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基於子空間映射與多臂吃角子老虎機技術之 實數最佳化

Real-valued Optimization by Subspace Projection and Multi-armed Bandit Techniques

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摘要

本論文提出了一新的實數多模態 (multimodal) 最佳化方法。大多 數的現實問題都可以由多個單峰問題合成。而對於實數最佳化,我 們較感興趣的是多模態且可被拆解成多個階層化的單峰問題。然而, 一旦面對多模態問題,我們就必須面對利用性 (exploitation) 與探索性 (exploration) 的問題。本論文提出一解決多模態問題的技巧。此技巧 結合了階層式聚合 (hierarchically clustering) 與最小描述長度 (minimum description length),來辨識搜尋空間中可能的單峰模型。接著,我們使 用一加一演化策略 ((1+1)-Evolutionary Strategy) 來最佳化一齊次座標 (homogeneous coordinate) 上的線性轉換矩陣,並利用此矩陣將原搜尋 空間投影到另一有良好邊界定義的子空間。此投影嘗試定義出只包含 一單峰之感興趣區域 (region of interest)。最後,我們提出一新的多臂吃 角子老虎 (multi-armed bandit) 技術,來分配給各個子空間的資源,以 最大化獲得全域最佳解的機率,我們將此結果與自適應共變異數矩陣 演化策略 (Covariance Matrix Adaptation Evolution Strategies, CMA-ES)、 標準粒子群演算法 (Standard Particle Swarm Optimization 2011)、與實數 域蟻群演算法 (Ant Colony Optimization for Continuous Domain, ACO_R) 結合,並利用 CEC 2005 實數最佳化測試問題來比較結果。

Abstract

This thesis presents a new technique for real-valued multimodal optimization. Most of the real-world problems can be described as a composition of uni-modal subproblems. For real-valued optimization, we are interested in problems that are composed of multiple hierarchically decomposable unimodals. However, allocating resources to exploit different uni-modals leads to the common delimma between exploration and exploitation. This thesis proposed some new techniques that aim to solve multimodal problems more efficiently. A new technique combining hierarchical clustering and minimum description length (MDL) is proposed to help identify potential uni-modals in the search space. Then, a linear transform matrix in homogeneous coordinate, optimized by the (1+1)-Evolutionary Strategy, projects the search space to a subspace with well-defined boundary. This projection aims to define a region of interest (ROI) that isolates a uni-modal subproblem. Finally, a new multi-armed bandit technique, aiming to maximize the probability of aquiring the global optimium, is proposed to allocate resources for each uni-modal. We combined our new techniques with Covariance Matrix Adaptation Evolution Strategy (CMA-ES), Standard Particle Swarm Optimization (SPSO) 2011, and Ant Colony Optimization for Continuous Domain (ACO_R). The results are evaluated with the CEC2005 Special Session on Real-Parameter Optimization benchmark problems.

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Chapter 1

Introduction

Multimodal problems that contain more than one region or more than one local optimum are common in the real world. *Rastrigin function*, a hierachical problem which is composed of a larger hill and multiple smaller hills are also well-known in real-valued optimization. In order to tackle these multimodal problems with a limited amount of resources, one needs to decide how much to invest for searching a better hill, while still maintain enough of exploitation on each potential hill.

The difficult part is that searching for a new hill and exploiting the best hill often require opposite searching behaviors. For a fixed amount of total evaluations, spending more time searching on one hill implies less attention on exploring other possible unimodals in the search space. This is the common dilemma between *exploration* and *exploitation* for real-valued optimization algorithms. Also, during different phases of optimization, the ratio between exploring new hills and exploiting the current best hill should be altered. Exploration is more important in the beginning, while exploitation is more required in the end. What makes this problem even harder is that in order to decrease the number of total evaluations, the size of the population are often limited in the real-valed optimization algorithms. This means that as the number of hills to invest increases, the number of particles on each hill decreases. This weakens the exploitation ability on each hill, making it less likely to identify new potential regions on each hill.

We are more interested in hierarchically decomposable problems. With a certain amount of hints, we should be able to focus on one of the many potential hills, and identify when

to invest more on certain hills to iteratively discover more smaller hills. Also, we would like to only focus on some regions instead of the whole search spaces, in order to gain performance. This requires detection of promising areas, and search space splitting techniques. Moreover, after identifying certain interesting regions, we also need to decide how to allocate our resources. Given the abilities described above, we can pay more attention on the promising regions and iteratively break down a difficult multimodal problem into smaller and easier unimodal problems.

1.1 Thesis Objectives

We propose some techniques to break down multimodal problems into several unimodal problems in smaller non-overlapping subspaces. We believe that solving one unimodal problem on a smaller subspace is easier than tackling the complete search space as a whole. This technique can **identify potential unimodal**, **define a region of interest** to exploit, and **allocate resources** according to remaining evaluations.

Identifying a *unimodal* depends on the sampling frequency and the underlying model. Clustering is a common way for discovering potentail unimodals. We adopt the hierarchical clustering technique to discover hills in the fitness landscape. We also use weighted normal distribution as the underlying consumption for unimodal, and trim overfitting models with minimum description length.

After identifying the unimodals by clustering, we can split the search space into different regions of interest with linear projection. We use different projection matrices to project the original search space onto several subspaces with well-defined boundaries. We also optimize the porjection matrix so that each region would have minimum overlapping with others. Searching in the smaller subspace enhances the probability of finding optimium. The well-defined boundaries are also needed for some algorithms. The projection might also rotate or shear the original search space, making some inseparable problems easier to solve.

With multiple subspace to search, we propose a new Multi-armed Bandit (MAB) technique that optimize the resource allocation to get the greatest probability of obtaining the

global optimum. Multi-armed Bandit algorithms are suitable for this scenario, since it learns models from outcomes while the actions do not change the state of the world. When there are still plenty of evaluations left, we should prefer exploration over exploitation and search equally on all subspaces. However, when few evaluations remain, we should focus on the current best hill and try to exploit it to get the best fitness with the remaining evaluations.

1.2 Roadmap

This thesis is composed of eight chapters.

Chapter 2 presents three optimization algorithms that are used for comparisons. We introduce the Covariance Matrix Adaptation Evolutionary Strategy (CMA-ES), the Standard Particle Swarm Optimization, and the Ant Colony Optimization for Continuus Domain. These three algorithms each have different characteristics due to different underlying models.

Chapter 3 presents some clustering techniques that guides the construction of ROIs and later becomes the initial points for algorithms in each arm.

Chapter 4 first describes four basic affine transformation: translation, rotation, scaling and shearing. Then the projective transformation and homogeneous coordinate are presented.

Chapter 5 briefly describes some common multi-armed bandit algorithms, including ... Then we present our new bandit techniques. Tranditional bandit algorithms focus on minimizing regret, while our new bandit focus on the probability of getting a rank 1 result.

Chapter 6 gives details of our new algorithms. First, the framework and pseudo code are given. Then a detailed process of initialization is given in ...

Chapter 8 summarizes this thesis. The conclusion and contributions are also given. Some further improvements and future works are also discussed at the end.

Chapter 2

Real-valued Optimization Algorithms

A real-valued optimization algorithm minimize a given *objective function* in the continuous domain. It searches the continuous domain by iteratively asking for the fitness of solutions. We often evaluate an algorithm by the *number of function evaluations* it consumes to obtain the global optimum solution. This task is often described as the *black box optimization*, since the algorithm has no prior knowledge about the problem characteristics or structures. That means the gradient information might not be useful or even not available, so hill-climbing algorithms do not work. The problem may be non-convex, multimodal, noisy and discontinuous. In order to solve these kinds of problems, different algorithms have different assumptions about the problem structure. They also use different stocastic models and update methods to try to converge to the global optimum as fast as possible. These algorithms are often used for parameter and model calibration. The following section describes three kinds of milestone evolutionary algorithms that are commonly used as the baseline for real-valued optimization.

