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A Reliable Method of Minimum Zone Evaluation of Cylindricity and Conicity from Coordinate Measurement Data

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Abstract

The form error evaluation of cylinders and cones is very important in precision coordinate metrology. The solution of the traditional least squares technique is prone to over-estimation, as a result unnecessary rejections may be caused. This paper proposes a reliable algorithm to calculate the minimum zone form errors of cylinders and cones, called a hybrid particle swarm optimization-differential evolution algorithm. The optimization is conducted in two stages, so that the program can hold a fast convergence rate, while effectively avoiding local minima. Experimental results demonstrate that the proposed algorithm can obtain very accurate and stable results for the calculation of cylindricity and conicity.

Keywords: Form error, Cylindricity, Conicity, Minimum zone, Differential evolution

2000 MSC: 90C47, 90C31, 65K10

1. Introduction

The evaluation of cylindricity and conicity is among the most important problems in computational metrology because cylindrical and conical shapes are ubiquitous in precision components. Most current commercial software evaluates the form errors of cylinders and cones using the least

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squares method due to its ease of implementation and the unbiasedness of its solution for uncorrelated normally distributed noise [1]. But it is likely to overestimate the form tolerance and lead to unnecessary rejections, therefore its solution is only an approximate one, rather than the optimum. According to ISO 1101 [2], a geometrical tolerance applied to a feature defines the tolerance zone within which that feature shall be contained. This means the cylindricity tolerance is a permissible deviation zone bounded by two coaxial cylinders within which the measured data must lie in between. Similarly, the conicity tolerance is a permissible deviation zone bounded by two coaxial cones. They are minimax problems and not continuously differentiable, thus very difficult to be solved.

A lot of linear programming techniques have been employed for the evaluation of straightness and flatness, e.g. simplex method [3], data exchange algorithm [4] and support vector machine [5]. The calculations of roundness and sphericity are quadratic programming problems. [3] and [6] approximately linearized the formulation and solved it using the simplex technique. Computational geometry methods were also utilized to calculate flatness [7] and sphericity [8, 9]. Venkaiah and Shunmugam calculated cylindricity by identifying 'extreme points' to form a limacon-cylinder [10]. These methods are developed for a particular shape. So far no computational geometry method has been found in literature for evaluating conicity. Recently Huang and Lee solved the conicity problem using minimum potential energy algorithms [11], which are very elaborate. Some researchers attempted to approximate the cylinder/cone iteratively and then solved a sequence of linear programs [12, 13], but these methods need a very good initial guess and are prone to sub-optimal solutions. Lai and Chen [14] converted a cylinder into a plane and obtained appropriate control points. The cylindricity is calculated by implementing a series of inverse transformations.

In recent years heuristic search optimizers have been employed to solve the minimum zone problem. Lai et al. [15] and Liu et al. [16] used genetic algorithms to evaluate the cylindricity and conicity errors, respectively, while Wen and Song adopted an immune evolutionary algorithm for sphericity [17]. The particle swarm optimization technique was utilized by [18, 19]. The advantages and shortcomings of these algorithms will be discussed in Section 3.

This paper presents a two-stage method called a hybrid particle swarm optimization-differential evolution algorithm for the evaluation of cylindricity and conicity errors. This method shows great superiorities on stability and

accuracy, and makes a good balance between exploration and exploitation.

2. Formulation of cylindricity and conicity errors

2.1. Cylindricity error

Fig.1 illustrates the cross section of a cylinder with axis direction $\mathbf{n}(m, n, 1)$ and a radius R. The projection of a measured point P onto the cylinder is F. Assuming the axis passes the point $Q(x_0, y_0, 0)$, then the axis function is $(x - x_0)/m = (y - y_0)/n = z$. The distance from P to the cylinder is,

$$e = |FP| = |EP| - |EF| = \frac{|\overrightarrow{QP} \times \mathbf{n}|}{|\mathbf{n}|} - R$$

In the equation $|\cdot|$ means the length of a vector in the Euclidean space. Given a point set $\mathbf{P} = \{P_k | k = 1, 2, ..., M\}$, the cylindricity error is,

$$d = \max_{k} e_{k} - \min_{k} e_{k}$$

$$= \max_{k} \frac{|\overrightarrow{QP_{k}} \times \mathbf{n}|}{|\mathbf{n}|} - \min_{k} \frac{|\overrightarrow{QP_{k}} \times \mathbf{n}|}{|\mathbf{n}|}$$
(1)

