A New Method to Evaluate Simulation Models: The Calibration Profile (CP) Algorithm

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Abstract

Models of social systems generally contain free parameters that cannot be evaluated directly from data. A calibration phase is therefore necessary to assess the capacity of the model to produce the expected dynamics. However, despite the high computational cost of this calibration it doesn’t produce a global picture of the relationship between the parameter space and the behaviour space of the model. The Calibration Profile (CP) algorithm is an innovative method extending the concept of automated calibration processes. It computes a profile that depicts the effect of each single parameter on the model behaviour, independently from the others. A 2-dimensional graph is thus produced exposing the impact of the parameter under study on the capacity of the model to produce expected dynamics. The first part of this paper is devoted to the formal description of the CP algorithm. In the second part, we apply it to an agent based geographical model (SimpopLocal).

The analysis of the results brings to light novel insights on the model.

Keywords: Calibration Profile, Model Evaluation

Introduction

1.1 Agent-based modelling is widely used in models of social systems because it is well suited to describe individual-centred dynamics with non-linear interactions. This modelling technique, however, introduces many free parameters which critically influence the behaviour of the model. Hand-picking parameters is a time-consuming process and makes it hard to evaluate the ability of a model to reproduce empirical data or formalised knowledge. Automatic calibration of models have the potential to free the modeller from this burden and to make the modelling process more rigorous by reducing the number of arbitrary choices.

1.2 The current prevalent method for automatic calibration is to see it as an optimisation problem in which the differences between known data and model predictions have to be minimised. This optimisation can be performed with different optimisation algorithms like “Approximate Bayesian Computation” (Lenormand et al. 2012) or evolutionary algorithms (Schmitt et al. 2014; Stonedahl 2011). Nevertheless, these methods find a single set of parameters, or, in the case of multi-objective evolutionary algorithms, several sets of parameters that represent optimal trade-off with regard to several model quality criteria. In both cases, the result of the calibration is solely that the model can reproduce the data with a given precision. These calibration processes do not say anything about how often parameter sets lead to realistic behaviours, and about how each parameter changes the behaviour of the model. For instance, it is often interesting to know when some parameter values would prevent the system to reach a realistic behaviour, rather than only knowing a single set of “optimal” parameter values.

1.3 To compute a more global view of the parameter space, we have designed a novel method that exposes the sensitivity of a single parameter on the calibration of a model independently of the other parameters. A high level representation of this method is shown on Figure 1. It computes the lowest calibration error that can possibly be obtained when the value of a given parameter is fixed and the others are free. It computes this minimal error for many values of the parameter under study. The values of the parameter are sampled all along its domain of definition to produce a so called calibration profile. For each sample value, the value of the remaining parameters are optimised in order to find the lowest possible calibration error. The profile can then be drawn on a 2-dimensional chart that depicts the influence of the parameter under study on the model calibration.

1.4 Conceptually, CP are similar to likelihood profiles (Pawitan 2001), a classic tool in statistics[1]. However, computing likelihood profiles requires an analytical expression of the model, which is unfortunately not available in most agent-based simulations, whereas calibration profiles can be used with any model. A calibration profile can thus be seen as a way to approximate the likelihood profile when the analytical expression of the model is not available or is too complicated.

1.5 To produce a calibration profile, a naive approach would consist in executing an entire calibration algorithm for each value of the parameter under study. Current automated calibration algorithms are too computationally intensive to make this approach tractable in practice. To tackle this problem we have designed an algorithm that computes the numerous points composing a calibration profile altogether. The design of this algorithm has been inspired by the recently published MOLE method.
Clune et al. (2013; Mouret & Clune 2012), which pictures 2-dimensional phenotype landscapes for evolutionary biology. MOLE and the CP method are both stochastic, population-based search algorithms that exploit the fact that a small change to a good solution is likely to lead to another similarly good solution (Deb et al. 2002). Nonetheless, while classic stochastic optimization algorithms search for a single optimum of a function, the CP method aims to fill a 1-dimensional array of "categories" (a 2-D map for MOLE) that should each contain the best performing sample for each possible value of the parameter under scrutiny (after a discretization). To do so, the algorithm starts by generating a few random samples, evaluates their performance, and stores the best sample of each category; these best samples are then randomly modified to generate new samples, thus having a chance to either improve the best samples of the current categories, or fill new categories in the neighborhood with high-performing samples.

1.6 The first section of this paper exposes a formal description of the CP algorithm as well as a reusable implementation of the method. In the second section we show how the application of the CP algorithm to a real-world geographical simulation model produces novel insights toward the validation of the model.

The Calibration Profile (CP) algorithm

Intuition

2.1 To compute the calibration profiles we propose a new algorithm designed in the framework of evolutionary algorithms. Figure 2 illustrates the progress of this algorithm through the example of the computation of a 9 points profile along x1 of a function f(x1, x2). In this example, the function f computes the evaluation of a 2-parameter model (x1 and x2). The calibration objective is to minimise f(x1, x2). In the step 1 the algorithm randomly samples points (random values of x1 and x2) and computes f(x1, x2). In the step 2, the algorithm divides the definition domain of x1 in 9 disjoint even intervals and keeps only the sampled point with the lowest f(x1, x2) in each of these intervals (this constitutes the elitism stage). The points that have been selected constitute a first approximation of the calibration profile of the model f along x1. In step 3 new samples (x1, x2) are generated by mutating the points in the current approximation of the calibration profile and f(x1, x2) is evaluated for each of these new points. The newly evaluated samples are merged with the existing ones and the algorithm iterates to the step 2. This iteration stops once a given stopping criterion is met after step 2. The projection of the last selected points along x1 constitutes an approximation of the theoretical continuous profile (step 4).
The algorithm is composed of the following steps:

1. **Initial sampling**
   - Sample points in the search space.