2.1 Covariance Matrix Adaptation Evolution Strategy

The *Evolutionary Strategies* (ES) samples new search points with a normal distribution. Each d-dimensional sample x_i in a population with size λ is drawn from a multivariate

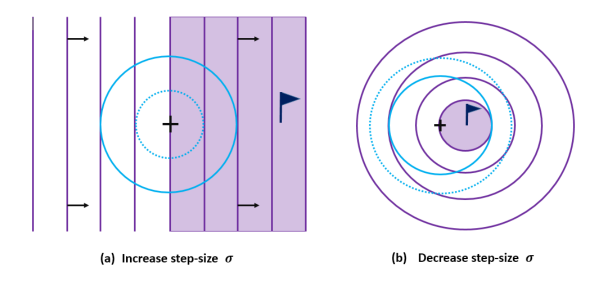


Figure 2.1: The one fifth rule of (1+1)-ES.

normal distribution as following:

$$x_i \sim m + \sigma N_i(0, C),$$

where $m \in \mathbb{R}^d$ is the mean, $\sigma \in \mathbb{R}_+$ is the step-size, and $C \in \mathbb{R}^{d \times d}$ is the covariance matrix. The mean vector represents the positions where the best solution is most likely to appear. The step-size determines the area to search and modifies exploration and exploitation behaviors. The covaraince matrix determines the shape of the distribution ellipsoid.

2.1.1 (1+1)-ES

One of the simpliest Evolutionary Strategy is (1+1)-ES. It samples one offspring x from parent m according to the equation described above. If the offspring x is better than m, it replaces m to be the new mean of the next sampling distribution. It also follows the *one-fifth success rule* as illustrated in Figure 2.1. The one-fifth success rule increases the step-size by 1.5 if the current sample is not better than the previous best sample, i.e. the mean m. We believe the best solution, denoted as the flag in Figure 2.1(a), should be somewhere outside of our current distribution and we should increase step-size to explore more search space. The one-fifth success rule decreases the step-size by $1.5^{-1/4}$ if the current sample is better than the previous best sample. This means that we believe the best solution,

illustrated as the flag in Figure 2.1(b), should be within the sampling distribution, and we should decrease the step-size to further focus on the current region. Here we describe the (1+1)-ES with one-fifth success rule with independent restarts [1]. The pseudo code of (1+1)-ES is given in Algorithm 1.

```
Algorithm 1: (1+1)-ES with 1/5 success-rule
```

 X_n : solution of the n^{th} iteration, σ_n : step size of the n^{th} iteration, $N(\mathbf{0}, \mathbf{I})$: multivariant normal distribution with mean vector $\mathbf{0}$ and identical covariance matrix \mathbf{I} .

input: f: evaluation function **output**: X_{n+1} : best solution

Initialize X_0, σ_0

while termination criterion not met do

$$\begin{vmatrix} \widetilde{\boldsymbol{X}}_n = \boldsymbol{X}_n + \sigma_n N(\boldsymbol{0}, \boldsymbol{I}) \\ \mathbf{if} \ \ f(\widetilde{\boldsymbol{X}}_n) \leq f(\boldsymbol{X}_n) \ \mathbf{then} \\ \mid \ \boldsymbol{X}_{n+1} = \widetilde{\boldsymbol{X}}_n \\ \mid \ \sigma_{n+1} = 1.5\sigma_n \\ \mathbf{else} \\ \mid \ \boldsymbol{X}_{n+1} = \boldsymbol{X}_n \\ \mid \ \sigma_{n+1} = 1.5^{-1/4}\sigma_n \end{vmatrix}$$

return X_{n+1}

2.1.2 **CMA-ES**

Covariance Matrix Adaptation Evolution Strategy (CMA-ES) is an Evolutionary Strategy proposed by Hansen in 2003 [7]. It adopts a multivariate normal distribution to generate *isotropic* search points which do not favor any direction. The multivariate normal distribution is widely observed in the nature and relatively easy to design and analysis for algorithms. CMA-ES modifies three parameters to control the multivariate normal distribution: the mean $m \in \mathbb{R}^d$, the step-size $\sigma \in \mathbb{R}_+$, and the covariance matrix $C \in \mathbb{R}^{d \times d}$.

The *mean* indicates the translation of the normal distribution model. It indicates the most favorable position that should be sampled with the largest density. It is updated with the weighted mean of all the positions according to their fitness. The *step-size* controls the state of exploration or exploitation. An ideal step-size control gradually decreases the searching region for exploitation. One of the easiest step-size control method is the one-

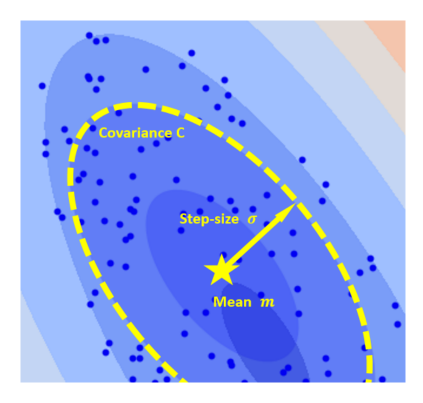


Figure 2.2: Three major elements of CMA-ES.

fifth success rule, mentioned above. Later a cumulative step-size adaptation is proposed in Cumulative Step-size Adaptation Evolutionary Strategy (CSA-ES) [8], a predecessor of CMA-ES. The cumulative step-size adaptation measures the pathway of the mean vector in the generation sequence. Adapting the step-size according to the *evolution path* enables translating more efficiently to a desired location. Later, it also allows perpendicular around the desired region. The *covariance matrix* indicates the shape of the normal distribution model. Adapting the covaraince matrix allows us to search in a region that is more coherent to the fitness landscape, This adaptation follows a natural gradient towards the direction of optimum solution. It can also learn a rotated representation of the problem and be coordinate independent. By adjusting these three parameters, shown in Figure 2.2, CMA-ES is extremely powerful on solving unimodal problems.

CMA-ES adopts a small *population size* to reduce function evaluations and increase the frequency of updating the underlying models. The default population size λ for a D-dimension problem, described in [6], is

$$\lambda = 4 + \lfloor 3\log(D) \rfloor$$

However, recent research suggest that it is benefitial to gradually increase the population size after random restart [2]. This parameterless adaptation enables few evaluations consumption for easy problems, yet is also able to solve more difficult problems with a larger population size later.

2.2 Standard Particle Swarm Optimization

Particle Swarm Optimization (PSO) is a swarm intelligence optimization algorithm. It was first proposed by J. Kennedy and R. C. Eberhart in 1995 [10] to simulate the foraging behavior of bird flocks. The swarm is composed of *particles* that move around in a given multi-dimensional *search space* to find the best solution. Each particle updates its velocity according to its historical experience, as well as the information of the neighboring particles. The neighborhood of a particle is a set of information links defined by the *swarm topology*. PSO iteratively updates the swarm topology, the velocities, and the positions of each particle until the global optimum solution is found.

Throughout the years, numerous varients of PSO have been proposed to improve performance. As a result, a *standard* PSO, composed of clear principals, is needed as the baseline for comparison. Standard PSO (SPSO) provides a well defined version that follows the common principals of PSO design. It is intended to be a milestone with simple and clear implementation for future comparison. So far, there have been three successive versions of standard PSO: SPSO 2006, SPSO 2007, and SPSO 2011. The underlying principals of these three algorithms are generally the same as all PSO varients. The exact formula and implementation are slightly different due to latest theoretical progress. A detailed description of SPSO 2011 [17] is given in the following paragrahs.

2.2.1 Initialization of the swarm

For a search space E with a given dimension D, E is confined by a set of minimum and maximum bounds in each dimension. The hyperparallelepid search space E can be

formally defined as the Euclidean product of D real intervals [4].

$$E = \bigotimes_{d=1}^{D} [min_d, max_d]$$

For each position x within the multi-dimensional search space E, there exists a corresponding numerical value f(x), i.e. *fitness*. A swarm is composed of particles, which explore different positions in the search space to find the best corresponding fitness value. At time t, each particle in the swarm possesses the following vectors with D coordinate:

- $x_i(t)$ is the **position** of the particle i at time t.
- $v_i(t)$ is the **velocity** of the particle i at time t.
- $p_i(t)$ is the **previous best position** the particle i had been to, at time t.
- l_i(t) is best position of all the previous best positions in the neighborhood of particle
 i at time t.