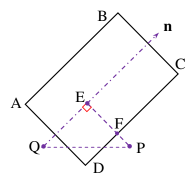


Figure 1: Calculation of cylindricity error

2.2. Conicity error

Fig. 2 shows the cross section of a cone with vertex angle 2θ , axis direction $\mathbf{n}(m, n, -1)$ and vertex $A(x_0, y_0, z_0)$. The projection point of a measured point P onto the cone is D. The distance from P to the cone is,

$$e = |DP| = \cos\theta(|BP| - |BC|) = \cos\theta|BP| - \sin\theta|AB|$$

Given a point set $\mathbf{P} = \{P_k | k = 1, 2, \dots, M\}$, the conicity error is,

$$d = \max_{k} e_{k} - \min_{k} e_{k}$$

$$= \max_{k} \left(\cos \theta \frac{\overrightarrow{AP_{k}} \cdot \mathbf{n}}{|\mathbf{n}|} - \sin \theta \frac{|\overrightarrow{AP_{k}} \times \mathbf{n}|}{|\mathbf{n}|} \right)$$

$$- \min_{k} \left(\cos \theta \frac{\overrightarrow{AP_{k}} \cdot \mathbf{n}}{|\mathbf{n}|} - \sin \theta \frac{|\overrightarrow{AP_{k}} \times \mathbf{n}|}{|\mathbf{n}|} \right)$$
(2)

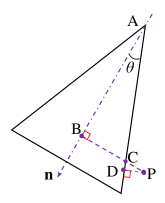


Figure 2: Calculation of conicity error

3. A hybrid particle swarm optimization-differential evolution algorithm

3.1. Particle swarm optimization

The particle swarm optimization (PSO) method is a swarm intelligence method for global optimization first proposed by Kennedy and Eberhart [20].

In PSO, each individual, named particle, of the population, called swarm, adjusts its trajectory toward its own previous best position, and toward the previous best position attained by any member of its topological neighbourhood. As a stochastic search scheme, PSO has properties of simple computation and rapid convergence capability. The individuals in evolutionary algorithms are primarily competitive. On the contrary, PSO adopts a more cooperative way. As a consequence it can find the optimal region of the solution very easily, i.e. it has outstanding capability of exploitation. Its populations move as a whole but do not evolve, thus no mutation or crossover is needed.

Suppose the search space is D-dimensional, and the positions of particles $\mathbf{Y} = \{\mathbf{y}_i | i = 1, \dots, N\}$ are initialized using a uniform distribution. The velocity (the position change per generation) of \mathbf{y}_i at the k-th generation is given as,

$$\mathbf{v}_i^k = \mathbf{v}_i^{k-1} + c_1 \mathbf{r}_1 \otimes (\mathbf{p}_i - \mathbf{y}_i) + c_2 \mathbf{r}_2 \otimes (\mathbf{p}_g - \mathbf{y}_i), k > 1$$

where c_1 and c_2 are acceleration coefficients regulating the relative velocities toward local best \mathbf{p}_i (exploration) and global best \mathbf{p}_g (exploitation), respectively. \mathbf{r}_1 and \mathbf{r}_2 are D-dimensional random vectors uniformly distributed in [0,1] and \otimes denotes element-wise multiplication.

To endure convergence and to improve the stability of the basic PSO, Clerc proposed the use of a constriction factor to the velocity [21],

$$\mathbf{v}_i^k = \chi[\mathbf{v}_i^{k-1} + c_1\mathbf{r}_1 \otimes (\mathbf{p}_i - \mathbf{y}_i) + c_2\mathbf{r}_2 \otimes (\mathbf{p}_q - \mathbf{y}_i)], k > 1$$
(3)

Following [21], the coefficients are set as $c_1 = c_2 = 2.05$ and $\chi = 0.7298$. Besides, a maximal allowable velocity vector \mathbf{V}^{max} is used to clamp velocities of particles on each dimension,

$$v_{ij} = \begin{cases} -V_j^{max} & v_{ij} < -V_j^{max} \\ v_{ij} & -V_j^{max} \le v_{ij} \le V_j^{max} \\ V_j^{max} & v_{ij} \ge V_j^{max} \end{cases}$$
(4)

here j denotes the j-th dimension of the vector. \mathbf{V}^{max} is set according to the range of the variable space,

$$\mathbf{V}^{max} = \gamma(\mathbf{y}_U - \mathbf{y}_L)$$

In the equation, \mathbf{y}_U and \mathbf{y}_L represent the upper bound and lower bound of the space, respectively [22]. It is found that the optimal setting of γ

is problem dependent. To improve the flexibility of the program, V^{max} is initially set using $\gamma = 0.5$ and then updated according to the actual situation of the running program [22],

$$\mathbf{V}^{max} \Leftarrow \begin{cases} \beta \mathbf{V}^{max} & \text{if solution stagnates for 3 generations} \\ \mathbf{V}^{max} & \text{otherwise} \end{cases}$$
 (5)

here β is set to be 0.95.

