ORIGINAL PAPER



# **Basin Hopping Networks of continuous global** optimization problems

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**Abstract** Characterization of optimization problems with respect to their solvability is one of the focal points of many research projects in the field of global optimization. Our study contributes to these efforts with the usage of the computational and mathematical tools of network science. Given an optimization problem, a network formed by all the minima found by an optimization method can be constructed. In this paper we use the Basin Hopping method on well-known benchmarking problems and investigate the resulting networks using several measures.

**Keywords** Benchmarking · Network science · Continuous global optimization · Basin Hopping

# **1** Introduction

The task of box-constrained global optimization (GO) is to find the solution to the problem

$$\min_{x \in S} f(x),\tag{1}$$

where  $f : S \subset \mathbb{R}^n \to \mathbb{R}$  is a continuous function and *S* is a box. The vast literature of GO contains several proposed algorithms for solving (1), and it is a question of high interest how these algorithms perform on different problems. To this end, several benchmarking techniques have already been proposed (see, e.g. Dolan and Moré 2002; Mittelmann 2017; Neumaier et al. 2005; Rios and Sahinidis 2013). Our method complements these works with the help of the emerging field of network science

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(Newman 2010). The proposed methodology follows the core idea of the early work of Stillinger and Weber (1984), in which potential energy landscapes of atom clusters were formed into graphs. This is done in a way that the landscapes can be divided into basins of attraction surrounding each locally minimal energy level. This approach was later applied in the analysis of network topology of small Lennard-Jones clusters (Doye 2002). In that paper, the so-called *inherent structure network* (ISN) was built in which vertices correspond to the minima and the edges link those minima which are directly connected by a transition state. The same idea can be used for combinatorial optimization problems (Scala et al. 2001; Tomassini et al. 2008). We give here a possible extension of these ideas to the space of continuous optimization problems<sup>1</sup>, under the assumption that the optimization method used is Basin Hopping (BH). BH is a primary heuristic method which could be considered as the basis of many elaborate heuristic-based global optimization algorithms.

Once the network representation G of a global optimization problem P is constructed, similarly to the above mentioned ISN, many interesting graph metrics and measures of G can be calculated which can shed a light on several detailed characteristics of P. The important questions we aim at answering in this paper are the following:

- What kind of graph representations can be constructed for continuous global optimization problems?
- Practically, how difficult is it to find these graphs?
- From the network science literature, what are the interesting and relevant measures and what are the interpretations of them in the context of continuous global optimization?
- Given the networks and their measures, how can these be meaningfully applied together on (well-known) optimization problems and what are their implications?

In the following we first give an overview of the methodology producing the graph models. Then, we discuss several graph metrics and measures together with their interpretation in the context of continuous global optimization problems. This is followed by numerical experiments in which some benchmark optimization problems from the literature are investigated. Details on the network models of the tested functions are given, which we believe give further contributions to the understanding of why some problems are easy or hard for a particularly efficient optimization scheme called Basin Hopping.

# 2 Methodology

### 2.1 Network representation of optimization problems

Interestingly, an early paper of Locatelli (2005) and the recent book of Locatelli and Schoen (2013) already contain the idea of the (possible) construction of the network representing a continuous global optimization problem. In the following, using the

<sup>&</sup>lt;sup>1</sup> Note that the optimization problem (1) can also be extended to have constraints, although in the experimental part of our paper we will investigate only box-constrained problems of form (1).

terminology from Locatelli and Schoen (2013), we give the necessary definitions of the graph construction.

First of all, we assume that a local search procedure  $\mathscr{L}(\cdot)$  is available which, given a starting point *y* returns a locally optimal solution *z* of *f* characterized by  $||x - z|| \le \varepsilon \implies f(z) \le f(x) \quad (\forall x \in S)$ . We associate a neighborhood structure  $\mathscr{N}(\cdot)$  to each point in the search space *S*: for a given point  $x \in S$ ,  $\mathscr{N}(x)$  contains those points of *S* which we get by perturbation of *x* and subsequently starting a local optimization method from the perturbed point. Practically, the structure  $\mathscr{N}$  depends on the underlying local optimization algorithm used to solve the global optimization problem (1). The *local optima network* G(V, E) can be defined in the following way. First of all, it is assumed that  $\mathscr{L}(x) = x$  if *x* is a local minimizer point of *f*.

- The set V of vertices are the local minimizer points of f:

$$V = \{ y \in S : \exists x \in S, y = \mathscr{L}(x) \}.$$

Note that we need to assume that  $|V| < \infty$ .

The set E of edges is defined as

$$E = \{(x, y) \in V \times V \mid \exists z \in \mathcal{N}(x) : \mathcal{L}(z) = y \text{ and } x \neq y\}.$$

Remark that the elements of set *E* are *directed*. Similarly to Locatelli and Schoen (2013), a *monotonic* graph  $G_m(V, E_m)$  can also be defined with the edge set

 $E_m = \{(x, y) \in V \times V \mid \exists z \in \mathscr{N}(x) : \mathscr{L}(z) = y \text{ and } f(y) \le f(x) \text{ and } x \ne y\}.$ 

We say that a local minimizer y is a *neighbor* of another local minimizer x iff  $(x, y) \in E$ . Note that in  $G_m(V, E_m)$  all nodes with no outgoing arcs are locally optimal solution of (1).