Let $U(min_d, max_d)$ be a random number drawn from a uniform distribution within $[min_d, max_d]$, and $N_i(t)$ be a set of neighbours of particle i at time t defined by the swarm topology. In SPSO 2011 [4], each particle is initialized with a random position and velocity defined as following:

$$x_{i}(0) = U(min_{d}, max_{d})$$

$$v_{i}(0) = U(min_{d} - x_{i,d}(0), max_{d} - x_{i,d}(0))$$

$$p_{i}(0) = x_{i}(0)$$

$$l_{i}(0) = argmin_{i \in N_{i}(0)}(f(p_{i}(0)))$$

The swarm size of SPSO 2011, denoted as S, is suggested as

$$S = 40,$$

yet it can also be defined by user [4].

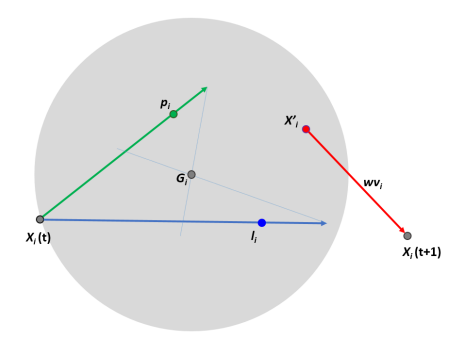


Figure 2.3: Velocity update of SPSO 2011.

2.2.2 Velocity update equations

Velocity update equations differs in different variations of PSO, since it is the core procedure of optimization that determines the performance of PSO. To prevent premature convergence, SPSO follows a random permutation order to update the positions of particles in the swarm [4]. The position of particle i at time t+1 depends on the previous position and the current velocity:

$$x_i(t+1) = x_i(t) + v_i(t+1).$$

The velocity of particle i at time t+1 can be described as a combination of three vectors:

$$v_i(t+1) = wv_i(t) + \alpha(p_i(t) - x_i(t)) + \beta(l_i(t) - x_i(t)),$$

where w mimics the momentum of the particle, and α , β describes how successive actions are effected by the personal previous best position and the neighborhood previous best position.

In SPSO 2011, the velocity update equations eliminates the coordinate dependency by

creating a hypersphere according to x_i , p_i and l_i , shown in Figure 2.3. The hypersphere is defined as:

$$H_i(G_i, ||G_i - x_i||),$$

with center G_i and radius $||G_i - x_i||$. When the personal best $p_i(t)$ is not the neighborhood previous best $l_i(t)$, the center G_i is defined as:

$$G_i = \frac{1}{3}(x_i + (x_i + c(p_i - x_i)) + (x_i + c(l_i - x_i))) = x_i + \frac{c}{3}(p_i + l_i - 2x_i)$$

If the personal best is the best in the neighborhood, i.e. $p_i(t) = l_i(t)$, the center G_i is defined as:

$$G_i = \frac{1}{2}(x_i + (x_i + c(p_i - x_i))) = x_i + \frac{c}{2}(p_i - x_i)$$

In both cases, the acceleration constant c is:

$$c = \frac{1}{2} + \ln(2) \simeq 1.193$$

As shown in Figure 2.3, a random sample x_i' is drawn from the hypershpere with uniform random direction and uniform radius.

$$r = U(0, ||G_i - x_i||)$$

Therefore, the velocity update equation and new position of SPSO 2011 is defined as:

$$v_{i}(t+1) = wv_{i}(t) + x'_{i}(t) - x_{i}(t)$$
$$x_{i}(t+1) = x_{i}(t) + v_{i}(t+1) = wv_{i}(t) + x'_{i}(t)$$

where the inertia weight is also:

$$w = \frac{1}{2\ln(2)} \simeq 0.721$$

As mentioned before, the search space is defined as a hyperparallelepid with confine-

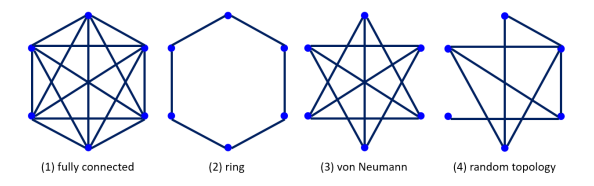


Figure 2.4: Different topologies used by PSO.

ment $[min_d, max_d]$ in each dimension d. Therefore, after calculating the new velocities and positions of particles, we have to go through confinement check to make sure all particles stay inside the search space. There are multiple options to handle confinement in SPSO 2011. The following method described in [4] treat boundary as "wall", so that all particles bounce back with half of previous velocity after they reach the border. For the new position $x_{i,d}(t+1)$ of particle i in dimension d,

$$if(x_{i,d}(t+1) < min_d), x_{i,d}(t+1) = min_d$$

$$v_{i,d}(t+1) = -0.5v_{i,d}(t+1)$$

$$if(x_{i,d}(t+1) > min_d), x_{i,d}(t+1) = max_d$$

$$v_{i,d}(t+1) = -0.5v_{i,d}(t+1)$$

2.2.3 The adaptive random topology

The swarm topology defines "who informs whom" to help distribute the information of optimum solution in the swarm. Multiple topologies shown in Figure 2.4 are adopted in different version of PSO. The random topology adopted in SPSO 2011 is updated under two circumstances [4]:

- at the very beginning
- after an iteration with no improvement of the best known fitness value

For each particle p_i in a swarm of n particles $S = \{p_1, p_2, ..., p_n\}$, a good topology

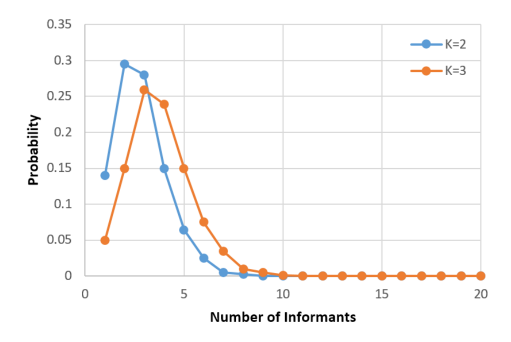


Figure 2.5: Probability of different numbers of informants in the random topology.

should allow each p_i to inform itself, along with a set of randomly selected particles from the other n-1 particles [3]. We would like most of the particles to be informed by two to four particles in the topology. Some of the particles can have no informant except for itself. There should also exist a non-zero probability for a particle to be informed by all the other particles.

In order to construct a random topology that satisfy the desirable features mentioned above, Clerc proposed a method [3] that results in a distribution of the sum of n-1 independent binary Bernouilli variables, shown in Figure 2.5. We use the distribution K=3, meaning that each particle tells three other particles about their previous best fitness and position.

2.3 Ant Colony Optimization for Continuous Domain

Ant Colony optimization (ACO) is first proposed by Dorigo [5] to solve combinatorial optimization problems, including scheduling, routing, and timetabling. These problems aim to find optimal *combinations* or *permutations* of finit sets of available components. Inspired by the foraging behavior of natural ants, ACO mimics the pheromone deposition

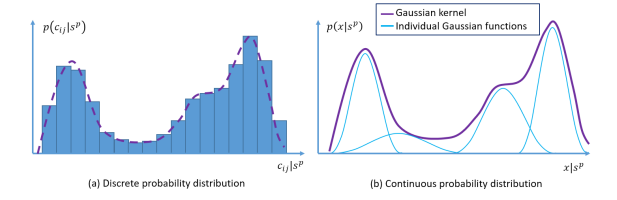


Figure 2.6: Comparison of discrete PDF continuous Gaussian kernel PDF.

of ants along the trail to a food source. The deposited pheromone, which indicates the quantity and quality of the food, creates an indirect communication among ants and enables them to find the shortest paths. Two major procedures: *solution construction* and *phermone update*, are detailed in the following paragraph.

2.3.1 Discrete solution construction and phermone update

For solution construction, we first consider a search space S defined over a finit set of all possible solution components, denoted by C. Each solution component, denoted by c_{ij} , is a decision variable X_i instantiated with value $v_i^j \in D_i = \{v_i^1, ..., v_i^{|D_i|}\}$. To construct a new solution, an artificial ants starts with an empty partial solution $s^p = \emptyset$. During each construction step, the partial solution s^p is extended with a feasible solution from the set $N(s^p) \in C \setminus s^p$. The probabilistic pheromone model adopted for selecting a feasible solution from $N(s^p)$ can be defined as follows:

$$p(c_{ij}|s^p) = \frac{\tau_{ij}^{\alpha} \cdot \eta(c_{ij})^{\beta}}{\sum_{c_{i\ell} \in N(s^p)} \tau_{i\ell}^{\alpha} \cdot \eta(c_{i\ell})^{\beta}}, \forall c_{ij} \in N(s^p),$$

where τ_{ij} is the pheromone value associated with component c_{ij} , and $\eta(\cdot)$ is a weighting function. α and β are positive parameters which determine the relation between phermone and heuristic information.