The basic PSO initializes particles using uniform pseudo-random numbers within the variable space. The random number sequences do not achieve the lowest possible discrepancy, thus the random points cannot evenly cover the whole space, as illustrated in Fig. 3(a). The discrepancy is used to measure how uniformly distributed the point set is. Various low discrepancy sequences have been proposed. Among them, the Hammersley sequence shows remarkable superiority on ease of implementation [23]. Thus it is adopted to generate initial particles.

Each nonnegative integer i can be expanded by a prime base p,

$$i = a_0 + a_1 p + a_2 p^2 + \dots + a_r p^r$$

with $0 \le a_j < p, j = 1, 2, \dots, r$. A function Φ_p is defined for i as,

$$\Phi_p(i) = \sum_{j=0}^r \frac{a_j}{p^{j+1}}$$

It can be proved that for any $i \geq 0$ and $p \geq 2$, the inequality $0 \leq \Phi_p(i) < 1$ always holds true.

A series of prime numbers $p_1 < p_2 < \cdots < p_{D-1}$ determine a sequence of functions $\{\Phi_{p_1}, \Phi_{p_2}, \cdots, \Phi_{p_{D-1}}\}$. Then a D-dimensional Hammersley point is defined as,

$$\left(\frac{i}{N}, \Phi_{p_1}(i), \cdots, \Phi_{p_{D-1}}(i)\right), i = 1, 2, \cdots, N$$

With p increasing, the point distribution becomes more and more regular [23]. To make the prime numbers as small as possible, we set $p_j = j + 1, j = 1, \dots, D - 1$.

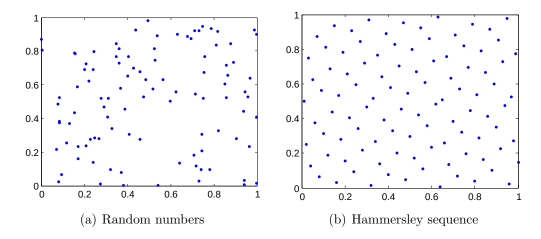


Figure 3: Uniform points generated by random numbers and Hammersley sequence

3.2. Differential evolution

Inspired by the natural evolution of biological species, Holland proposed a popular algorithm called the *genetic algorithm* (GA) [24]. In 1995, Price and Storn replaced the classical crossover and mutation operators in genetic algorithms by a differential operator, which leads to an algorithm called differential evolution (DE) [25]. Compared to GA, DE is more simple but performs better on many numerical optimization problems [25]. Compared to PSO, it can cover the variable space more sufficiently and has greater probability to find the global optimum, *i.e.* it possesses superior capability of exploration.

At each generation, a Donor vector \mathbf{v}_i is generated for each individual of the population (called *genome* or *chromosome*) $\{\mathbf{y}_i|i=1,\cdots,N\}$. It is the method of creating this Donor vector that demarcates between various DE schemes. Two mutation schemes 'DE/rand/1/bin' and 'DE/current to best/2/bin' are applied [26] here,

$$\mathbf{v}_{i} = \begin{cases} \mathbf{y}_{r} + F(\mathbf{y}_{s} - \mathbf{y}_{t}) & \text{rand}[0, 1] (6)$$

where r, s and t are integers randomly selected from the range [1, N] (excluding i). $F \in [0, 2]$ is used to scale the differential vector.

These two strategies are used very commonly in literature and perform well on problems with distinct characteristics. 'DE/rand/1/bin' demon-

strates good diversity while 'DE/current to best/2/bin' shows good convergence property. Here $p \in [0, 1)$ is a user-set parameter.

After the mutation phase, a 'binomial' crossover operation is applied,

$$u_{ij} = \begin{cases} v_{ij} & \text{if } \operatorname{rand}_{j}[0,1] \leq CR \text{ or } j = j_{rand} \\ y_{ij} & \text{otherwise} \end{cases}$$
 (7)

where $CR \in [0, 1)$ is a user-specified crossover constant and j_{rand} is a randomly chosen integer within the range [1, N] to ensure that the trial vector \mathbf{u}_i will differ from \mathbf{y}_i by at least one component. The subscript j refers to the j-th dimension. rand[0, 1] is a random number uniformly generated in the interval [0, 1].