2. **Elitism**
   - Update the best solutions per category.

3. **New sampling**
   - Select new samples based on the profile.
   - Check if the stopping criterion is met.

4. **Profile approximation**
   - Compute the profile of each category.
   - Update the profile for each category.

5. **Stopping criterion met?**
   - If yes, stop.
   - If no, go to step 1.

**Objective**

2.1 The CP method takes inspiration from evolutionary algorithms to provide a thorough analysis of the role played by each parameter. Such analysis is independent from one parameter to the other. A function that evaluates the performance of each particular model is supposed to be given (typically such evaluation takes into account the fitting of the model to the whole space $\mathcal{A}$).

2.2 Let $\mathcal{M}$ be the set of possible models under consideration and let $\mathcal{A} = A_1 \times \ldots \times A_n$ be the domain of the parameter vectors that we consider as determining the models, where $n$ is the number of parameters. We denote by $\mu : \mathcal{A} \rightarrow \mathbb{R}$ the map that provides a model for a given parameter vector. Such map can be viewed as a meta-model, that is, a model with free parameters.

2.3 For each category of parameters, we propose a new algorithm that converges directly to the whole set $\mathcal{A}$, and that such intervals form a partition of $\mathcal{A}$.

2.4 Let $f : \mathcal{M} \rightarrow \mathbb{R}$ be the function requiring an evaluation for each model. We can obviously evaluate a parameter vector with the function $\hat{f} = f \circ \mu : \mathcal{A} \rightarrow \mathbb{R}_+$ in the context when we use the word solution to just mean a parameter vector $\hat{a} \in \mathcal{A}$. By convention we suppose that best solutions are closer to 0.

2.5 Let $\mathcal{I}_i = C_i \cap \mathbb{A}$ be the set of parameter vectors that we consider as determining the models.

2.6 Our aim is then to compute, for each $k \in \{1, \ldots, n\}$, a good approximation of $h_k(x)$, the best possible performance of the model. In our view, this function is a good indicator of the impact of a parameter and provides valuable information for further analysis.

2.7 In order to explore $\mathcal{A}_k$ uniformly, we use an evolutionary algorithm with diversity pressure mechanism (Goldberg et al. 1992; Toffolo & Benini 2003; Mauw 1997; Mouret & Doncieux 2012). Let $C_1, \ldots, C_n$ be $n$ different semantic categories to which one can associate an element belonging to $\mathcal{A}_k$. In our approach, we consider that categories can be identified with intervals $[a_i^k, \ldots, a_j^k]$, and that such intervals form a partition of $\mathcal{A}_k$.

2.8 In this section we consider $k \in \{1, \ldots, n\}$ to be fixed and we describe the CP algorithm for the analysis of the $k$-th parameter. As mentioned earlier, the goal is to approximate $h_k(x)$, that is to provide sufficiently good approximations of $h_k(a_k)$, for certain $a_i^k, \ldots, a_j^k \in \mathcal{A}_k$. To overcome the computational limits of this approach, we propose a new algorithm that converges directly to the whole set $\{h_k(a_i^k), \ldots, h_k(a_j^k)\}$.

2.9 To estimate the impact of the $k$-th parameter on the model, we consider the function $h_k : \mathcal{A}_k \rightarrow \mathbb{R}_+$ defined by

$$h_k(a_k) = \inf\{f(a_1, \ldots, a_{k-1}, a_k, a_{k+1}, \ldots, a_n) : q \in \mathcal{A}\}.$$

2.10 Let $\mathcal{A}_k = \{a_1, \ldots, a_n\}$ be the domain of the parameter vectors that we consider as determining the model. We denote by $\mu : \mathcal{A}_k \rightarrow \mathbb{R}$ the map defined by $\mu(\hat{a}_k) = \hat{a}_k$, where $\hat{a}_k = (a_1, \ldots, a_{k-1}, a_k, \ldots, a_n)$, with $a_{k-1}$ and $a_{k+1}$ such that $\mu(\hat{a}_k) = (\mu(a_{k-1}), \mu(a_{k+1}))$.

2.11 The idea of the algorithm is to generate successive sets of solutions and to keep only the best solution per category. At each iteration the set of best solutions is updated.

The algorithm is composed of the following steps:
Step 1 (Initialisation)

We first generate a random set $X_0$ of parameter vectors, that is $X_0 = \{\tilde{\mathbf{a}}^1, \ldots, \tilde{\mathbf{a}}^M\}$, with $\tilde{\mathbf{a}}^i = (d_{\mathbf{a}}^i, \ldots, d_{\mathbf{a}}^i)$ and $d_{\mathbf{a}} \in A$.

Step 2 (Elitism)

We can consider, by induction, that $X_n$ has already been constructed at this point.

Let us define the sets $S_i = \{\tilde{\mathbf{a}} \in X_n : \text{Cat}(\tilde{\mathbf{a}}) = C_i\}$ for $i = 1, \ldots, J$ (we may get $S_i = \emptyset$ for some values of $i$). Then we apply elitism, that is we select only the best parameter vector in each category. We define the sets $E_i = \{\tilde{\mathbf{a}} \in S_i : f(\tilde{\mathbf{a}}) < f(\tilde{\mathbf{a}}^5) \vee \tilde{\mathbf{a}} \in S_j \}$ for $i = 1, \ldots, J$ and $\Delta_i = \bigcup_{j=1}^i E_j$. In practice all the sets $E_i$ will be either empty, if $S_i = \emptyset$, or containing only one element (if $|E_i| > 2$ we can randomly choose one element in $E_i$ and consider $E_i$ composed of just that element).