The *pheromone update*, also known as phermone evaporation, acts as solution selection. It increases the phermone values associated with promising solutions and decreases

the phermone values associated to the bad ones. This is achieved by increasing the phermone levels of more preferred solutions by $\Delta \tau$. Let $\rho \in (0,1]$ be the *evaporation rate*. If the phermone value is associated with a chosen good solution s_{ch} , the phermone value is updated as following:

$$\tau_{ij} = (1 - \rho)\tau_{ij} + \rho\Delta\tau,$$

while the other unrelated phermone values are updated as:

$$\tau_{ij} = (1 - \rho)\tau_{ij}.$$

Thus, the discrete phermone model shown in Figure 2.6(a) can be updated to emphasize the good solutions.

2.3.2 Continuous solution construction and phermone update

Over the years, multiple approaches of extending the ACO on continous domain have been given. One of the most successful version is ACO_R , proposed by Socha and Dorigo in 2008 [15]. It extends ACO to the continuous domain without making any major conceptual change to its structure. The fundamental idea underlying ACO_R is to substitue the discrete probability distribution with a continuous PDF in each dimension, demonstrated in Figure 2.6. ACO_R keeps a solution archive of **top-k solutions** it has seen so far, shown in Figure 2.7. All the solutions are sorted(ties are broken randomly) according to their fitness f(s), so that the best solution stays at the top of the archive. For the *i*-th dimension, a Gaussian kernel PDF $G^i(x)$ is constructed with multiple weighted normal distributions $g_i^i(x)$:

$$G^{i}(x) = \sum_{j=1}^{k} \omega_{\ell} g_{\ell}^{i}(x) = \sum_{\ell=1}^{k} \omega_{\ell} \frac{1}{\sigma_{\ell}^{i} \sqrt{2\pi}} e^{-\frac{(x-\mu^{i})^{2}}{2\sigma_{\ell}^{i}}^{2}}$$

Such a PDF is easy to sample and provides a better flexibility to describe the landscape. Three elements decide the shape of the Gaussian kernel:

- ω_{ℓ} is the **weight** of the solution s_{ℓ} .
- μ^i is *i*-th value of all the solutions in the archive.

Solution Archive	Fitness	Weight
$\begin{bmatrix} s_1^1 & s_1^2 & \dots & s_1^i & \dots & s_1^d \end{bmatrix}$	$f(\mathbf{s_1})$	ω_1
$\begin{bmatrix} s_2^1 & s_2^2 & \dots & s_2^i & \dots & s_2^d \end{bmatrix}$	$f(s_2)$	ω_2
$\begin{bmatrix} s_3^1 & s_3^2 & \dots & s_3^i & \dots & s_3^d \end{bmatrix}$	$f(s_3)$	ω_3
į.	÷	÷
$\begin{bmatrix} s_k^1 & s_k^2 & \dots & s_k^i & \dots & s_k^d \end{bmatrix}$	$f(s_k)$	ω_k

$$G^{1} \cdots G^{i} G^{k} G^{i}(x) = \sum_{l=1}^{k} \omega_{l} \frac{1}{\sigma_{l}^{i} \sqrt{2\pi}} e^{-\frac{(x-\mu_{l}^{i})^{2}}{2\sigma_{l}^{i}^{2}}}$$

$$\omega_{l} = \frac{1}{qk\sqrt{2\pi}} e^{-\frac{(l-1)^{2}}{2q^{2}k^{2}}}$$

$$\mu^{i} = \left\{ s_{1}^{i}, \dots, s_{k}^{i} \right\}$$

$$\sigma_{l}^{i} = \xi \sum_{e=1}^{k} \frac{|s_{e}^{i} - s_{l}^{i}|}{k-1}$$

Figure 2.7: The solution archieve of ACO_R

• σ^i is standard deviation that determines that new samples.

Following are some parameters used by ACO_R that also effects the Gaussian kernel construction. k is the size of the solution archive, which works like the memory of the swarm. It may not be smaller than the number of dimensions. q represents the locality of the search process, i.e. how often should we not pick the best solution in the archive. It controls exploration so that give a higher value of q leads to slower yet more robust convergence. $\xi > 0$ represents the speed of convergence that has effect similar to the phermone evaporation rate in ACO. It works like selection and enhance the good solutions in the archive.

The initial solution construction process proceed as following. After evaluating k randomly generates solutions, they are stored along with their fitnesses and weights in the solution archive. The weights are caluated according to the equation in Figure 2.7. Then the solution archive is sorted horizontally according to the fitness so that the best solution s_1 is on the top

For general solution construction, we first define m as the number of ants used in an iteration for sampling. During each iteration, m new solutions are constructed dimension by dimension by sampling the i-th dimension Gaussian kernel G^i A more detailed and efficient process of sampling the Gaussian kernel PDF is described in [15].

Phermone update is accomplised by adding the new samples, their fitness, and the corresponding weight into the archive. Then, the solution archive is sorted according to the fitness and we only keep the top-k solutions in the archive. This selection process ensures only the best solutions are kept in the archive. Therefore, future samples will be constructed near these best solutions and guide the ants toward the optimum.

Chapter 3

Clustering Techniques

We are interested in solving real-valued multimodal problems composed of observable unimodals. Given the case, it is easier to solve an isolated unimodal within a subspace, than tackling the complete search space with the multimodal problem. Therefore, the first thing for solving a multimodal problem is to **identify** and **isolate** the potential uni-modals within the given search space.

We tried to isolate potential *fitness hills*, i.e. uni-modals, by **clustering** the initial samples points, and consider each cluster as a multi-dimension normal distribution. Different clustering techniques are often applied to identify different characteristics of clusters. Here we proposed a *hierarchical clustering* techniques to identify *fitness hills*, since we consider not only the density of the particles, but also the fitness values of different search points. Our basic assumption for the under lying uni-modal is a weighted normal distribution, since we need to take fitness into account instead of viewing each point with the same weight. We tend to focus on the particles with better fitness, than a dense cluster with less fitness. It is also easier to calculate the weighted mean vector and weighted covariance matrix in higher dimension.

Later, we applied the **Minimum Description Length** (MDL) to reduce the number of clusters. Although a complex Gaussian Mixture Modal is able to describe the sample distribution better, we believe that a more compact model, in terms of information entropy, is the better choice when multiple models can describe the same distribution. This also allows us to define a more stable subspace for further searching.

3.1 K-Means Clustering

In this section, we first describe the K-Means clustering techniques and its limitation for identifying the fitness hills. K-Means clustering aims to partition n observation data points into k clusters. There are two main procedures for K-Means clustering: the *assignment* step and the *update* step.

During the *assignment* step, an initial set of k means positions are given. Then, each data point is assigned to the cluster with the *nearest mean*. The assignment to the *nearest mean* can be formally described as creating clusters whose mean yields the least within-cluster sum of squares (WCSS), i.e. the sum of squared Euclidean distance. During the *update* step, the new centroids of each new clusters are calculated and assigned as the new means. This can be also be described as minimizing the WCSS. The algorithm proceeds by alternating between these two steps until the means no longer change. Since there only exists a finite states of partitions, the algorithm must converge to a local optimum. However, different initial mean positions results in different clusters.

One of the main problem for using the K-Means clustering to identify *fitness hills* is that it considers only the *spatial density* and does not utilize the fitness value of each particle. Different sets of initial centers might result in different kinds of clusters. Meanwhile, each cluster might be centering at a point that does not necessarily possess the best fitness in the neighborhood. Therefore, the K-Means clustering often results in **unstable clusters** depending on initial conditions, and is highly sensitive to particle density. Another common phenomenon for being sensitive to spatial density is that the points on the *border of clusters* might belong to different clusters after each update. This makes the clusters unstable and costs unnecessary evaluations and computation for redefining the borders of ROI after each update.