We will also use the concept of the *adjacency matrix* A of a graph G in the later notations, which is defined as

$$A_{ij} = \begin{cases} 1 & \text{if } (i, j) \in E(G), \text{ and} \\ 0 & \text{otherwise.} \end{cases}$$

Finally, we define the *natural* local optima network (NLON). In this representation, two nodes are connected if they are separated by a critical point (i.e. a stationary point where the Hessian has a single negative eigenvalue More and Munson 2002). Separation of two local minima  $x_1$  and  $x_2$  means that starting a gradient descent local search  $\mathcal{L}$  from a point which is given by arbitrarily small perturbation of the critical point can lead to either  $x_1$  or  $x_2$ .

*Illustration* As an illustrative example, NLON of the classical, two dimensional Six Hump Camel Back (SHCB) global optimization problem is shown on Fig. 1. This problem has 6 local optima among which two of them are global optima (shown as larger (blue) nodes). The labels on the nodes represent the two dimensional coordinates of the corresponding local optima. Size of the nodes are proportional to their degree.



Fig. 1 The natural local optima network of the SHCB global optimization problem

### 2.2 Basin Hopping method

The Basin Hopping (BH) method is a metaheuristic, which proved to be very efficient in solving global optimization problems (Leary 2000; Locatelli and Schoen 2013; Wales and Doye 1997). Using the terminology of Locatelli and Schoen (2013) the high level description is given in Algorithm 1. In the following, we refer to the lines of Algorithm 1 to give a detailed description. It is assumed that a uniform pseudorandom generator  $\mathscr{U}(\cdot)$  is provided and the input is a continuous global optimization problem of form (1). In Line 1 a starting point y is generated uniformly at random in the search space S. Using a local search procedure  $\mathscr{L}$  a local minimizer point x is found in Line 2. Line 4 selects a new starting point from the global neighborhood (to be defined later) of x. In order to do so, we let d be an n-dimensional Gaussian(0, 1) random vector with ||d|| = 1 (e.g. d is a random direction), and  $r_2$  be a positive fixed step size. The new starting point z is generated as being  $x + r_2 d$ . In Line 5 a local search is performed starting from z and its result is stored as x (a local minimizer point). Line 7 selects a new starting point z from the local neighborhood of x. This is done by sampling a uniformly random point over  $S \cap B[x', r_1]$ , where B[x, r] is a box centered at x and having half-edge length r > 0. We start a local search from z and its result is stored as y (Line 8). In Line 9 we check whether y is a better solution than x (being 'better' is to be defined later). In Lines 12 and 13 we check whether the local and global stopping criteria are satisfied, respectively. The algorithm returns with the local minimizer point x and the corresponding function value f(x) in Line 14.

The conditional statement in Line 9 requires the procedure IsAcceptable(x, y) to be given. This procedure can be implemented in different ways, the most common approaches are as follows:

Monotonic: the procedure IsAcceptable(x, y) returns whether f(y) < f(x).

1:  $y := \mathscr{U}(S);$ 2:  $x := \mathscr{L}(y);$ 3: repeat 4:  $z := \mathscr{U}(\mathscr{N}_{\varrho}(x));$ 5:  $x := \mathscr{L}(z);$ 6: repeat 7:  $z := \mathscr{U}(\mathscr{N}_{\ell}(x));$ 8:  $y := \mathscr{L}(z);$ 9: if IsAcceptable(x, y) then 10: x := y;11: end if 12: until local stopping rule is not satisfied 13: until global stopping rule is not satisfied 14: return x, f(x)

Generic: the procedure IsAcceptable(x, y) returns whether

 $\mathscr{U}[0,1] \le \exp(-(f(y) - f(x))/T),$ 

where T is a nonnegative parameter (called *temperature* in the literature), which iteratively gets decreased during the execution of Algorithm 1. Note that this version of the algorithm occasionally accepts non-improving local solutions as well.

Furthermore, there are two procedures in Algorithm 1, namely  $\mathcal{N}_g(\cdot)$  and  $\mathcal{N}_\ell(\cdot)$ , which needed to be defined in detail. These procedures correspond to the local search at Level 3 and Level 2, respectively, of the multi level optimization approach of Locatelli (2005). We employ the scheme from Locatelli (2005), where the neighbors of a local minimum  $x_0$  are all the local minima whose basins of attraction have a nonempty intersection with the box  $B[x_0, r] \cap S$ . Here  $B[x_0, r] := [x_0 - r\mathbf{1}, x_0 + r\mathbf{1}]$ , with half-edge length r > 0 and centered at  $x_0$  (and  $\mathbf{1}$  is the vector whose components are all equal to 1). As this definition depends on the parameter r (which appears to be either  $r_1$  or  $r_2$  in Algorithm 1) an adaptive scheme can be used which iteratively updates its value—for full details see Locatelli (2005).

#### 2.3 Building the Basin Hopping Network

In order to build the local optima network for a particular optimization problem we applied an optimization scheme based on the BH method. Using the same terminology as in Sect. 2.2 the high level description is given in Algorithm 2.