Stopping criterion

Check if a predefined stopping criterion is met (number of iterations, stabilisation of the solutions, etc.). Go to Step 4 if it is, continue to Steps 3 if it is not.

Step 3 (New sampling)

Generate $H'$ from $H$ by randomly modifying the parameter vectors in $H$. We can for instance generate each vector in $H'$ by applying a gaussian variation to a vector in $H$. Define the set $X_{n+1}$ as the union of $H$ and $H'$ and go to Step 2.

Step 4 (Ending)

When the stopping criterion is met, we can output our approximation to the function $h_k(x)$: If $\{\tilde{\mathbf{a}}^1, \ldots, \tilde{\mathbf{a}}^m\}$ is the last $H$ computed, our approximation to $h_k(x)$ is the set of 2-tuples $(\tilde{\mathbf{a}}^i, f(\tilde{\mathbf{a}}^i))$. Let us note that $m \geq k$ by construction, but in practice, when the number of iterations is high enough (see section 2.1 point 1) we get $m = k$.

Validity

As we said in paragraph 2.7, our approach consists in estimating the impact of the $k$-th parameter on the performance of the model, by finding a good approximation to the function $h_k(x)$. Why our method works is because such a good approximation, we make the following remarks:

1. At each iteration we randomly generate new solutions from old ones. Given a solution $\tilde{\mathbf{a}}^i$ such that that $\text{Cat}(\tilde{\mathbf{a}}^i) = C_i$, the probability of getting a new solution $\tilde{\mathbf{a}}^j$ from $\tilde{\mathbf{a}}^i$ such that $\text{Cat}(\tilde{\mathbf{a}}^j) = C_i$ with $j \neq i$ is different from 0 (this is clearly the case if we use, for instance, a gaussian variation). Hence, if enough iterations are repeated, every category will eventually be reached by as many solutions as we wish.

2. Let us consider $u = \inf\{f(\tilde{\mathbf{a}}^i) : \text{Cat}(\tilde{\mathbf{a}}^i) = C_i\} = \inf\{h(x) : \text{Cat}(x) = C_i\}$. If the function is sufficiently regular (is it enough if it is continuous, or if it is just semi-continuous on its local minima), then, by the analogous argument to that in point 1, we will eventually find an element $\tilde{\mathbf{a}}$ such that $\text{Cat}(\tilde{\mathbf{a}}) = C_i$ and $f(\tilde{\mathbf{a}})$ is as close to $u$ as we want.

These remarks, together with the fact that we can consider multiple categories as needed, justify our claim. We should note that, as it is usually the case with evolutionary algorithms, we do not provide a precise speed of convergence for our algorithm; however, experimental results are very satisfactory (see section 3).

Heuristic improvement

In paragraph 2.11 we have presented the most basic version of our algorithm, for the sake of clarity. However in practice we use a variable version, heuristically devised, to improve its performance. In this section we present these modifications. We use the same notation as in paragraph 2.11.

The main point of this modified version is the random generation of new solutions (step 3). In this part we rely on the standard techniques used in genetic algorithms, in which the random variation comes from two biologically-inspired operators: mutation and crossover. These operators are not applied to the whole set of solutions, but rather on a subset $H_{GA}$ which is generated by a certain selection mechanism (we will explain below this selection mechanism). From $H_0$ we generate $H'$ in two steps: We first generate a new set of solutions $H_0$ by applying a SBX-crossover to $H_0$ (see Deb & Agrawal 1994), and then we apply a self-adapting Gaussian mutation to $H_0$ (see Hinterding 1995). The result is our $H'$. In the selection of $H_{GA}$ we follow a standard selection mechanism: Binary tournament selection. It consists in randomly choosing two solutions $x, y \in H$, comparing them via a certain function $g : H \to \mathbb{R}^+$, and then keeping the best ($x$ is kept if $g(x) < g(y)$); we repeat this procedure until we reach the desired size of $H_0$.

Note: Although we have defined $H_0$ as a set (elements in $H_0$ are unique), we can also define $H_0$ as a tuple, allowing a same solution to appear more than once.

One feature that is specific to our approach is the definition of the function $g$ that we call local contribution.

Definition Let $H = \{\tilde{\mathbf{a}}^1, \ldots, \tilde{\mathbf{a}}^m\}$ and $\alpha_1^1 < \ldots < \alpha_m^m$. Given a solution $\tilde{\mathbf{a}}^i$, let $D_i$ denote the area of the triangle in $\mathbb{R}^2$ formed by the 2-tuples $(\tilde{\mathbf{a}}^i, f(\tilde{\mathbf{a}}^i))$, $(\tilde{\mathbf{a}}^j, f(\tilde{\mathbf{a}}^j))$ and $D_i = f(\tilde{\mathbf{a}}^i) - f(\tilde{\mathbf{a}}^j)$. Then we define the local contribution $g(\tilde{\mathbf{a}}^i) = (\mathbf{1}^t \alpha^2)^t$, where if $0^t \alpha^2$ is not extreme, i.e. $\alpha_i^2 < \alpha_j^2$, then we define $g(\tilde{\mathbf{a}}^i) = \alpha_i^2$.

