the predator Š, stood in front of the table and searched for the discs for one minute by tapping with her finger. As each disc was found, it was removed, set to one side and searching continued. Each experiment was replicated eight times at densities of discs ranging from four to 256 per nine sq. ft. The results of one such experiment are shown in Fig. 1 [see Fig. 4 here], where it can be seen that the number of discs picked up increased at a progressively decreasing rate as the density of discs rose. [12, p. 385]

Mathematically, it has been proved by Real [24] that the functional responses of Holling type II and III are analogous to the function developed in 1913 by Leonor Michaelis and Maud Menten [21] to describe the kinetics of enzymatic reactions. All these functional responses are summarized Table 2.

| Volterra | Gause | Holling type II | Holling type III |
|----------|-------|----------------|------------------|
| \( N_1(t) N_2(t) \) | \( N_1(t) N_2(t) \) | \( \frac{N_1(t)}{h + N_1(t)} N_2(t) \) | \( \frac{N_1^2(t)}{h^2 + N_1^2(t)} N_2(t) \) |

Table 2. Functional responses for predation

2. The Origin of the Paradox

According to Yuri A. Kuznetsov [17], in the context of a Volterra predator-prey model, the functional responses that limit growth (see Table 1) have a stabilizing effect for the prey whereas the functional responses that limit predation (see Table 2) have a destabilizing effect for predators. Thus, in the Volterra predator-prey model [8] by restricting the growth of the prey \( N_1(t) \) by a Verhulst functional response and the growth of the predator \( N_2(t) \) by a functional response of Holling type II, we obtain the celebrated Rosenzweig-MacArthur model [25]:

\[
\begin{align*}
\frac{dN_1}{dt} &= \varepsilon_1 N_1 - \lambda N_1^2 - \gamma_1 \frac{N_1}{h + N_1} N_2 \\
\frac{dN_2}{dt} &= -\varepsilon_2 N_2 + \gamma_2 \frac{N_1}{h + N_1} N_2
\end{align*}
\]

where \( \varepsilon_1 \) represents the prey’s growth rate in the absence of the predator, \( \gamma_1 \) the predator’s predation rate on the prey, \( \varepsilon_2 \) the predator’s mortality rate in the absence of prey, and \( \gamma_2 \) the predator’s growth rate due to its predation. The combination of these two effects then leads to the existence of a periodic solution. Henri Poincaré [23] named this solution, or more exactly this periodic oscillation, limit cycle. This terminology derives from the fact that in the phase space \((N_1, N_2)\), in this case the prey-predator space, it takes the form of a cycle towards which every
solution converges asymptotically. Thus, the solution of the Rosenzweig-MacArthur model (6) is periodic, like that of the Volterra model (3). Nevertheless, there is a fundamental difference in that it is totally independent of the initial conditions, that is, of the initial density of prey and predators, which is consistent with reality. Indeed, in nature, the amplitude and period of the periodic oscillations of prey and predators cannot depend on their initial densities considered at a given time arbitrarily taken as the origin of the time [6, 7, 8].

3. Discussion

Volterra’s model, which in the study of nonlinear dynamic systems has become a kind of paradigm, has been the subject of many books and articles criticizing it for its lack of realism. The main flaws of this model are considered to be the absence of limitation in the growth of prey and predator, the fact of not having taken into account seasonality (that is, having considered the growth rate of prey and predator as constant), and finally having proposed an ideal, simplified model by limiting it to two species. The analysis of these flaws will shed light on the “paradox” of Volterra’s model. In the first place, it should be recalled that even today it is impossible to formulate a simple expression of the periodic solutions of Volterra’s model (3), that is, an expression that uses only elementary functions. Consequently, criticizing Volterra for having proposed an ideal, simplified model is inadmissible. Indeed, it is precisely thanks to the simplicity of his model that Volterra was able to solve the problem posed by D’Ancona and explain the phenomenon by establishing the law of perturbation of means. In fact, Volterra was well aware of the limitations of the model and the assumptions attached to it, including the one assuming the environment as invariable and without influence on the growth rate and the one that considers the homogeneity of the individuals of each species. Indeed, in [32] Volterra laid the foundations for what would be much later called “population dynamics” and considered all the aspects of the problem: the influence of environment or pollution, the heterogeneity of individuals or age groups (on hereditary actions [32, p. 141]), diffusion or migration (contribution of a small number of individuals [32, p. 118]), seasonality (variation of exterior conditions with time, [32, p. 131]). In the introduction, he writes:

Certainly there exist periodic circumstances relating to environment, as would be those, for example, which depend upon the changing of the seasons, which produce forced oscillations of an external character in the number of individuals of the various species. These actions of external periodic nature were those which were specially studied from the statistical point of view, but are there others of internal character, having periods of their own which add their action to these external causes and would exist even if these were withdrawn? [32, p. 5].