In the following, we refer to the lines of Algorithm 2 to give a detailed description. The algorithm starts with an empty graph  $G_w$ , which iteratively gets expanded if new nodes and edges are found. In Line 1 a starting point y is generated uniformly at random in the search space S. The first node x of the graph  $G_w$  is found in Line 2. Line 4 selects a new starting point from the global neighborhood of x using the same technique in Algorithm 1. In Line 5 a local search is performed starting from z and its result x (as a local minimizer point) is added to the set of vertices. Note that it is

### Algorithm 2 Basin Hopping Network builder algorithm

Require: Global optimization problem P 1:  $v := \mathcal{U}(S)$ ; 2:  $x := \mathscr{L}(y); V := \{x\};$ 3: repeat  $z := \mathscr{U}(\mathscr{N}_{\varrho}(x));$ 4:  $x := \mathscr{L}(z); V := V \cup \{x\};$ 5: 6: repeat x' := x;7: 8:  $y := \mathscr{U}(\mathscr{N}_{\ell}(x));$ Q٠  $x := \mathscr{L}(y); V := V \cup \{x\}; E := E \cup (x', x)$ 10: until local stopping rule is not satisfied 11: until global stopping rule is not satisfied 12: return  $G_w(V, E)$ 

possible that the local search finds a solution which has already been found earlier. In a computer implementation using floating-point arithmetic, one needs to apply  $\varepsilon$ tolerance here, e.g. to check if  $||x - \tilde{x}||_2 < \varepsilon$  for any  $\tilde{x} \in V$  and prescribed  $\varepsilon > 0$ . Thus it is not given that the set *V* gets expanded in each iteration. In Line 7 we store the previously found local solution *x* in a temporary variable *x'*. This will be needed to construct new edges of the graph  $G_w$ . Line 8 selects a new starting point *y* from the local neighborhood of *x'*, similarly to Line 7 Algorithm 1. What is done in Line 9 is that we start a new local search from *y*, and its result *x* is added to the set of nodes *V*, as well as the edge (x', x) to the set of edges *E*. In Line 10 and 11 we check whether the local and global stopping criteria are satisfied, respectively.

It is important to note that the output graph of Algorithm 2 is usually an approximation of the natural local optima network of the input problem P. This is due to the fact that finding the natural LON is a computationally intractable task, especially for higher dimensions. Moreover, a computer implementation is based on floating-point numbers, thus checking if a new node is found can only be done with pre-defined and fixed precision only.

The efficiency of Algorithm 2 highly depends on the parameters  $r_1, r_2$ , on the stopping criteria used in Line 10 and 11, and on the local search procedure  $\mathcal{L}$ . The algorithm needs to find *all* local minima, thus it is usually better to let it run for longer time while allowing a larger number of iterations. According to our experiments, this usually leads to an output graph that has all the local minima of the optimization problem but with more edges than the natural LON. This means that, depending on  $\mathcal{L}$ , nodes which are not neighbors of each other in the natural LON get connected by an edge in the Basin Hopping Network. Thus, post-processing is necessary, which needs a slight modification of Algorithm 2 in the following way. When a potentially new edge is added to the graph in Line 9 we count how many times this edge has been found already. In this way, each edge in the resulting graph has a weight. The post-processing procedure then iterates through the list of edges and removes those ones whose weight is below a certain threshold. This threshold is chosen to be the *P*-th percentile calculated by the nearest rank method. In the numerical examples (see Sect. 4) we experimented with different values of P. Note that a similar procedure was proposed in Daolio et al. (2011).



Fig. 2 A Basin Hopping Network of the SHCB global optimization problem

*Illustration* A possible Basin Hopping Network of the two dimensional Six Hump Camel Back function is shown in Fig. 2. Note the differences between Figs. 1 and 2.

# **3** Graph measures

In the following we give a list of relevant graph measures, taken from network science literature, together with their interpretations in the context of LONs.

Size of the network This measure is defined as the number of nodes, i.e. |V|. Clearly, this represents the number of local minima. As it has been argued, e.g., in Locatelli (2005) a higher number of minima does *not* imply that the problem at hand is more difficult to solve.

*Neighborhood of a node* Besides the size of the network, this is also a critical feature to be found by Algorithm 2, as these two provide the basis for the following measures which are to capture the structural characteristics of the corresponding network. Put it differently, if Algorithm 2 is not able to find the correct network representation of the investigated global optimization problem P, then the measures listed in this section can lead to incorrect claims on P. The neighborhood set of node  $i \in V$  in graph G(V, E) is denoted by  $N_i(G)$ .

*Path and shortest path* These are important definitions for further measures. The series of nodes  $x = x_0, x_1, \ldots, x_k = y$ , where  $x_i$  is adjacent to  $x_{i+1}$ , is called a *walk* between the nodes x and y. If  $x_i \neq x_j$  ( $\forall i, j$ ), then it is called a *path*. The *path length* is k. Given all paths between nodes x and y, a *shortest path* is a path with fewest edges. Shortest paths are usually not unique between two nodes. Note that most of the

heuristic based global optimization methods basically do random walks on paths in a specific underlying graph. If the method is of monotonic type (like Monotonic Basin Hopping Wales and Doye 1997 or Differential evolution Storn and Price 1997) then it walks on  $G_m$ . Some methods, like Simulated Annealing (Kirkpatrick et al. 1983), allow steps towards non-improving solutions, thus they walk on graph G.