We have just described how to generate a new set of solutions $H'$ from $H$. The heuristics behind this procedure is to give some priority to the generation of new solutions that are either locally good, which accelerates the approximation to the local minima in each category, or on the extremes, which widens the exploration of the space.

In order to describe the version of the algorithm that we use in the experiments, the other point we should mention is the stopping criterion. A natural choice would be based on the stability of the successive $H$ computed: If we detect no important changes from one $H$ to the next one (during several iterations), the algorithm stops. We can measure these changes in different ways, for instance considering the function $\tilde{R}H = \sum_{\tilde{\mathbf{a}} \in H} f(\tilde{\mathbf{a}})$. However, in practice, we consider a maximal
computation time as the stopping criterion and the stability becomes an element of subsequent analysis.

Generic implementation

2.20 In order to make the CP algorithm usable we provide along with this paper a generic implementation in the free and open source library called MGO (Multi Goal Optimisation)\(^\text{[2]}\) programmed in Scala\(^\text{[3]}\). MGO is a flexible library of evolutionary algorithms. It leverages the Scala "trait" system to provide a high level of modularity.

2.21 This section shows how to compute a calibration profile on a toy model with MGO. In this example, the model is the Rastrigin function and the objective is to minimise this function. The objective function in MGO should be implemented through a specialised trait of the "Problem" trait. Listing 1 shows how to implement the Rastrigin function into MGO.

```scala
import fr.iscpif.mgo._
import math._
import ultil.Random

trait Rastrigin <: GAProblem with MGFitness {
  def min = Seq.fill(genomeSize)(-5.12)
  def max = Seq.fill(genomeSize)(5.12)
  type P = Double

  def express(genome: Seq[Double], rng: Random) =
    10 * genome.size + genome.map(x => (x * x) - 10 * math.cos(2 * math.Pi * x)).sum

  def evaluate(phenotype: P, rng: Random) = Seq(phenotype)
}

Listing 1. Implementation of the Rastrigin function in MGO.

2.22 To compute a calibration profile of this Rastrigin function it should be composed with the "Profile" trait that provides the CP algorithm. Listing 2 shows how to achieve this composition. In this example two other traits are used: A termination criterion based on a maximum number of steps ("CounterTermination") and a trait to specify that the profile is computed on a part of the genome ("ProfileGenomePlotter").

```scala
import fr.iscpif.mgo._
import scalax.io._

object TestProfile extends App {
  val profile = new Rastrigin
    with Profile
    with CounterTermination
    with ProfileGenomePlotter {
    def genomeSize: Int = 6
    def lambda: Int = 200
    def steps = 200
    def x: Int = 0
    def nX: Int = 1000
  }

  implicit val rng = newRNG(42)

  val res =
    m.evolve.untilConverged {s => println(s.generation)}
  .population

  val output = Resource.fromFile("/tmp/profile.csv")
  for {
    i <- res.toIndividuals
    x = m.plot(i)
    v = m.aggregate(i.fitness)
    if !v.isPosInfinity
  } output.append(s"$x,$v\n")
}

Listing 2. Computing the profile of the Rastrigin function in MGO.

2.23 Like most evolutionary algorithms, the CP algorithm compute the evaluation function many times. Those numerous evaluations could be computationaly costly. To cope with this, MGO takes advantages of the multi-core parallelism available in modern computers. This kind of parallelism is suitable for many models but can be limiting in case of large stochastic agent based models for instance. To overpass this limit we have integrated the CP algorithm in the OpenMOLE\(^\text{[4]}\) (Open Model Exploration) framework (Reuillon et al. 2013). OpenMOLE enables the distribution of the CP algorithm on large scale distributed computing infrastructures including multi-core computers, computing clusters and even world wide computational grids. In the following section we used this distributed implementation of CP framework to study a real world agent based model.

Application the SimpopLocal geographical model

A model to simulate the emergence of structured urban settlement system

http://jasss.soc.surrey.ac.uk/18/1/12.html

20/10/2015
3.1 SimpopLocal is a stylised model of an agrarian society in the Neolithic period, during the primary “urban transition” manifested by the appearance of the first cities (Schmitt et al. 2014). It is designed to study the emergence of a structured and hierarchical urban settlement system by simulating the growth dynamics of a system of settlements whose development remains hampered by strong environmental constraints that are progressively overcome by successive innovations. This exploratory model seeks to reproduce a particular structure of the Rank-Size distribution of settlements well defined in the literature as a generalised stylised-fact: For any given settlement system throughout time and continents, the distribution of sizes is strongly differentiated, exhibiting a very large number of small settlements and a much smaller number of large settlements (Liu 1996; Durand-Dastes et al. 1998; Berry 1964; Fletcher 1986). This kind of distributions can be modeled by a power-law when small settlements are not considered or a log-normal when small settlements are considered (Robson 1973; Favaro & Pumain 2011). Such distributions are easily simulated by rather simple and non spatial statistical stochastic models (Gibrat 1931; Simon 1955). But for theoretical considerations and to specify the model in a variety of geographical frames, we think it is necessary to make explicit the spatial interaction processes that generate the evolution of city systems. According to the evolutionary theory of system of cities (Pumain 1997, 2009), the growth dynamics of each settlement is controlled by its ability to generate interurban interactions. The multi-agent system modeling framework enables us to include mechanisms, derived from this theory, that govern the interactions between settlements (Bretagnolle et al. 2006; Pumain & Sanders 2013). The application of this concept resulted in several Simpop models (Bura et al. 1996; Bretagnolle & Pumain 2010; Sanders et al. 1997) in which the expected macro-structure of the log-normal distribution of sizes emerges from the differentiated settlement growth dynamics induced by the heterogeneous ability of interurban interactions. Therefore the aim of SimpopLocal is to simulate the hierarchy process via the explicit modelling of a growth distribution that is not entirely stochastic as in the Gibrat model (Gibrat 1931) but that emerges from the spatial interactions between micro-level entities.