Then, he adds:

Later, it will be observed that it is closer to reality to suppose that the growth coefficients depend not only, at each moment, on the current values of the quantities \(N_i\) (characterizing the species \(i\)), but also on past values up to a more or less remote period. It will no longer be sufficient to consider them as functions of the
\(N_i\), but as “functionals”, and this will lead us to integro-differential equations that we will approach from the equations we are led to in the so-called “hereditary” mechanics [32, p. 5].

It is in this context that he invents the famous Volterra equations of the first and second type. With regard to the limitation to a two-species model of predator-prey type, in chapter II of [32] Volterra presents the “study of the coexistence of any number of species”. Moreover, the chapter ends with the analysis of a “very remarkable case: that of three species of which the first feeds on the second and this on the third”. Different types of interactions between species are also considered outside of predation, including competition, cooperation and migration. Regarding the absence of limitation in the growth of prey and predators, Volterra proposes in chapter III of [32] to study a model of \(n\) coexisting species having reciprocal actions, in which he replaces the Malthusian growth by a Verhulst logistic growth. Thus, Italian or French authors who claim that Volterra’s model is unrealistic have obviously not read Volterra’s texts [30, 31, 32] and this is clearly one of the origins of this paradox. Indeed, it seems that Volterra’s model and criticisms of it are known worldwide, but his work has not even been read by his detractors! For authors who only read English, this paradox is rooted in another problem. While the original Italian version of Volterra’s study [29] comprised 84 pages, the very first English translation published in the prestigious journal Nature [30] was only two pages long. It is easy to understand that for many Volterra’s study may have seemed very limited given the conciseness of this summary, which did not allow the exposition of analytical developments and considerably reduced the scope of Volterra’s results. Nevertheless, and this is again a paradox, even though by 1928 there was an integral English translation of the original Italian text by Volterra [31], it seems to have been almost totally ignored by English-speaking authors. Thus, Volterra’s work is perhaps the most quoted and the most criticized in the world while being the least read and studied.

From 1926, Volterra’s publications intensified in the field of population dynamics. It was in 1931 that his work entitled \textit{Leçons sur la théorie mathématique de la lutte pour la vie} [32] was published in French, following a series of lectures at the Institut Henri Poincaré where Volterra had been invited by Borel. Compiled by Marcel Brelot, this book contains Volterra’s entire memoir on biological fluctuations [29] as well as a part about the case where heredity plays a role. From 1936, Volterra questioned the validity of his model and the possibility of experimental verification. The experiments of Gause (1910-1989) seemed to confirm the first law, and he showed Volterra his acceptance of the model’s forecasts. But what followed was disappointing because they could not find irrefutable cases of a cyclic behaviour in predator-prey ecosystems. Moreover, voiced by biologists Karl Pearson (1857-1936) and Friedrich Simon Bodenheimer (1897-1959), objections multiplied to the point of questioning the interpretation of D’Ancona’s statistics, which led to D’Ancona himself coming to doubt their merits. At present, the only example of a predator-prey ecosystem showing a cyclic evolution is the famous set of statistics by Hudson’s Bay Company about hares and lynxes in Canada. According to Giorgio Israel, it is thus important to note that unlike Van der Pol’s model [27]:

6. Under the same title, in a later issue of the same journal, two letters were published, one by Lotka and one by Volterra himself [19, 30].

7. To the best of our knowledge, this book has never been translated into English.
Volterra’s model is not deduced from an analogy but from a more traditional approach that consists in starting from the analysis of a real phenomenon, making some abstraction of the accessory aspects such as friction, determining the state variables, formulate a mathematical hypothesis concerning the pace of the phenomenon. . . . For Volterra’s model, the problem of the experimental verification consists in finding empirical evidence directly verifying the laws deduced from the model and not justifying its validity in an indirect way, that is, from the effectiveness of some of its consequences. Thus, for Volterra, the justification of D’Ancona’s hypothesis on the effects of fishing was not sufficient to prove the empirical validity of the model. It is because of this conviction that he sought for the rest of his life a direct empirical proof of the validity of the first law according to which the evolution of populations presents periodic oscillations [27].

Thus, if Volterra sought to make his mathematical research work at the service of biological sciences, it seems that it always was with the aim of explaining phenomena and describing reality as faithfully as possible:

“ . . . the hypotheses will be seen to become more and more complex in order to get closer to reality . . . ”

Translated from the French by Daniele A. Gewurz

Acknowledgments

I wish to express my sincerest thanks to my friend Christian Gérini, Agrégé de Mathématiques and Docteur en Histoire des Sciences, who encouraged and supported me in my work.