The other problem for using the K-Means clustering is that one needs to **decide the number of clusters**. Determining the number of clusters is also a highly studied subjects. We breifly discuss three common estimation methods that we have tried, the *silhouette coefficient*, the *gap statistics* and the *dip test* in the following paragraph.

The silhouette coefficient [14] measures how cohesive an object is to its own cluster

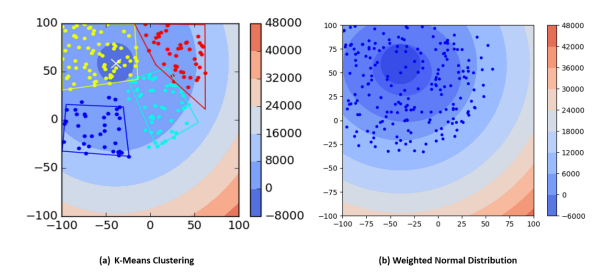


Figure 3.1: Comparing K-Means clustering with weighted Gaussian Distribution on CEC2005 F1 Problem

and how seperate it is to the other clusters. Ranging from -1 to 1, the silhouette coefficient with a higher value indicates it is more likely to belong to its own cluster. The gap statistics [16] compares the change in within-cluster dispersion with that of an reference null distribution. It calculates the gap statistics for different number of clusters and select the one with the maximum gap statistics. The Hartigans' dip test [9] uses the empirical cumulative distribution function (ECDF) to measure the maximum difference between multimodal samples and the unimodal samples. It calculates maximum deviation of the sample ECDF from the unimodal CDF and check if the difference is statistically significant.

We wish that as the algorithms proceed, particles should gather around the good solutions and gives a clear density signal. However, they seldom identify a unimodal at the beginning. It means that we need to add up more particles to meet the population size requirement for each clusters. This increases the density in that specific region, and creates an artificial cluster that should not exist in the first place.

Figure 3.1 shows how K-Means fails to identify a unimodal and results in four clusters due to density. A more preferable clustering result is shown on the right in Figure 3.1. We would like our clustering method to be capable of identifying the uni-modal and be able to center around the particle with the best fitness.

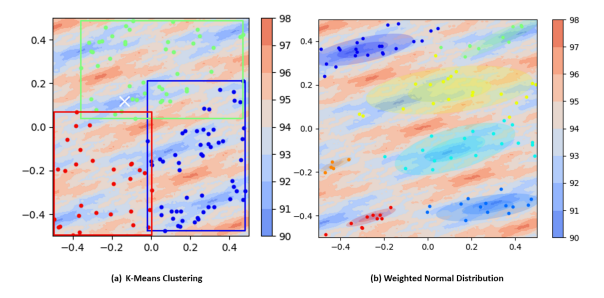


Figure 3.2: Comparing K-Means clustering with weighted Gaussian Distribution on CEC2005 F11 Problem

Figure 3.2 shows another example of how K-Means fails to identify the right multi-modals. Since K-Means only consider the density, it is hard to recognize the dense hills in the landscape. This initial clusters leads to more unnecassy evaluations for merging and reidentifying the hills. However, we can see that on the right, our technique successfully recognize almost all the hills in the problem. It also helps to locate the local optimum solutions approximately at the center. This gives alot of advantage for the algorithms to search, since the task becomes exploiting a single hill. Therefore, later we adopted the weighted normal distribution as the underlying model.

3.2 Heirarchical Clustering

Since we would like to recognize the *fitness hills*, we modified the agglomerative clustering technique to create hierarchical clusters. For a problem in D dimension, we defined the neighborhood of a particle to be the $4 + \lfloor 3\log(D) \rfloor$ particles with the minimum Euclidean distance, including the particle itself. Given the neighborhood of each particle, we create a directed graph by pointing each particle to the particle with the highest fitness in the neighborhood. If a particle does not possess the best fitness in its neighborhood, it points toward the direction with the maximum gradient. If it is the best particle in the

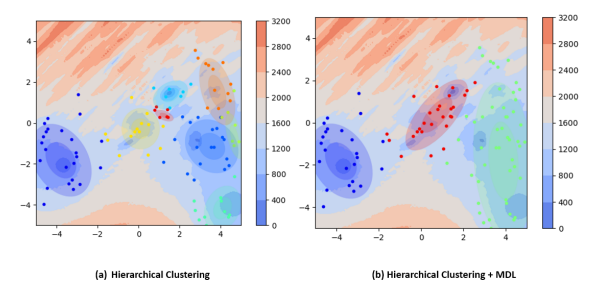


Figure 3.3: Before and after Minimal Description Length trimming on CEC2005 F19 Problem

neighborhood, it is very likely to be at the approximate location of local minimum.

We then create trees with the directed graph, where each tree represents a cluster with a potential fitness hill. The best position within this cluster is indicated by the root node. This way, the fitness signal guides the construction of clusters, so that this method is able to identify more *fitness hills* than the K-Means clustering. Moreover, we do not need to provide the estimate number of clusters, since the agglomerative clustering method *intrinsically identifies hills* These characteristics solves some fundamental problems that we faced when using the K-Means clustering along with different number-of-clusters-estimation techniques.

We can see that for a multimodal problem depicted in Figure 3.3(a), the hierarchical clustering technique successfully identifies several fitness hills. However, for the unimodal problem in Figure 3.4(a), the hierarchical clustering creates multiple "ripples" on the marginal area. These clusters are created when the best particle is a bit further from the next best particle. This indicates that the using the Euclidean distance as the distance metric is not suitable for problems with larger hills.

In order to solve this problem, we assume the underlying hills are composed of multivariate normal distribution. Given the distribution and a sample position, we can use the *Manhanoblis distance* to serve as a more expressive metric. We would also like to

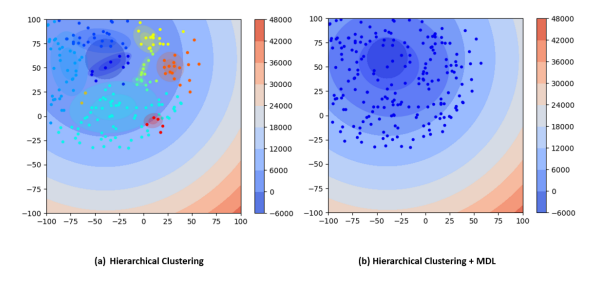


Figure 3.4: Before and after Minimal Description Length trimming on CEC2005 F1 Problem

trim down the number of clusters in the beginning, since we believe that the problems are composed of hierarchical hills. We do not need to invest so much resources in so many regions during the initialization. Reclustering after a few iterations shall give us a better signal about where to focus.

3.3 Minimum Description Length

The Minimum Description Length (MDL) principle was introduced by Rissanen in 1978 [12]. It adopts the idea of Occams's razor by selecting the codec that gives the shortest codelength of the data. It provides a natural safeguard against overfitting, since it considers not only the goodness of the model, but also the complexity of the data to give the hypothesis. Here we modified the MDL described in [11] to help us determine the optimum number of clusters that provides the most efficient codes for the sample data. The MDL considers both *data description accuracy* and *model description efficiency*.

3.3.1 Data description accuracy

Since we would like to take the fitness into account, we use the weighted Gaussian distribution as the underlying model. For a cluster with N particles, each particle X_i has a rank r_i between 1 and N (ties are broken randomly) that corresponds to the fitness. The weight

 ω_i can be defined as:

$$\omega_i = \log(N + 0.5) - \log(r_i) \tag{3.1}$$

The mean μ and covariance matrix Σ in a weighted Gaussian distribution N can be denoted as:

$$\mu = \frac{1}{\sum_{i=1}^{N} \omega_i} \sum_{i=1}^{N} \omega_i X_i$$
 (3.2)

$$\Sigma = \frac{1}{\sum_{i=1}^{N} \omega_i} \sum_{i=1}^{N} \omega_i (X_i - \mu)^T (X_i - \mu)$$
 (3.3)

Let $X = \{X_1, X_2, ..., X_I\}$ denote the data set, composed of I samples. Each sample is a D-dimensional vector $X_i = \{X_{i1}, ..., X_{iD}\}$. The class-probability of observing sample X_i in class j, described by a parameter set Θ_j , can be denoted as $P_j(X_i|\Theta_j)$. The probability of observing the sample X_i in a finite mixture model with J components can be denoted as:

$$P(X_i|\Theta) = \sum_{j=1}^{J} \alpha_j P_j(X_i|\Theta_j)$$
(3.4)

$$\alpha_j = n_j / I \tag{3.5}$$

where α_j is the prior probability that X_i belongs to the j-th model, and n_j denotes the number of samples in cluster j.