Agents, attributes, and mechanisms of the model

3.2 The SimpopLocal model is a multi-agent model developed with the Scala software programming language. The landscape of the simulation space is composed of a hundred of settlements. Each settlement is considered as a fixed agent and is described by three attributes: The location of its permanent habitat, the size of its population, and the available resources in its local environment. The amount of available resources is quantified in units of inhabitants and can be understood as the carrying capacity of the local environment for sustaining a population. This carrying capacity depends on the resources exploitation skills that the local population has acquired from inventing or acquiring innovation. Each new innovation acquired by a settlement develops its exploitation skills. This resource exploitation is done locally and sharing or trade is not explicitly represented in the model. The growth dynamics of a settlement is modelled according to the assumption that its population size is dependent on the amount of available resources in the local environment and is inspired from the Verhulst model (Verhulst 1845) or logistic growth. For this experiment, we assume a continuous general growth trend for population - this may be different in another application of the model. The annual growth factor \(g^{\text{growth}}\) is expressed on an annual basis, thus one iteration or step of the model simulates one year of demographic growth. \(g^{\text{growth}}\) has been empirically estimated to 0.02. The limiting factor of growth \(K\) is the amount of available resource that depends on the number \(M\) of innovations the settlement \(i\) has acquired by the end of the simulation step \(t\). \(K\) is the population of the settlement \(i\) at the simulation step \(t\):

\[
K = K_i = \frac{P_i}{R_{\text{max}}} = P_i \left[1 - \frac{P_i}{R_{\text{max}}}\right]
\]

3.3 The acquisition of a new innovation by a settlement allows it to overcome its previous growth limitation by allowing a more efficient extraction of resources and thus a gain in population size sustainability. With the acquisition of innovations the amount of available resources tends to the maximal carrying capacity \(R_{\text{max}}\) of the simulation environment:

\[
R_{\text{max}} = \left(\frac{1}{n} \right) \sum_{j=1}^{n} R^j_{\text{settlements}}
\]

3.4 The mechanism of this impact follows the Ricardo model of diminishing returns, also a logistic model (Turchin 2003). The \(\text{InnovationImpact}\) parameter represents the impact of the acquisition of an innovation and has a decreasing effect on the amount of available resources \(R^j_{i}\) with the acquisition of innovations:

\[
R^i_{\text{diffusion}} = R^i_{\text{diffusion}} \left[1 + \text{InnovationImpact} \left(1 - \frac{R^i_{\text{diffusion}}}{R_{\text{max}}}\right)\right]
\]

3.5 Acquisition of innovations can occur in two ways, either by the emergence of innovation within a settlement or by its diffusion through the settlement system. In both cases, interaction between people inside a settlement or between settlements is the driving force of the dynamics of the settlement system. It is a probabilistic mechanism, depending on the size of the settlement. Indeed, innovation scales super-linearly: The greater the number of innovations acquired, the bigger the settlement and the higher the probability of innovating (Lene et al. 2009; Diamond 1997). To model the super-linearity of the emergence of innovation within a settlement, we model its probability by a binomial law. If \(P_{\text{creation}}\) is the probability that the interaction between two individuals of a same settlement is fruitful, i.e. leads to the creation of an innovation, and \(K\) the number of possible interactions, then by the binomial law, the probability \(\gamma\) of the emergence of \(m_{\text{creation}}\) innovations can be calculated. It is thus possible to compute the probability \(P(\text{emerge } > 0)\) that at least one innovation appears in a given settlement \(i\). This probability then can be used in a random drawing:

\[
P(\text{emerge } > 0) = 1 - P(0) = 1 - \left(1 - P_{\text{creation}}\right)^K
\]

3.6 If the size of the settlement \(i\) considered is \(R^i\) then the number \(N\) of possible pairwise interactions between individuals of that settlement is:

\[
N = \frac{R^i \cdot (R^i - 1)}{2}
\]

3.7 The diffusion of an innovation between two settlements depends both on the size of populations and the distance between them. If \(P_{\text{diffusion}}\) is the probability that the interaction of two individuals of two different settlements is fruitful, i.e. leads to the transmission of the innovation, and \(K\) the number of possible interactions, then by the Binomial law the probability \(\gamma\) of the diffusion of \(m_{\text{diffusion}}\) innovations can be calculated. It is thus possible to compute the probability \(P(\text{diffuse } > 0)\) that at least one innovation appears in a given settlement \(i\). This probability then can be used in a random drawing:

\[
P(\text{diffuse } > 0) = 1 - (1 - P_{\text{diffusion}})^K
\]

3.8 But in this case, the size \(K\) of the total population interacting is a fraction of the population of the two settlements \(i\) and \(j\) which is decreasing by a factor \(\text{DistanceDecay}\) with the distance \(D_{ij}\) between the settlements, as in the gravity model (Wilson 1971).
3.9 However another mechanism controls this innovation diffusion process. Each innovation can only be diffused during a time lag just after its acquisition by a settlement. This time lag of diffusion is controlled by the parameter InnovationLife. According to this depreciation mechanism, an innovation is considered obsolete after InnovationLife simulated years and cannot be diffused any more.