Average path length This is defined as the average value of all shortest paths in the network, denoted by  $\ell$ . Networks with low average path length are called *small worlds*. More specifically, in small world networks the average path length grows proportionally to  $\log(|V|)$ . Intuitively, the small world property is a desirable feature in graphs corresponding to global optimization problems.

*Diameter* The size of the longest of all shortest paths is called diameter, and it is denoted by D. This gives a worst-case scenario regarding the number of jumps that have to be taken to reach the global optimum. Similar to the average path length, the smaller the diameter is, the better it is.

*Clustering coefficient* It measures the average probability that two neighbors of a node are themselves neighbors of each other. Formally, the *local clustering coefficient* of node i is

$$C_i = \frac{|\{(x, y) \in E : x, y \in N_i\}|}{k_i(k_i - 1)},$$

where  $k_i = |N_i|$ . The definition of global clustering coefficient is based on triplets. A *triplet* consists of three nodes that are connected by either two (open triplet) or three (closed triplet) undirected ties. The *global clustering coefficient* C is the number of closed triplets over the total number of triplets (both open and closed).

Note that small world networks tend to have high clustering coefficient. Intuitively, networks with high C value correspond to easier to solve global optimization problems.

*Node degree* The neighborhood structure  $\mathscr{N}$  can be quantified. This gives the definition of node degree, which is the number of edges adjacent to a node. In our case, this measures the number of adjacent local optima. Since our graphs are directed, we have indegree and outdegree for a given node. Formally, the *outdegree* is a function  $d^+: V \to \mathbb{N}_0$  which for a node x gives  $d^+(x) = |\{y \in V : (x, y) \in E\}|$ . The *indegree* is defined as  $d^-(x) = |\{y \in V : (y, x) \in E\}|$ . Nodes with degree that greatly exceeds the average degree in the graph are called *hubs*. It is known that high degree nodes are easier to be found by random walks (Newman 2010). Hence, if the global optimum vertex is a hub, then a heuristic method can perform well on the problem.

Average degree This measure is the ratio  $\frac{1}{|V|} \sum_{x \in V} d(x)$ , where d(x) is either the indegree and outdegree (the average is the same value in both cases); and it is denoted by  $\langle k \rangle$ .

*Degree distribution* This measure is defined as the probability distribution of all degrees in the graph. Formally,  $p_k$  is the fraction of nodes with degree k:

$$p_k = \frac{|\{x \in V : d(x) = k|}{|V|},$$

where d(x) can be indegree or outdegree, or the sum of the two (i.e. the graph is made undirected). Degree distributions have two categories of particular interest: (i) *random networks* [also called Erdős–Rényi graphs (Erdős and Rényi 1959)] have binomial distribution of degree k:

$$p_k = {\binom{|V|-1}{k}} p^k (1-p)^{|V|-1-k},$$

where *p* is the probability that two nodes are connected; and (ii) *scale-free networks* (Albert and Barabási 2002), which follow a power law distribution of the form  $p_k \sim k^{-\alpha}$ , where  $\alpha$  is a parameter typically in the range  $2 < \alpha < 3$ .

The degree distribution is an important global measure of a network. Both random and scale-free networks have advantages and disadvantages. These networks tend to have small clustering coefficients and short average path length. By definition, scalefree networks contain a few hubs with high degree and lots of nodes with low degree. In contrast, random networks contain very similar nodes.

*Community structure* It can be informally defined as a partition of vertices into groups in such a way that nodes are more connected within a group and sparsely connected between different groups (Radicchi et al. 2004). Let H be a subgraph of G including node i. If the graph is directed, then define

$$k_i^{in}(H) := N_i(H), \text{ and } k_i^{out}(H) = N_i(G) \setminus N_i(H).$$

Moreover,  $k_i(H) := k_i^{in}(H) + k_i^{out}(H)$ . Now, one can define a subgraph H as a community in a *strong sense*, which is the case when  $k_i^{in}(H) > k_i^{out}(H)$  holds  $\forall i \in V(H)$ ; and also in a *weak sense*, when  $\sum_{i \in H} k_i^{in}(H) > \sum_{i \in H} k_i^{out}(H)$ . The number of communities we find in a network is denoted by K. Note that most of the community detection algorithms treat the graph as undirected. A high number of communities in G does not necessary imply a hard-to-solve optimization problem. However, if the problem is multimodal and the local minima are located in different communities then the Monotonic Basin Hopping method can have difficulties to find the global minimum.

*Modularity* This quantity, denoted by Q, measures the fraction of the edges in the network that connect vertices of the same type (i.e., within-community edges) minus the expected value of the same quantity in a network with the same community divisions but random connections between the vertices (Newman 2006). Formally,

$$Q=\sum_i \left(e_{ii}-a_i^2\right),\,$$

where  $e_{ij}$  is the fraction of edges with one end vertices in community *i* and the other in community *j*, and  $a_i$  is the fraction of ends of edges that are attached to vertices in community *i*. Modularity intends to measure the strength of the community structure in a graph.