Then a selection operation follows,

$$\mathbf{y}_{i}^{k+1} = \begin{cases} \mathbf{u}_{i}^{k} & \text{if } f(\mathbf{u}_{i}^{k}) < f(\mathbf{y}_{i}^{k}) \\ \mathbf{y}_{i}^{k} & \text{otherwise} \end{cases}$$
(8)

with k and k+1 denoting the individuals at the k-th and (k+1)-th generations, respectively, and f representing the fitness (here it is the cylindricity or conicity error).

The optimal configuration, *i.e.* the values of F, CR and p, is problem-dependent. To obtain relatively good performance in different situations, a self-adaptive DE is employed [27]. The parameters are trained through the optimization process. The training procedure will not be presented here for the sake of concision.

3.3. A two-stage optimization algorithm

The PSO program possesses an advantage of fast convergence, but it has a tendency of premature convergence because of the possibility of overwhelming, *i.e.* sub-optimal groups best influence early in the search. While DE can maintain a greater diversity in the population and the individuals can travel more amply within the variable space, thus the local minimum problem can be overcome more effectively. However DE updates the individuals somehow 'blindly' and finds the optimal region more slowly than PSO. Therefore a two-stage approach is used: first search for the optimal region using PSO and then refine the solution by DE. The criterion adopted for switching between these two stages is the diversity, which is defined as [28]

$$\operatorname{diversity}(\mathbf{Y}) = \frac{1}{N \cdot L} \sum_{i=1}^{N} \sqrt{\sum_{j=1}^{D} (y_{ij} - \bar{y}_j)^2}$$
(9)

where $\overline{\mathbf{y}} = \sum_{i=1}^{N} \mathbf{y}_i / N$ is the centroid of population and L is the length of the diagonal in the search space.

The pseudocode for the two-stage optimization program is shown in Algorithm 1.

3.4. Implementation issues

It is obvious that Eq. (1) has only four independent variables $\{x_0, y_0, m, n\}$ whereas the nominal radius R can vary arbitrarily without altering the cylindricity error. The reason is only the difference between $\max_k e_k$ and $\min_k e_k$ is used but their own values are not taken into account. In practice, the value $\alpha = \max_k e_k / \min_k e_k$ will be supplied in different situations of surface mating. In this paper we enforce the tolerances are symmetric, *i.e.* $\alpha = -1$. As a result the individuals in the program need only four dimensions, rather than five. The radius is enforced R = 0 during optimization and then adjusted as,

$$R \Leftarrow \frac{\max_k e_k + \min_k e_k}{2}$$

For the same reason, the conicity program has only five independent variables, rather than six. The vertex position of the nominal cone cannot be determined through the optimization program. To eliminate the ambiguity in the solution, the vertex is forced located on the X-Y plane, i.e. $z_0=0$ during optimization and then worked out after that. For symmetric tolerances,

$$\begin{cases} x_0 \Leftarrow x_0 - \frac{m(\max_k e_k + \min_k e_k)}{2|\mathbf{n}|\sin\theta} \\ y_0 \Leftarrow y_0 - \frac{n(\max_k e_k + \min_k e_k)}{2|\mathbf{n}|\sin\theta} \\ z_0 \Leftarrow - \frac{\max_k e_k + \min_k e_k}{2|\mathbf{n}|\sin\theta} \end{cases}$$

4. Experimental validation and discussion

To validate the proposed algorithm, four benchmark data sets are used. They are two cylinders from [12] (dataset 2) and [15], and two cones from [16] and [29]. Many researchers compared their own calculation results using these data. The best fitting results found in literature are presented in Table 1.

The two cylinders are checked first. To reveal the reliability of this algorithm, the variable space of the cylinder centre (x_0, y_0) was set a half of the data range, and m and n were generated within [-1, 1]. The number of

```
Input: X
// data points
Initialize population \mathbf{Y} \in \mathbb{R}^{N \times D}, velocities \mathbf{V} \in \mathbb{R}^{N \times D} and \mathbf{V}^{max} \in \mathbb{R}^{D};
k=0; // generation number for PSO
while k < k_1^{max} do
    k + +;
    for i = 1 to N do
        Evaluate f(\mathbf{y}_i);
        Update \mathbf{p}_i and \mathbf{p}_g if possible;
      Adapt \mathbf{v}_i with (3) and (4);
     \mathbf{y}_i \Leftarrow \mathbf{y}_i + \mathbf{v}_i;
    \operatorname{end}
    Update \mathbf{V}^{max} with (5);
    if diversity(\mathbf{Y}) < d_0 then
        Break;
    end
end
Initialize F, CR and p;
k=0 // generation number for DE
while k < k_2^{max} do
    k + +;
    Train F, CR and p;
    for i = 1 to N do
        Mutation and crossover of \mathbf{y}_i with (6) and (7);
        Selection of \mathbf{y}_i with (8);
    end
    if termination condition satisfied then
        Break;
    end
end
Output: p_q
           Algorithm 1: A Two-Stage PSO-DE Algorithm
```

particles was N=20. The fitness value associated with the global optimal solution \mathbf{p}_g at every generation was stored. If the program stagnates for 30 iterations, i.e. $f(\mathbf{p}_g^k) - f(\mathbf{p}_g^{k+30}) < \epsilon$, the program will be terminated. Here it is set $\epsilon = 10^{-6}$.