3.10 SimpopLocal involves six free parameters that cannot be evaluated using empirical values: InnovationLife, InnovationImpact, \( R_{\text{max}} \), \( P_{\text{creation}} \), \( P_{\text{diffusion}} \) and DistanceDecay. Those parameters are constrained by the definition domains shown in Table 1.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition domain</th>
</tr>
</thead>
<tbody>
<tr>
<td>( R_{\text{max}} )</td>
<td>[1,40,000]</td>
</tr>
<tr>
<td>InnovationImpact</td>
<td>[0,2]</td>
</tr>
<tr>
<td>( P_{\text{creation}} )</td>
<td>[0,0.1]</td>
</tr>
<tr>
<td>( P_{\text{diffusion}} )</td>
<td>[0,0.1]</td>
</tr>
<tr>
<td>DistanceDecay</td>
<td>[0,4]</td>
</tr>
</tbody>
</table>

3.11 In order to ensure the replicability of the model, a portable implementation has been designed and the source code of SimpopLocal used in this experiment has been filed in a public repository [8].

Objective function

3.12 As defined in paragraph 2.2, CP requires the definition of an objective function (called \( f \)). This function computes a scalar value that is a measure of the quality of the model dynamics for a given set of parameter values. In case of SimpopLocal, \( f \) has been designed by aggregating three objective functions. Each of these objective functions evaluate a different aspect of the stylised facts we seek to simulate with the model.

3.13 The SimpopLocal model being stochastic, the simulation outputs vary from one simulation to the next for a given parameter setting. Therefore, the evaluation of a single parameter setting on the three objectives must take into account this variability. Each evaluation is therefore computed on the output of 100 simulations. We have shown in Schmitt et al. (2014) that it was a suitable compromise between capturing the variability and not increasing too much the computation duration.

3.14 Contrary to some calibration methods such as ABC (Lenormand et al. 2012; Beaumont 2010) which make direct use of stochasticity, genetic algorithms can be strongly biased by noisy evaluation functions. To compensate the bias of over-evaluated set of parameters (that could occur despite the 100 replications), we periodically re-evaluate them. The newly computed evaluation simply replace the old one. In the following experiments 1 percent of the computing time has been dedicated these re-evaluations in order to avoid keeping mistakenly-well-evaluated set of parameters in the population for too long. This re-evaluation scheme has the advantage of simplicity but may be improved in the future.

3.15 Here is how we test each set of parameters according to the three objective functions (a more detailed explanation of the calibration objectives can be found in Schmitt et al. 2014):

- The objective of distribution quantifies the ability of the model to produce settlement size distributions that fit a log-normal distribution. First we evaluate the outcome of each subset of 100 simulations corresponding to one parameter setting by computing, according to a 2-sample Kolmogorov-Smirnov test, the deviation between the simulated distribution and a theoretical log-normal distribution having the same mean and standard deviation. Two criteria are reported, with value 1 if the test is rejected and 0 otherwise: The likelihood of the distribution (the test returns 0 if \( p \cdot \text{value} > 5\% \)) and the difference between the two distributions (the test returns 0 if \( D \cdot \text{value} < D0 \)). In order to summarise those tests in a single quantified evaluation, we add the results of the two tests on the 100 simulations. The best possible score on this objective is thus 0 (all tests returning 0) and the worst 200 (all tests returning 1). By dividing this score by the worst possible value (200), we get a normalised error objective noted \( o \).
- The objective of population (noted \( q2 \)) quantifies the ability of the model to generate large settlements that have an expected size. The outcome of one simulation is tested by computing the deviation between the size of the largest settlement and the expected value of 10,000 inhabitants: \( \frac{\text{populationOfLargestSettlement} - 10 000}{10 000} \). The evaluation of the parameter setting reports the median of this test on the 100 simulations.
- The objective of simulation duration (noted \( q3 \)) quantifies the ability of the model to generate an expected configuration in a suitable length of time (in simulation steps). The duration of one simulation is tested by computing the deviation between the number of steps of the simulation and the expected value of 4000 simulation steps: \( \frac{\text{simulationSteps} - 4000}{4000} \). The evaluation of the parameter setting reports the median of this test on the 100 simulations.

3.16 Given these three objectives, \( f \) is defined as an aggregated objective value that summarises the calibration error of a set of parameter values. This summarised objective is defined as: \( f = \max (\text{error}, \text{q2}, \text{q3}) \). The lowest \( f \) the better a set of parameter values. This function compels the algorithm to select sets of parameter values that score well on the three objectives all together. It prevents from selecting parameter values that would provides efficiency on solely a subset of the objectives as it would be the case with compensative aggregation such as average or a sum function. Indeed, a good calibration requires low scores on all three objectives, none should be disregarded.

3.17 To analyse the produced results, we have established that only input parameters leading to values of \( f \) of less than 0.1 of error are considered valid. Indeed, the empirical data and theoretical knowledge that led to the definition of the objective function are not precise enough to justify a more thorough analysis of the model. This threshold is largely exceeded for some parameter values, however rendering this threshold of acceptability explicit enables the definition of credible bounds for each of the free parameters of the model. These bounds define a validity domain of each of the parameters. Note that the profile algorithm iteratively refines the computed profiles from high values toward lower ones through an iterative process, therefore the proposed bounds are more restrictive than the exact ones.