Références

[1] Darwin, C., “The Origin of the Species,” W. Clowes & Sons, London, (1859).
[2] Fibonacci, L., Liber abaci, reproduit dans, “A Translation into Modern English of Leonardo Pisano’s Book of Calculation,” Springer-Verlag, New-York, (2004).
[3] Frontier, S. & Pichod-Viale, D., Ecosystèmes, structure, fonctionnement, évolution, Dunod, Paris, (2001).
[4] Gause, G.F., The struggle for existence, Williams and Wilkins, Baltimore, (1935).
[5] Gause, G.F. vériﬁcations expérimentales de la théorie mathématique de la lutte pour la vie, Hermann, Paris, (1935).
[6] Ginoux, J.M. Analyse mathématiques des phénomènes oscillatoires non linéaires, Thèse, Université de Paris VI (2011).
[7] Ginoux, J.M. Histoire de la Théorie des Oscillations Non Linéaires, Hermann, Paris, (2015).
[8] Ginoux, J.M. History of Nonlinear Oscillations Theory, Archimedes Series, Vol. 49, Springer International Publishing (2017).
[9] Gompertz, B., “On the Nature of the Function Expressive of the Law of Human Mortality, and on a New Mode of Determining the Value of Life Contingencies,” Phil. Trans. Roy. Soc. London, 123, 513-585, (1832).
[10] Guerraggio, A. & Paolini, P., Vito Volterra. Trans. Kim Williams. Springer-Verlag, Berlin (2013).
[11] Holling, C.S., “The components of predation as revealed by a study of small-mammal predation of the European pine sawfly,” Canadian Entomologist, 91, 293-320, (1959).
[12] Holling, C.S., “Some characteristics of simple types of predation and parasitism,” Canadian Entomologist, 91, 385-398, (1959).
[13] Israel G., “Volterra’s Analytical Mechanics of Biological Associations”, Arch. Int. Hist. Sci., 41, 57-104 ; 307-352, (1991).
[14] Israel G., “The Emergence of Biomathematics and the Case of Population Dynamics : A Revival of Mechanical Reductionism and Darwinism”, Sci. in Context, 6, 469-509, (1993).
[15] Israel, G., La Mathématisation du réel : essai sur la modélisation mathématique, Seuil, Paris, (1996).
[16] Israel, G., Millán GASCA, A., The Biology of Numbers. The Correspondence of Vito Volterra on Mathematical Biology, Basel, Birkhäuser, Boston, (2002).
[17] Kuznetsov Y., Elements of Applied Bifurcation Theory, Springer-Verlag, Berlin, (1998).
[18] Lotka, A., Elements of physical biology, Williams & Wilkins Co, Baltimore, (1925).
[19] Lotka, A., “Fluctuations in the abundance of a species considered mathematically,” Nature 119, 12 (1927)
[20] Malthus, T.R., An Essay on the Principle of Population, printed for J. Johnson, in St. Paul’s Church-Yard, London (1798).
[21] Michaelis, L. & Menten, M. L., “Die Kinetik der Invertinwirkung,” Biochemische Zeitschrift 49, 333-369 (1913).
[22] Pearl, R. & Reed, L.J., “On the rate of growth of the population of the United States since 1790 and its mathematical representation,” Proceedings of the National Academy of Science (U.S.A.), Vol. 6, 6, 275-288, (1920).
[23] Poincaré, H., “Sur les courbes définies par une équation différentielle,” Journal de mathématiques pures et appliquées, (III) 8, 251-296, (1882).
[24] Reall, L., “The kinetics of functional response,” American Naturalist, 111, 289-300, (1977).
[25] Rosenzweig, M. & MacArthur, R., “Graphical representation and stability conditions of predator-prey interaction,” American Naturalist, 97, 209-223, (1963).
[26] Scudo, F.M. & Ziegler, J.R., The Golden Age of Theoretical Ecology : 1923-1940, Springer-Verlag, Berlin, (1978).
[27] Van der Pol, B., “On 'Relaxation-Oscillations',' Phil. Mag., 7, Vol. 2, 978-992, (1926).
[28] Verhulst, P.F., “Notice sur la loi que suit la population dans son accroissement,” Corresp. Math. Phys., X, 113-121, (1838).
[29] Volterra, V., “Variazioni e fluttuazioni del numero d’individui in specie animali conviventi,” Mem. Acad. Lincei III, 6, 31-113 (1926).
[30] Volterra, V., Fluctuations in the abundance of a species considered mathematically (“ Nature “, vol. CXVIII, 1926, pp. 558—560). Sotto lo stesso titolo furono poi pubblicate due lettere, una del LOTKA e una del VOLterra (Ibidem, vol. CXIX, 1927, pp. 12—13).
[31] Volterra, V., Variation and fluctuations of the number of individuals in animal species living together. Translated by Miss MARY EVELYN WELLS (“Journal du Conseil international l’exploration de la mer”, Copenhague, vol. III, n. 1, 1928, pp. 3—51).
[32] Volterra, V., Leçons sur la Théorie Mathématique de la Lutte pour la Vie, Gauthier-Villars, Paris, (1931).

1 Laboratoire LSIS, CNRS, UMR 7296, Université de Toulon, BP 20132, F-83957 La Garde cedex, France
2 Archives Henri Poincaré, Université de de Lorraine, CNRS, UMR 7117, BP 454, F-54001 Nancy, Cedex, France
E-mail address: ginoux@univ-tln.fr