Betweenness centrality This measure gives a local score to vertices by measuring the extent to which a vertex lies on paths between other vertices (Freeman 1977). Mathematically, let  $n_{st}^i$  be the number of shortest paths from *s* to *t* that pass through *i*, and define  $g_{st}$  as the total number of shortest paths from *s* to *t*. Then the betweenness centrality (BC) of vertex *i* is  $\sum_{st} \frac{n_{st}^i}{g_{st}}$ . BC is usually calculated on undirected graphs. Since a global optimization method does not necessarily take shortest paths on *G*, a variant called Random Walk BC will instead be investigated in Sect. 4.

*PageRank* This local measure is used on directed graphs, where the score of a vertex is derived from the scores of its network neighbors and it is proportional to their centrality divided by their out-degree. Formally, we need to calculate the vector  $D(D - \alpha A)^{-1}\mathbf{1}$ , where A is the adjacency matrix of the graph  $G_m$ , D is a diagonal matrix with elements  $D_{ii} = \max\{d^+(i), 1\}, \mathbf{1}$  is again the vector whose components are all equal to 1 and  $\alpha$  is a damping parameter (default  $\alpha = 0.85$ ). PageRank was originally designed as an algorithm to rank web pages (Brin and Page 1998) and essentially the score it gives to a page reflects the chance that the random surfer will land on that page by clicking on a link. In the context of global optimization, higher PageRank score means higher chance to be found by the Monotonic Basin Hopping algorithm, which performs random walks on the directed network representing the optimization problem to be solved.

# **4** Numerical results

In this section we demonstrate the usage and implications of the analysis of the Basin Hopping Networks of global optimization problems. For this purpose, two well-known benchmarking problems have been selected from the literature which we discuss in Sects. 4.1 and 4.2 in full details. Further test functions are also analyzed in Sect. 4.3. We are interested to see if the global and local measures listed in Sect. 3 are able to characterize the solvability of the problems.

The implementation of Algorithm 2 was done in AMPL (Fourer and Kernighan 2002), which allows to use a very general class of objective functions and a large selection of local optimizer methods. In our tests we used MINOS (Murtagh and Saunders 2003) as local optimizer  $\mathcal{L}$ . The parameters were:

- the local stopping rule (in Line 10) was: 10,000 iterations;
- the global stopping rule (in Line 11) was: 50 iterations;
- the parameter  $\gamma$  (see Locatelli (2005) for details) was set to 0.5;

- and the values of P in the post-processing were starting from 20 up to 70 with increment 5.

In order to compute the measures listed in Sect. 3, we used the igraph package in R and the NetworkX package in Python. Modularity Q and number of communities K were calculated with the method called Multi Level (Blondel et al. 2008), which is based on local optimization of the modularity measure around a node.

As we have already discussed in Sect. 2.3, the output of the implemented procedure for a given global optimization problem is a set of graphs. These graphs are then used for two types of analysis.

- First, we need to select one of them, which gives the BHN representation of the problem. The selection of this graph is done in the following way. It is assumed that the global optimization problem is continuous, hence the BHN representation must be a connected graph. Furthermore, as a general rule, we select that connected graph which corresponds to a *P* value at which the diameter of the graph gets increased in case of choosing a larger *P* value. This is motivated by aiming at getting such BHN which is close to the natural LON of the problem. If the diameter of the graph gets increased then it is an implication that we just removed a significant amount of edges than before. On the other hand, if the diameter does not change by removing edges, that means we have removed edges from the short ones from all shortest paths (i.e. we have removed unrealistic huge jumps between nodes which are far away from each other in the natural LON).

The graph which represents the optimization problem can then be analyzed using the measures from Sect. 3.

- Secondly, the series of graphs can be considered as results of a certain edge-deleting procedure. This way the *robustness* of the graphs can be measured with respect to a particular metric called *random walk betweenness centrality* (RWBC) (Newman 2005). RWBC is a local measure, a particular variation of the betweenness centrality (see Sect. 3). It is based on random walks, counting how often a node is traversed by a random walk between two other nodes. Calculation of RWBC values are done on the vertices of graph *G* using the edge weights obtained by executing Algorithm 2, i.e., where we count how many times this edge has been found already. In particular, we essentially associate a relative quantity to the node corresponding to the global optimum and thus it can be seen and compared how it relates to the other nodes' RWBC values.