3.18 This section exposes results of the CP algorithm on each of the six free parameters of SimpopLocal. Each profile presented in this paper has required the achievement of 100,000 CPU hours time (meaning that it would have required 100,000 hours of computation on a single core of a modern processor). To achieve such a huge computational load we have distributed it across the numerous CPU of the European computation grid EGI (European Grid Infrastructure) using the distributed implementation of CP that we have integrated into OpenMOLE. The effective user time was around 15 days to compute all the profiles achieving a gain of 2200. The OpenMOLE script files used to compute those profiles have been made available [8].

**Results**

4.1 The value of the parameter InnovationLife depicts the duration during which an innovation can diffuse in the system of settlements. Figure 3 shows the application of the CP algorithm this parameter. When InnovationLife is set to 0 the model can not be calibrated. For such a value the innovations are instantly deprecated just after their creation, meaning that no diffusion of innovation is possible. This particular point shows that the diffusion process of the model is mandatory to produce acceptable dynamics.
4.2 Under the value 150 the calibration error is higher than our validity threshold and then we consider that the model cannot be calibrated. From a thematical point of view: The deprecation of the innovations is too fast, thus the innovations cannot percolate in the system leading to unrealistic dynamics.

4.3 Over the value 150 the obsolescence of the innovations is slow enough to have no noticeable effect on the calibration of the model. In particular when InnovationLife is set to 4000, the calibration error is lower than our acceptance threshold. For this particular value, the matching mechanism of innovation obsolescence could be considered as disabled. The maximum duration of a simulation is 4000 steps, thus when InnovationLife is set to 4000, the innovations are never considered deprecated. This profile shows that this mechanism does not constrain much the model and is useless to reproduce the expected dynamics. Seeking for a more parsimonious model we have removed this mechanism. The following of the article is based on the SimpopLocal model without the deprecation mechanism.

$R_{\text{max}}$

4.4 The parameter $R_{\text{max}}$ constrains the maximum size of settlements by limiting the impact of the acquisition of an innovation on the size of a settlement. The calibration profile for this parameter is shown on Figure 4.
4.5 The calibration objective for the size of the biggest settlement has been set to 10,000 inhabitants. When \( R_{\text{max}} \) is set to a value lower than 10,000 it is impossible for the model to reach the 10,000 population objective. Thus, by construction, low values of this parameter can not achieve acceptable calibration errors.

4.6 The profile exposes a narrowed range of very low calibration error between [9500;11500]. Surprisingly the calibration algorithm is not able to find low calibration errors for \( R_{\text{max}} \) above 10,500. It indicates that this mechanism is necessary to produce acceptable dynamics and reaching settlements of a size matching empirical evidences.

5.1 The \textit{InnovationImpact} parameter controls the impact of the acquisition of an innovation on the growth of a settlement. Figure 5 show the calibration profile for this parameter. The section of the profile containing low calibration errors is very narrowed, only 2 solutions with calibration errors under 0.1 have been found. This profile has been computed using the definition domain provided by the modellers [0;2]. From Figure 5 it can be deduced that the definition domain is too wide compared to the validity domain for this parameter. Therefore another profile has been computed for a much more restraint definition domain: [0;2 \times 10^{-2}].
5.2 The zoomed profile is shown on Figure 6. It shows that when the impact on the settlement growth of the innovation is too low the dynamic is too slow to reach the calibration objectives. On the contrary, when InnovationImpact is too high the growth of the settlements is too fast to match credible dynamics (whatever the value of the other parameters can be). This profile shows that the validity domain of this parameter is [0.006;0.01].

6.1 The $P_{creation}$ parameter determines the capacity of a settlement to create a new innovations. Figure 7 shows the calibration profile for this parameter. Once again the definition domain given by the modellers [0;0.01] is too wide to establish directly a validity domain. However, this profile shows that for high values of $P_{creation}$ the model can not reach low calibration errors. The inflexion of the profile near the value 0 indicates that low calibration errors are located in a much more narrowed domain close to 0.

6.2 Figure 8 shows the calibration profile of the $P_{creation}$ parameter within the domain $[0;10^{-5}].$ From this profile it can be deduced that the validity domain of $P_{creation}$ is $[0.4 \times 10^{-6};2 \times 10^{-6}]$. For low values of $P_{creation}$ the model can not be calibrated. The innovations are generated too slowly to engender a sufficient growth (whatever the value of the other parameters can be). For high values of $P_{creation}$ the system races and growth is too fast.
7.1 The parameter $P_{\text{diffusion}}$ governs the exchange of innovation between settlements. Figure 9 shows the calibration profile for this parameter which is very flat and contains only high calibration errors. This profile exposes a shortcoming of the algorithm. If the definition domain is very wide compared to the validity domain, the method might take a lot of computation time to find the parameters settings matching low calibration errors and give the false impression that they don’t exist. In the case of SimpopLocal we know that low calibration errors exists because they were found when computing the other profiles. By deduction from the model mechanisms, low calibration errors should be found in a domain closer to 0.

7.2 The zoomed profile using the definition domain $[0;10^{-5}]$ is shown on Figure 10. At this scale the validity domain is revealed: $[0.3 \times 10^{-6};1.8 \times 10^{-6}]$. 

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http://jasss.soc.surrey.ac.uk/18/1/12.html
20/10/2015
7.3 A very interesting aspect of this profile is that low values of $P_{\text{diffusion}}$ prevent the model from producing acceptable dynamics. Noticeably when $P_{\text{diffusion}} = 0$ it disables entirely the diffusion mechanism of the model. At this particular point the calibration error is unsatisfying, thus this profile shows that the diffusion mechanism is mandatory in order to produce realistic behaviours.