Therefore, the probability of observing sample X_i in cluster j can be further described by the j-th weighted Gaussain distribution:

$$P(X_i|\Theta_j) = N(X_i|\mu_j, \Sigma_j)$$
(3.6)

With the assumption that the data instance X_i are independently distributed, the joint data probability is the product of the individual instance probabilities:

$$P(X|\Theta) = \prod_{i=1}^{I} \sum_{j=1}^{J} \alpha_j P_j(X_i|\Theta_j)$$
(3.7)

3.3.2 Model description efficiency

Although a more complex model can describe the data distribution better, we believe that a simpler model with approximately the same accuracy is better. Therefore, the number of parameters that a model requires, is also considered in MDL. The free parameters in the Gaussian mixture model are:

- J-1 paramters for J weights α_j , since $\sum \alpha_j = 1$
- D parameters for each mean μ_i
- D(D+1)/2 parameters for each covariance matrix Σ_j

Therefore, the total number of free parameters is

$$\mathbb{K} = J - 1 + J(D + D(D+1)/2) = J(D^2 + 3D + 2)/2 - 1 \tag{3.8}$$

Combining the *data description accuracy* and *model description efficiency*, the MDL defined in [13] is denoted as:

$$\min_{\mathbb{K},\Theta} -\log(P(X|\Theta)) + \frac{1}{2}\mathbb{K}\log(I)$$
(3.9)

3.3.3 Trimming model with MDL

As mentioned before, we need to solve the problem of having too many "ripple" clusters on the margin area. Here we would like to use the MDL Equation to calculate model efficiency, and trim down unnecessary clusters. Therefore, after identifying potential unimodals with hierarchical clustering in Section 3.2, we use Equation 3.3.2 to calculate model efficiency. We created mutliple weighted Gaussian distribution according to the hierarchical clusters, as shown in Figure 3.4(a) and Figure 3.3(a). Then we try all combinations of merging two clusters and calculate the corresponding MDL. Therefore, we compare the MDL score between the original hierarchical clusters and the new clusters

with the minimal MDL score. If the MDL score of the new model is smaller than the previous model, we repeat the same procedure and keep merging. This trimming procedure terminates when there is only one cluster remain, or when the minimum MDL score of all possible new modals is larger than the MDL score of the previous model. Then we get fewer yet still expressive clusters, shown in Figure 3.4(b) and Figure 3.3(b).

Linear Projection

We would like to define a region of interest that contains only one uni-model for exploitation. Also, some algorithms, e.g. SPSO, requires a well-defined hypercube search space with fixed-value constraints in each dimension. Defining the projection onto subspace can be described as a model selection. In our case, this process does not require high accuracy, yet has a strict demand on time consumption. Combining the requirements, we use a linear projection matrix to project the original search space onto a subspace with feasible solutions bounded within [0,1] in each dimension.

One of the advantage for using projection matrix is that for a problem with ℓ variables, a linear projection matrix on homogeneous coordinate only requires $(\ell+1)^2$ hyperparameters to be optimized. Therefore, less hyperparameters are assigned than directly define each hyperplane borders or vertices. This reduces the parameters that we need to optimize and results in less time consumption during model selection. We also use the (1+1)-ES, a fast and simple evolutionary strategy, to optimize the matrix. It allows us to rapidly approximate a high dimensional projection matrix within given number of iterations.

Furthermore, subspace projection also gives advantage for solving *inseparable problems*. For variabels that are not independent, projection allows the algorithm to solve an easier, rotated and sheared problem on subspace, as shown in Figure 4.1. The homogeneous projection matrix separates the original ROI into less overlapped ROI, which reduces redundant search and sometimes enlarges the crutial regions.

In the following sections, we first describe the cannonical affine transformation. Then

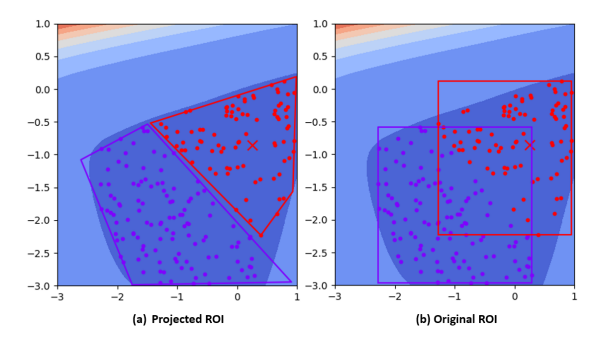


Figure 4.1: Projecting inseparable problems onto subspace.

we discuss how the projection matrix allows linear projection in a homogeneous coordinate. Finally, we give details of how we design the cost function and how we utilize the (1+1)-ES to optimize the projection matrix.

4.1 Affine Transformation

In geometry, an affine transformation preserves points, straight lines and planes. For two affine spaces A and B and any pair of points $P,Q\in A$, an affine transformation f determines a linear transformation φ that can formally defined as:

$$\overrightarrow{f(P)f(Q)} = \varphi(\overrightarrow{PQ})$$

Here, we list some common 2D affine transformation matrix in Figure 4.2.

1. Translation

Translation moves every point in the cluster by the same amount in a given direction. It allows us to reallocate the center of ROI to the positions with the best fitness.

2. Rotation

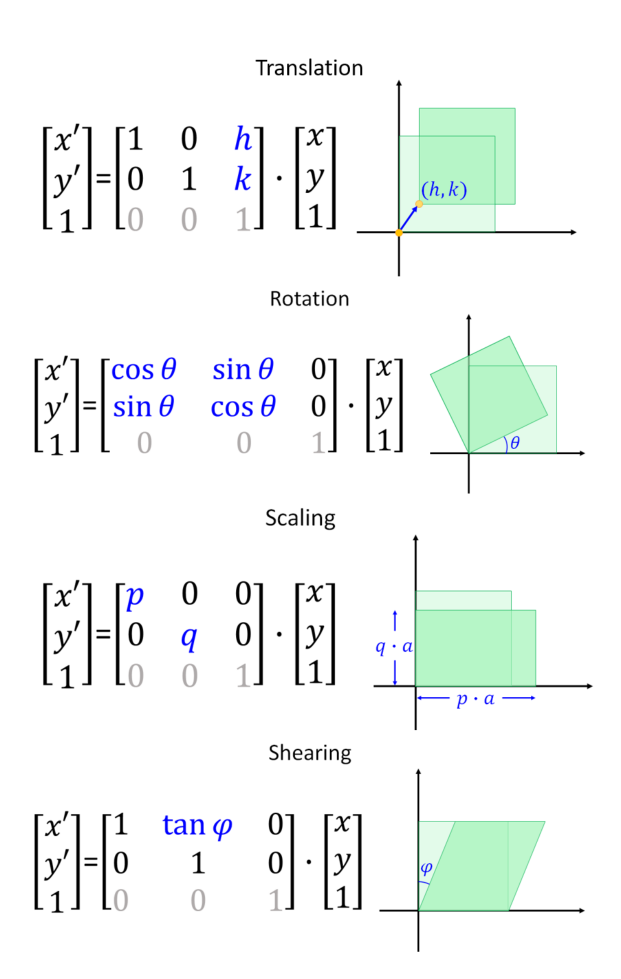


Figure 4.2: Some common affine transformation

Rotation is a common method to make a problem inseperable, by creating dependencies among variables. Therefore, using a rotation matrix to rotate the problem allows us to alter the problem characteristics. In higher dimensions, rotating abouth different axis requires different form of matrix. Here we only demonstrate a simple two dimensional case in Figure 4.2.

3. Scaling

Scaling transformation can stretch or shrink a given search space. Since it changes the lengths and angles, scaling is not an Euclidean transformation. In search space projection, it allows us to enlarge certain areas that we are interested in, and shink some area that we tend to ignore on the subspace.

4. Shearing

Shearing transform acts like "pushing" the search space in a direction parallel to a certain plane or axis. It changes the lengths and angles of certain area on the subspace. This transformation also allows us to expend certain interesting areas on subspace for a more thorough search.