#### 4.1 Griewank function

The first test function we study is proposed by Griewank (1981) and it has the form

$$Griewank_n(x) = \sum_{i=1}^n \frac{x_i^2}{4000} - \prod_{i=1}^n \cos\left(\frac{x_i}{\sqrt{i}}\right) + 1.$$

Usually the search space used in the literature is  $x_i \in [-600, 600]$ , (i = 1, ..., n). However, as this function has a huge amount of local minima we restrict the search

Graph	Size	$\langle k \rangle$	l	D	С	Q	K
G (n = 2, P = 30)	123	7.4796	4.7419	12	0.4810	0.6152	7
$G_m \ (n=2, P=30)$	123	3.7642	3.7609	11	0.4629	0.6179	7
G(n = 3, P = 45)	1359	6.8286	8.7206	20	0.1551	0.7019	12
$G_m \ (n = 3, P = 55)$	1359	2.8182	5.9961	17	0.0330	0.7180	13

Table 1 Network properties of Griewank graphs

Fig. 3 A BSN of Griewank<sub>2</sub> function. *Colors* represent community structure, size of a node corresponds to its PageRank value (color figure online)



space to a much smaller one:  $x \in [-28, 28]^n$ . This restriction results in a smaller network, whose size can be justified by the literature (Cho et al. 2008).

The *Griewank<sub>n</sub>* function, independently from its dimension *n*, has exactly one global minimizer point with value 0, located at the origin. Although the number of its local minima is growing exponentially with *n*, the locations of these minima follow a regular pattern. This makes the corresponding network of simple form. Namely, in n = 2 it is a regular lattice, whose structure remains the same in higher dimensions as well.

*Graph measures* The summary of the graph measures are listed in Table 1. Note that the sizes of the networks reported here are in accordance with the (estimated) number of local optima reported in Cho et al. (2008) if the search space is restricted to  $[-28, 28]^n$ . We chose to study this test function first, mainly because of its regular structure, which is well illustrated on Fig. 3. As we can see, almost all the nodes (apart from those at the edge) have the same degree, so this graph is a typical example of the Erdős–Rényi random networks (see Sect. 3).

It can be immediately noticed that the BHNs have relatively large diameters. This indicates that an optimization method needs to take a large number of iteration steps to guarantee success. This fact is already known from the literature, see, e.g. Locatelli (2003). It is worth mentioning here that although these graphs have large modularity values, which implicates the presence of communities in the network, their nodes



Fig. 4 Degree investigation of Griewank networks; the points are jittered for better visibility.  $\mathbf{a} \ n = 2$ ,  $\mathbf{b} \ n = 3$ 

are very similar to each other with respect to their degree. Thus high Q values are misleading in these cases. We can also notice that the clustering coefficient C is much smaller for n = 3 than for n = 2, which should also be treated with care. In fact the simple reason for this is the BSN we found for n = 3 is incomplete compared to the natural LON representation. As we have already discussed, finding the natural LON representation of an optimization problem is practically impossible in general. Still, it can be constructed easily for the Griewank problem given its regular structure.

Concluding the analysis with the graph measures we can say that they do not give us any particular insights about the Griewank test problems.

*Degree investigation* For investigating the degree distribution of the BHNs we propose the usage of a scatter plot on which the degree of the vertex of the undirected graph and the in-degree of the same vertex of the directed graph can be compared. This kind of visualization gives a very interesting landscape of the problem's local optima. Figure 4 shows the corresponding plots for the Griewank test function. By definition, no points can be above the red line. Note that in both cases the point representing the global optimum (which must be on the red line) is at the top right corner of the figure and the other points are beneath. This implies that the Monotonic Basin Hopping method has a much better chance to find the global optimizer point than the Generic BH method in which steps towards non-improving solutions are allowed.

*Robustness of BHNs* Using the graph sequences we obtained from Algorithm 2 we calculated the random walk betweenness centrality (RWBC) values. The results of these experiments are shown on Fig. 5. Note that a higher P value means a sparser graph, thus higher P values correspond to such runnings of the Basin Hopping method where the number of iterations are relatively small (compared to those represented by lower P values). For both cases the RWBC value of the global optimum is higher than the nodes' average RWBC value. We can also see that for many P values the global optimum vertex has the highest RWBC value, especially for low P values. Clearly,



Fig. 5 Random walk betweenness centralities of Griewank networks.  $\mathbf{a} n = 2$ ,  $\mathbf{b} n = 3$ 



Fig. 6 PageRank values of Griewank networks. Note that the global optimum vertex has the highest PageRank score.  $\mathbf{a} n = 2$ ,  $\mathbf{b} n = 3$ 

nodes with high RWBC values are easier to be found by random walks. Thus, we can conclude that finding the global optimum by Basin Hopping using the general approach is not hopeless, it is only a matter of allowing large numbers of iterations. On the other hand, it is also indicated by these figures that the RBWS values do not really change for lower P values, thus, by only letting the BH search run for a longer time does not guarantee success in global optimization.

Turning now our attention to the *monotonic* network representations, we have already seen in Fig. 3 that due to the special structure of the Griewank functions the global optimum node has the highest PageRank score. Figure 6 shows the calculated values for the different P levels together with the mean PageRank scores. Note that the PageRank value of the global optimum is the highest, hence there are overlaps on the figures. It is clearly advised that using the BH method for solving the Griewank problems should be done using the Monotonic approach.