DistanceDecay

8.1 The parameter DistanceDecay acts on the mechanism of diffusion of the innovations. It introduces a decaying effect of the distance between settlements on the probability to diffuse an innovation from one settlement to another. Figure 11 shows the calibration profile for this parameter. High values of this parameter prevent the diffusion. The calibration errors computed with high values of DistanceDecay are almost equal to the calibration errors computed with $P_{\text{diffusion}}$ set to 0 (that also inhibits the diffusion process). This matching calibration error values provide an additional hint that the model and its implementation are coherent.
of interest, then the profile likelihood is the function defined as

\[ L(\theta | y) = \max_\xi L(\xi, \theta | y) \]

If \( \delta \) is the parameter of interest, then the profile likelihood is the function defined as \( L_p(\delta) = \max_\xi L(\xi, \theta | y) \). See Pawitan (2001, p. 64) for a detailed description.

9.2 Conclusion

This paper proposes a new method to explore models. This method, called Calibration Profile, computes 2-dimensional graphs that depict the effect of each parameter on the model dynamics using a calibration objective. It thus provides a global view of the local effect of each parameter on the expected behaviour. This constitutes a new form of sensitivity analysis of parameters on a calibration objective that differs from classical sensitivity analysis methods (Saltelli et al. 2000). Classical sensitivity analysis methods are based on statistical analysis that are either local or global. Local sensitivity analysis provides a measurement of the effect of the model parameters on the outputs in a narrowed portion of the parameter space. Global sensitivity analysis provides few aggregated global indicators of the effect of each parameter in the whole parameter space. The CP method produces a global picture of the local effect of a parameter on the calibration objective all along its definition domain. Furthermore the CP method seeks for extreme behaviours (the best ones) of the model and that's why it is based on an optimisation algorithm. This kind of information is not captured by classical sampling-based sensitivity analysis. This CP method has been shown very useful to better understand the dynamical behaviours of a geographical multi-agent simulation model. It has lead to important novel understandings:

- The mechanism linked to the parameter InnovationLife of the model has been shown useless. This mechanism has thus been removed from the model, making it more parsimonious.
- Except from this mechanism all the other mechanisms (i.e spatial heterogeneity, diffusion of the innovations, creation of the innovations...) have been shown mandatory to produce acceptable dynamics. From a geographical point of view, these results are extremely interesting. They show how the consideration of space and of spatial interaction plays a central role in the simulation of credible settlement system growth dynamics.
- For all the parameters (InnovationLife excepted) a connected validity domain has been proposed, these boundaries are summarised in Table 2. The seemingly very narrow validity domain of several of those parameters (in particular InnovationImpact, pCreation, pDiffusion) is not preoccupying to our senses. On the one hand, this difference between the exploration domain and the validity domain can be directly assigned to the lack of empirical knowledge on the processes of innovation. Indeed the probability of innovation or even the impact of innovation adoption are not easily measurable variables in the real world, especially for the ancient periods that the model focuses on. Unsurprisingly the exploration domains that we defined a priori for these parameters has been found much larger than the actual validity domain. In the framework of the model hypothesis, it seems correct that these values are very small. Hence that seemingly very narrow validity domain of several parameters is not due to a lack of robustness of the model but rather to lack of knowledge and empirical measures that prevent the definition of a relevant observational scale. On the other hand, the value of those parameters is also directly linked to the definition of the concept of innovation implemented in the geographical model and to how we can quantify it (such as the average number of innovation per time period for example). Therefore the impact of each innovation is strongly related to the number of all considered/simulated innovations.

9.2.1 This model exploration using the CP method has shown very instructive on the model behaviour. More methodological development is being achieved to include this method in an extensive modelling framework encompassing several complementary tools for model evaluations. We think that CP can play a significant role in modelling methodologies such as KISS (Keep It Simple Stupid) for which the simplicity of the models prevails. Using CP during the modelling process may constitute a very efficient tool to achieve progressive complexification of a very simple model towards a realistic one.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition</th>
<th>Validity domain</th>
</tr>
</thead>
<tbody>
<tr>
<td>( R_{\text{max}} )</td>
<td>[1;40, 000]</td>
<td>[9, 500;11, 500]</td>
</tr>
<tr>
<td>InnovationImpact</td>
<td>[0;2]</td>
<td>[8×10^{-3}, 1×10^{-2}]</td>
</tr>
<tr>
<td>pCreation</td>
<td>[0;0.1]</td>
<td>[4×10^{-7};2×10^{-9}]</td>
</tr>
<tr>
<td>pDiffusion</td>
<td>[0;0.1]</td>
<td>[3×10^{-7};8×10^{-6}]</td>
</tr>
<tr>
<td>DistanceDecay</td>
<td>[0;4]</td>
<td>[0;2;1.1]</td>
</tr>
</tbody>
</table>

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Notes

1Let \( y \) be the data and \( \theta \) a vector of parameters that can be decomposed as \( \theta = (\delta, \xi) \). The likelihood function can be written as \( L(\theta | y) = L(\delta, \xi | y) \). If \( \delta \) is the parameter of interest, then the profile likelihood is the function defined as \( L_p(\delta) = \max_\xi L(\xi, \theta | y) \). See Pawitan (2001, p. 64) for a detailed description.

2https://github.com/ISCPIF/mgo

3http://scala-lang.org/

4www.openmole.org

5https://github.com/ISCPIF/profiles-simpopLocal