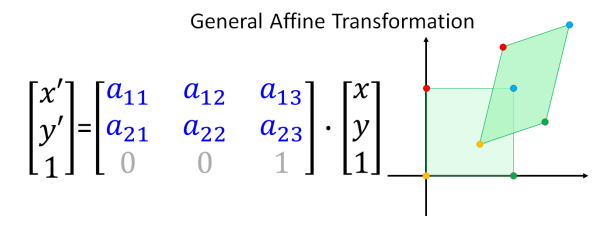


Figure 4.3: General affine transformation in 2D

The general affine transformation can be expressed as the form in Figure 4.3. All the common affine transformation matrix mentioned above, including translation, rotation, scaling and shearing can fit into this form. The general affine transformation can be seen

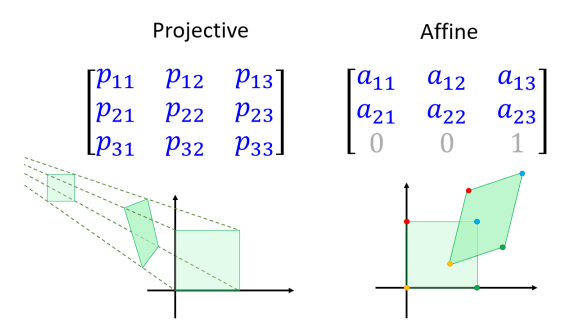


Figure 4.4: Affine transformation and projective transformation comparison.

as the matrix product of several affine transformation matices. Although the affine transformation does not preserve the lengths and angles of certain shape, it does not alter the degree of a polynomial. That means parallel lines and planes are transformed to parallel lines and planes, and intersecting lines and planes are also transformed into intersecting lines and planes. It preserved certian geometric characteristics, which is benefitial for us to do searching on subspace.

However, the affine transformation can only creat a hyperparallelepid ROI, as shown in Figure 4.4 The parallelepid shape is sometimes not flexible enough to define a non-overlapping ROI that matches the shape of the fitness landscape. In order to aquire a more flexible quadrilateral ROI, we need the projective transformation. Projective transformation is more expressive than affine transformation since the last row does not have to be 0, 0, and 1.

4.2 Projective Transformation

Projective transformations are the most general linear transformations and require the use of *homogeneous coordinates*. The *homogeneous coorinate* is also called *projective coordinates*, since it is a system of coordinates widely adopted in projective geometry. Given

Projective Transformation

$$\begin{bmatrix} p_{11} & p_{12} & p_{13} \\ p_{21} & p_{22} & p_{23} \\ p_{31} & p_{32} & p_{33} \end{bmatrix} \cdot \begin{bmatrix} x \\ y \\ 1 \end{bmatrix} = \begin{bmatrix} x' \\ y' \\ z' \end{bmatrix} = \begin{bmatrix} x'' \\ y'' \\ 1 \end{bmatrix} \cdot z'$$

Figure 4.5: Projective transformation matrix.

Inverse Affine Transformation

$$\begin{bmatrix} a_{11} & a_{12} & a_{13} \\ a_{21} & a_{22} & a_{23} \\ 0 & 0 & 1 \end{bmatrix}^{-1} \cdot \begin{bmatrix} x' \\ y' \\ 1 \end{bmatrix} = \begin{bmatrix} x \\ y \\ 1 \end{bmatrix}$$

Inverse Projective Transformation

$$\begin{bmatrix} p_{11} & p_{12} & p_{13} \\ p_{21} & p_{22} & p_{23} \\ p_{31} & p_{32} & p_{33} \end{bmatrix}^{-1} \cdot \begin{bmatrix} x'' \\ y'' \\ 1 \end{bmatrix} = \begin{bmatrix} x_r \\ y_r \\ z_r \end{bmatrix} = \begin{bmatrix} x + e_x \\ y + e_y \\ 1 \end{bmatrix} \cdot z_r$$

Figure 4.6: Comparison between inverse affine transform and inverse projective transformation.

a point (x, y) on the Euclidean plane, than (xZ, yZ, Z) is called a set of homogeneous coordinates for the point, for any non-zero real number Z. One of the great adventage of homogeneous coordinate is that the points, including the ones at infinity, can be represented with simplier, finite coordinates. That is, when Z is 0, the homogeneous coordinates of the point is defined at infinity.

Therefore, as shown in Figure ??, for given a position (x, y) on the original search space, we get a projected homogeneous coordinate (x'', y'') on the subsapce. We can generalize this to any d-dimensional space. The *homogeneous convention* is to convert the projected position, a vector with (d+1) variables, to a d variable vector by dividing first d elements with the last element.

However, unlike the affine transformation, the projective transformation does not guarantee inverse transform. A $d \times d$ projection matrix does not guarantee a non-zero matrix

determinant. Also, after inverse transform, there could be an error (e_x, e_y) that deviates from the original position (x, y), as shown in Figure 4.6. Therefore, we need to *optimize* the projection matrix in order to aquire a more suitable ROI, as well as small reconstruction errors for each samples.

4.3 Optimization for Projection Matrix

After identifying different clusters of samples, which represents a uni-modal, we would like to further define a reasonably good ROI. In order to obtain a nice ROI for a problem with D variables, we need to optimize the $(D+1)^2$ parameters in the projection matrix with a loss function. The loss function should cost little computational time and should suggest the right direction for optimization, in order to match the restrictions of a well-defined ROI. Also, the optimization only needs to define a reasonably well ROI for the alogirthms to search. In the later iterations, the matrix update procedure allows us to define a more accurate ROI with a better insight of the uni-modal subproblem. In our case, it is acceptable to sacrefice a bit of accuracy for computational time in hyperparameters tuning. We will describe the desgin of our loss function and the advantage of using a (1+1)-ES to optimize the matrix in the following sections.

4.3.1 Loss function

The goal of the ROI is to seperate the search space into non-overlapping subspaces, each containing an approximately normalized fitness hill. In our loss function, we considered the following features:

- 1. Distances to the boundary for the **points within the cluster** yet excluded in the ROI.
- 2. Distances to the boundary for the **random samples within other ROIs** yet included in the ROI.
- 3. Distances to the boundary for the **random samples** in the subspace that are outside of the original search space boundary.

- 4. Distance of the weighted **mean** to the center of subspace $[0.5]^D$
- 5. The Mean Absolute Error (MAE) for each element in the weighted **covariance matrix** to a scaled identicle matrix

6. The sum of **reconstruction** error for each particle

We add up all the errors mentioned above and try to minimize that error. Following part explains the reason for considering each features.

First, we would like to keep all the particles that belong to this cluster in the boundary $[0,1]^D$. It can easily be done by projecting all points within the cluster onto the subspace, and check whether if any of the projected position is outside of the boundary $[0,1]^D$.

Second, we would like to avoid ROIs from overlapping by adopting the Monte Carlo method for region collision detection. When optimizing one ROI, we randomly generate a given amount of samples within the subspace for all the other ROIs. Then, we use the inverse projection matrix of each other ROIs to project these sample points back to the original search space. After that, we use the projection matrix of the ROI that we are optimizing to project all the exterior samples onto the subspace. This way, we can easily discover which exterior samples are within the $[0,1]^D$ boundary. Figure 4.7 shows a screen shot of the red ROI before and after optimizing the projection matrix of the red ROI. We randomly generates 200 samples within the green and blue ROI and tries to exclude them in the red ROI.

Third, we also use the Monte Carlo method described before to make sure that the ROI does not exceed the boundaries of the original subspace. We randomly generate samples from the subspace and inverse project it back to the search space to make sure that all points are within boundaries. If not, we would like to minimize the distance of sample points to the nearest border.

Forth, we would also like each underlying model for each subproblem to be a multivariate gaussian distribution with mean around the center $[0.5]^D$. If the algorithm searches near the boundary, it indicates that ROI should be updated since the optimum solution might be outside of the border. Placing the optimum solution at the center of the search

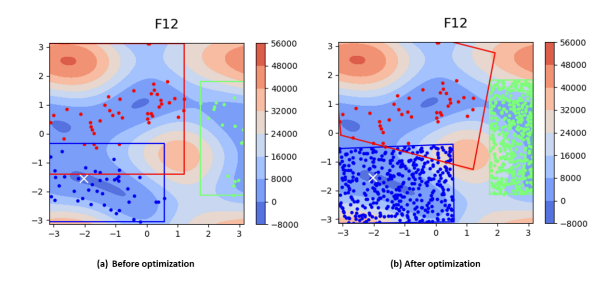


Figure 4.7: Optimization of projection matrix.

space creates a more stable model for the algorithm to search. This allows a thorough search around the center, and decreases the frequency of ROI update. Therefore, we calculate the weighted mean of the points within the cluster on the subspace. Then we minimize the distance of weighted mean to the center.