#### 4.2 Schwefel

Another test problem we study is the Schwefel function which is defined as follows:

N	C.	(1)	0	D	0		
Name	Size	$\langle K \rangle$	ł	D	C	Q	K
G (n = 2, P = 50)	64	14.9688	2.0761	4	0.5712	0.3679	4
$G_m \ (n=2, P=45)$	64	7.8281	2.0447	5	0.5478	0.4039	4
G (n = 3, P = 30)	502	17.4522	3.4073	7	0.3877	0.5345	6
$G_m \ (n = 3, P = 40)$	492	7.849593	3.6651	10	0.3655	0.5501	7

 Table 2
 Network properties of Schwefel graphs (directed graph)

Schwefel<sub>n</sub>(x) = 
$$\sum_{i=1}^{n} -x_i \sin(\sqrt{|x_i|})$$
  $x_i \in [-500, 500].$ 

This problem differs from the previous one in a sense that it has exponentially growing number of local minimizer points whose values are very close to the global optimum and, more importantly, they are located at different regions of the search domain. Thus, this function is considered as a hard problem instance for global optimization methods.

*Graph measures* The properties of the BHNs we found for the Schwefel problems are listed in Table 2. Comparing the different quantities to the ones we obtained for the Griewank functions, we can immediately see the differences everywhere. First of all, the Schwefel networks have very small diameter as well as small average path lengths. This means that the BH method can discover the entire network in reasonable time. However, it must be emphasized that this is true for the BH using the General approach. The modularity values are not that high compared to those of the Griewank networks. Still, the community structure is clearly there in these Schwefel networks, as it is even shown on Fig. 7. Note that the vertices representing the local optima are moved to the periphery for better visibility. We can see here a very interesting fact, namely that 3 out of 4 local optimizer points are in different communities. This is certainly an indication that the Schwefel functions are difficult problems for global optimization methods. In particular, applying the Monotonic approach for BH search is not advised in this case.

*Degree investigation* Figure 8 shows the degree investigation of the Schwefel problems. In order to understand what makes this problem difficult to be solved (at least for BH) we note that the point representing the global optimum is always the one which has the lowest degree, i.e., it is the bottom left point on the red line, indicated by a label 'GO'. In particular, for n = 3, where the number of local optimum is 8, there are many vertices having larger degree than that of the global optimum vertex and hence they are having higher probabilities to be found by random walk. Hence, this is another evidence for indicating the usefulness of applying the Generic BH approach for the Schwefel problems.

*Robustness of BHNs* Finally, we have calculated the RWBC and PageRank scores for the series of Schwefel networks. Figure 9 shows the undirected case, thus it corresponds



Fig. 8 Degree investigation of Schwefel networks; points are jittered for better visibility.  $\mathbf{a} n = 2$ ,  $\mathbf{b} n = 3$ 



Fig. 9 Random walk betweenness centralities of Schwefel networks; *black lines* with *square* markers represent local optima.  $\mathbf{a} n = 2$ ,  $\mathbf{b} n = 3$ 



Fig. 10 PageRank values of Schwefel networks; *black lines* with *square* markers represent local optima. Note the different scales on the *y*-axes.  $\mathbf{a} n = 2$ ,  $\mathbf{b} n = 3$ 

<b>Table 3</b> Network properties of additional test functions								
Name	Size	$\langle k \rangle$	l	D	С	Q		
Levy8 $(n = 2, P = 40)$	47	13.0426	1.9172	4	0.5917	0.2035		
Levy8 <sub><i>m</i></sub> $(n = 2, P = 70)$	45	3.8222	1.8422	4	0.4386	0.3217		
Levy8 ( $n = 3, P = 35$ )	97	9.4124	2.4099	5	0.4728	0.2612		
Levy8 <sub><i>m</i></sub> $(n = 3, P = 50)$	78	4.3333	2.1189	5	0.4353	0.3928		
Ackely $(n = 2, P = 30)$	111	14.5225	2.4985	6	0.5988	0.2103		
$Ackley_m \ (n = 2, P = 20)$	109	7.1927	2.3597	7	0.5766	0.3638		
Ackley $(n = 3, P = 30)$	358	13.9469	3.0894	7	0.4427	0.2452		
$Ackley_m \ (n = 3, P = 30)$	356	7.5365	2.7845	9	0.3928	0.4361		
Rastrigin ( $n = 2, P = 20$ )	118	21.6102	2.2024	6	0.5933	0.1704		
Rastrigin <sub>m</sub> $(n = 2, P = 30)$	116	10.0086	2.0473	6	0.5394	0.2714		
Rastrigin ( $n = 3, P = 65$ )	335	13.2298	2.8954	8	0.3728	0.2548		
Rastrigin <sub>m</sub> $(n = 3, P = 60)$	351	9.3988	3.0543	14	0.3717	0.2905		
Sinusoidal ( $n = 2, P = 25$ )	178	22.6348	2.3764	6	0.5455	0.2010		
Sinusoidal <sub>m</sub> $(n = 2, P = 25)$	167	10.3353	2.5047	6	0.4918	0.3723		
Sinusoidal ( $n = 3, P = 65$ )	912	12.2983	3.9024	12	0.3365	0.3557		

7.7833

3.1646

10

946

т

Sinusoidal<sub>m</sub> (n = 3, P = 60)

to the Generic Basin Hopping. We can immediately see that in these cases the global optimum vertex has lower value that those representing the local minima. Moreover, the node having the maximum RWBC score is a different one. For small P values (representing longer runs of the optimizer method) and n = 3, interestingly, the differences between the GO and the local minima are vanishing. However, this is not the case for n = 2. Though this does not imply that finding the global optimum of the Schwefel function is easier for higher dimension, it only indicates that for higher dimension the probabilities of finding any local minima (including the global one) are roughly equal. Hence, the advice here is to use the Generic Basin Hopping, which can more easily escape from local minimizer points compared to the Monotonic approach.

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0.4276

0.2892



Fig. 11 Degree investigation of Levy8 networks; the points are jittered for better visibility. **a** n = 2, **b** n = 3



Fig. 12 Degree investigation of Ackley networks; the points are jittered for better visibility.  $\mathbf{a} n = 2$ ,  $\mathbf{b} n = 3$ 

Regarding PageRank values on the directed networks, we obtain a completely different result, see Fig. 10. In this case we include networks for higher P values, which represent shorter BH runs. Although all the local optima have higher score than the average, the global optimum node ranks lower than the other optima. For large P values all of them are below the maximum score. When the P value is low, i.e., when the BH algorithm is allowed to take larger amount of iterations, the global optimum vertex has the highest PageRank score. The reason for this is very simple: being stuck in a local optimum by the Monotonic Basin Hopping, the only vertex to which we can jump is the global optimum node. Due to the recursive definition of PageRank, the global optimum node becomes the vertex of highest rank. Note that this happens when letting the MBH algorithm run for exceptionally long time.



Fig. 13 Degree investigation of Rastrigin networks; the points are jittered for better visibility. **a** n = 2, **b** n = 3

### 4.3 Further test functions

In this section we show the analysis of further global optimization test functions. These functions are also extensively used as benchmarks in the GO literature, hence we do not give here the full definitions, only the references: Ackley (2012), Levy8 (Levy et al. 1981), Rastrigin (Torn and Zilinskas 1989), and Sinusoidal (Zabinsky and Smith 1992). As for the Griewank and Schwefel problems, the 2 and 3 dimensional versions of these additional functions were investigated. The results of the network measures are shown in Table 3.

We start with the discussion on Levy8. These functions have the smallest number of local minima, the smallest average path length and diameter, large clustering coefficients and the smallest number of communities. The degree investigation of Levy8 graphs are shown on Fig. 11. For n = 2 the global optimizer node has the highest indegree in  $G_m$  and there is only one node which has higher indegree in G. Similar trend can be noticed for n = 3. We conclude that the Levy8 functions are the most simple ones for MBH. These indicators are in lines with the experiments done in Locatelli et al. (2014) using MBH.

The Ackely and Rastrigin problems are similar to the already analyzed Griewank problem with respect to their landscape, their corresponding BH networks show rather regular grid structure. On the other hand, as we can see from the graph measures, the Ackely and Rastrigin functions have less number of nodes, larger average degree, smaller average path length and diameter compared to Griewank. The degree investigation figures of Ackely functions (see Fig. 12) are similar to Griewank in the sense that there are only a few nodes which have higher degree than the global minimizer. In line with the experiments done in Locatelli et al. (2014) using MBH, Rastrigin functions are slightly more difficult to solve, which can also be demonstrated by the degree investigation, see Fig. 13. We conclude that these test problems can be solved easier than the Griewank problem.



Fig. 14 Degree investigation of Sinusoidal networks; the points are jittered for better visibility.  $\mathbf{a} \ n = 2$ ,  $\mathbf{b} \ n = 3$ 

Finally, the Sinusoidal test problem has the largest number of nodes. This simple fact does not make it difficult to solve. As it can be seen in Fig. 14, especially for n = 3, the global minimizer node has the highest degree.

# **5** Conclusions

Basin Hopping Networks are interesting representations of global optimization problems. Using the rich set of measures and metrics from network science lots of properties can be analyzed regarding the solvability of continuous problems by the fundamental heuristic method Basin Hopping. In this paper we have investigated some well-known benchmark problems, hence our contribution here can be regarded as '*telling classical optimization stories in the language of network science*'. It needs to be emphasized that we did not want to solve the optimization problems but to analyze their structural properties. Hence, we proposed and successfully applied a graph building scheme which, in order to discover how the heuristic BH method performs its search, results in a series of (weighted) networks representing possible outcomes of BH run with different parameter setups.

As future works we can outline two main directions. Based on the results shown in this paper, it is worth dealing with the development of an extension of the Basin Hopping method. That version would work as follows. During its run the algorithm would build up the BHN representation of the global optimization problem. Using that network it would adaptively change its parameters (local stopping rule, direction of search, length of the jumps, acceptance criterion, etc) according to the characteristics of the BHN. For example, if it detects strong community structure in the network then the algorithm should make bigger jumps in the search space to discover further details. This and further techniques might result in a Basin Hopping approach which, although for a price of larger computational cost, would give higher level of guarantee that the best solution found is the real global minimum. This has particular relevance in case of multimodal optimization.

Another line of research is to discover such network representations of global optimization problems which correspond to other optimization methods. Although many heuristic methods share similarities to BH, it would be interesting to see and compare the different graphs and develop benchmarking methodologies based on network science.

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