Fifth, we would like the weighted covariance matrix to be approximately 0.2I, where I is an identicle matrix. This enables the projection matrix to rotate, scale, and shear a fitness hill. The normalization preprocess is a common technique to make future optimization easier. Therefore, we calculate the MAE between the weighted covariance matrix and 0.2I.

Finally, since the we project the particle positions in a homogeneous coordinate, the correlations between reconstructed positions on the subspace might not be identical to the original ones. We would like to minimize the transformation error while keeping the ROI within the original boundaries. Therefore, for each possible matrix solution, we use it to project all points in the corresponding cluster onto a subsapce. Later, we use the inverse matrix to project all the points back to the original search space to get the reconstructed positions. Then, we calculate the MAE of all points in all dimensions between the original position and the reconstructed positions.

4.3.2 Optimization Algorithms

With the loss function defined in the previous section, we still need an optimization algorithm to optimize the projection matrix. The difficulty for *hyperparameters optimization* is to balance between speed and optimization ability. We need the hyperparameters to define a subspace that contains the possible optimum solution while costing few computation time.

We can simplify and stablelize the optimization process by giving a fixed initial solution. Here, the initial solution is a simple dot product of a translation matrix and a scaling matrix. The matrix defines a hypercube, which is bounded by the minimum and maximum value in each dimension of all positions in the cluster. This allows the algorithm to start the search around a more prefered neighborhood.

We first tried CMA-ES to optimize the projection matrix. However, we found that it cost too much time to optimize the projection matrix with 961 variables in the 30-D problem Later, we utilize the (1+1)-ES described in Algorithm 1. We would like to reduce the computation time by giving the (1+1)-ES only one hundred iterations for each run of three random restarts. (1+1)-ES works well with one of the result shown in Figure 4.7

Finally, the infeasible solutions that belong to the cluster yet being outside of ROI after the optimization are repaired with the following steps. We normalize the distance between the infeasible solution and the center of ROI, to project it onto a hypershpere centering at $[0.5]^D$ and with radias 0.5. This way, although the reconstructed position is not accurate on the original search space, it is still able to indicate the approximate fitness in that direction in the subspace. Also, our basic assumption is a unimodal hill that puts the best fitness point in the center. Therefore, marginal points should have lower fitness and should not spend more resources to exploit. It just needs to give us a hint of the gradient in that direction.

Multi-armed Bandit Algorithms

After identifying the unimodals in Chapter 3 and defining a ROI for exploitation in Chapter 4, we still need to decide how to allocate our resources. Describe the exploration vs. exploitation dilemma. One should take different strategies according to evaluations left. Instead of letting the algorithms handling both eploration and exploitation, we propose a bandit technique to help manage exploitation.

Multi-armed Bandit Algorithms are suitable for this scenario. It learns model from outcomes and the actions do not change the state of the world.

Unlike canonical MAB algorithms that minimize regrets, we wish to maximize the probability of gaining the maximum rank.

5.1 The Multi-armed Bandit Problem

Multi-aremed Bandit (MAB) Problem describes an agent needs to decide in K arms to pull at time t and receives a reward.

Describe policy and regret.

The goal is to minimize regret.

5.2 Some common MAB Algorithm

5.2.1 UCB

5.2.2 POKER

The Price of Knowledge and Estimated Reward (POKER) strategy considers three ideas: pricing uncertainty, exploiting the lever distribution and taking into account the horizon

5.3 The New Bandit Technique

The New Bandit Technique

In this chapter, we illustrate the proposed algorithm step-by-step.

Overview of real-valued optimization

6.1 Framework of the New Bandit Algorithm

The goal is to identify the ROI for exploitation and maintain exploration through resource allocation.

Each algorithm needs to be modified to satisfy the following conditions:

- 1. Update one individual at a time
- 2. The algorithm can be projected onto a subspace and continue iterating
- 3. Replace one individual with a given position and fitness

The pseudocode of our new algorithm is given in Algorithm ??

6.2 Initialization and Unimodal Identification

For a problem with D dimension, initialize with 100D points.

Keep only top 50% of points for unimodal identification. Use clustering techniques mentioned in Chapter 3.

Iteratively add 10D points until estimated cluster number is identical.

Algorithm 2: Framework of the new Bandit Algorithm

```
oldsymbol{X}_n: solution of the n^{th} iteration, \sigma_n: step size of the n^{th} iteration, N(\mathbf{0}, \boldsymbol{I}): multivariant normal distribution with mean vector \mathbf{0} and identical covariance matrix \boldsymbol{I}.

input: f: evaluation function output: X_{n+1}: best solution

Initialize oldsymbol{X}_0, \sigma_0

while termination criterion not met \mathbf{do}

oldsymbol{\widetilde{X}}_n = oldsymbol{X}_n + \sigma_n N(\mathbf{0}, oldsymbol{I})

if f(\widetilde{oldsymbol{X}}_n) \leq f(oldsymbol{X}_n) then

oldsymbol{X}_{n+1} = \widetilde{oldsymbol{X}}_n

\sigma_{n+1} = 1.5\sigma_n

else

oldsymbol{X}_{n+1} = oldsymbol{X}_n

\sigma_{n+1} = 1.5^{-1/4}\sigma_n
```

With a given cluster number k, do K-Means clustering.

6.3 Define Region of Interest

return X_{n+1}

Each arm is composed of an algorithm and a projection matrix. Initial matrix is set as a tight hyperbox that contains all points in the given cluster. Optimize matrix one-by-one. Resize each cluster to match the required population for each algorithm. Start algorithm with given initial positions and fitnesses.

6.4 Remain Evaluations Allocation

Calcualte remain evaluations allocatoin and normalize the value to 0 1. Add newly calculated allocatoin to record. Choose arm the argmax allocation to pull, i.e. update one individual. Max arm allocation -1.

6.5 Recluster

Experiments

Overview of real-valued optimization

7.1 Test Problems

7.1.1 CEC2005 25 benchmark problems

Describe termination criterion and evaluation method for CEC2005 Describe reason for 25 repeat runs. Describe how initial setting effect real-valued optimization, so that all algorithms should start with identicle initial status.

7.2 Experiment Settings

Describe the parameters setting for CMA-ES, SPSO and ACOR.

We use the *Covariance Matrix Adaptation Evolution Strategy for non-linear numerical optimization in Python* package ¹ provided by Hensen, the author of CMA-ES. CMA-ES initial mean and std, population settings.

SPSO2011 parameters (c, w) settings, and population settings.

For ACOR, we set our parameters according to the original paper [15]. The parameters are shown in Table 7.2.

¹https://pypi.python.org/pypi/cma

Parameter	Symbol	Value
No. of ants used in an iteration	m	2
Speed of convergence	ξ	0.85
Locality of the search process	q	10^{-4}
Archive size	k	50

Table 7.1: Summary of the parameters used by ACO_R

Describe our bandit parameters setting, including the initial population, maximum number of arms, and (1+1)-ES step size.

Conclusion

Contribution: find potential region to search allocate resources

Weakness: Potentially more NFE on unimodals and moving regions. Requires a better

clustering techniques to identify underlying unimodals.

Future work: Non-linear transformation

Attention plays an important role in human vision. For example, when we look at an

image, our eye movements comprise a succession of fixations (repetitive positioning of

eyes to parts of the image) and saccades (rapid eye jump). Those parts of the image that

cause eye fixations and capture primary attention are called regions of interest (ROIs).

Studies in visual attention and eye movement have shown that humans generally only

attend to a few ROIs. Detecting these visually attentive regions in images is challenging

but useful in many multimedia applications, such as automatic thumbnail cropping, object

recognition, content-based image retrieval, adaptive image compression and automatic

browsing in small-screen devices.

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