The program was run 400 times and the minimal, maximal and mean values of the cylindricity errors are listed in Table 1. The standard deviation σ of the 400 errors is applied to measure the uncertainty. It is directly related with the stability of the program. The optimal solutions obtained via this PSO-DE algorithm are listed in Table 2.

It can be seen that the program can find better results than the cited papers. It behaves very stably for the second cylinder and the global optimal solution can always be found. On the contrary, the uncertainty of the first cylinder is a little greater. This is because there are only 20 data points and the data range in the z direction is only a third of the cylinder radius, which makes the evaluation program less stable. But the relative uncertainty of the cylindricity is 0.0000092, which is sufficient for practical applications. In fact, by altering the termination conditions and increasing the particle population, the individuals can search more sufficiently within the variable space and the solution can be made more stable, of course, with a cost of more computation time.

It has to be mentioned that [19] gave form errors 0.174635 and 0.002788 for cylinders 1 and 2, respectively. They introduced a vertex angle in the optimization program, which in turn added one extra degree of freedom in the variable space. Consequently the form errors they gave were actually conicity, rather than 'cylindricity'. The result of cylinder 2 appears correct. It is because the vertex angle is very small, which by chance has little influence on the evaluation result.

Table 1: Form errors of PSO-DE							
data	published	PSO-DE					
		min	max	mean	σ		
Cylinder1	0.18396 [12]	0.183957	0.183991	0.183958	1.69e-6		
Cylinder2	0.002788 [15]	0.002788	0.002788	0.002788	2.800e-9		
Cone1	0.0280 [19]	0.026911	0.027030	0.026918	1.59e-5		
Cone2	0.0032 [11]	0.003154	0.003178	0.003156	4.03e-6		

The conicity evaluation program applied the same settings, except the half vertex angle $0 < \theta < \pi/2$. The minimal, maximal and mean values

Table 2: Best solutions of PSO-DE to cylinders

data	x_0	y_0	m	n	R
Cylinder1	0.010650	0.046918	-0.000619	-0.002915	59.989505
Cylinder2	0.003390	-0.008381	-0.013961	0.007043	12.001374

Table 3: Best solutions of PSO-DE to cones

data	x_0	y_0	z_0	\overline{m}	\overline{n}	θ
Cone1	22.249169	10.087816	-15.768426	9.874e-3	2.337e-3	0.240671
Cone2	8.806048	3.989779	-6.169093	1.497e-3	-5.96e-4	0.239001

as well as standard deviation of 400 results are given in Table 1 and the optimal solutions are shown in Table 3. The superiority of this algorithm is more remarkable for cones. Even the worst results are better than the cited papers. This means those algorithms were trapped at local minima. Another point worth noting is that the vertices of both two cones are above the point sets, so that it is proper to set the direction vector of the axis to be (m, n, -1). If vertices are below, the direction vectors should be (m, n, 1) accordingly, or equivalently make the half vertex angle $-\pi/2 < \theta < 0$. In fact, to make the optimization program more flexible, the variation range of the angle can be set as $-\pi/2 < \theta < \pi/2$, which needs no prior information on the orientation of the cone.

To reveal the behaviour of the algorithm more sufficiently, Two parameters y_0 and n of cylinder 2 were fixed at the optimal solution, and only the other two parameters x_0 and m were altered, as illustrated in Fig. 4(a) and 4(b). The gray background represents the values of the objective function associated with different (x_0, m) . The error map is not smooth. Instead, it contains many 'grooves', which imply local minima. Randomly generating 10 particles in the variable space, Fig. 4(a) and 4(b) illustrate the trajectories of the 10 particles during PSO and DE optimization, respectively. The 'stars' in the graphs denote the global optimal solutions.

It can be seen that the PSO particles assemble toward the optimum very quickly and the program shows good exploitation behaviour, thus a good initial guess is not required. While at the DE phase, the individuals may move along any direction. The optimal region can be searched amply and local minima can be avoided to the greatest extent. As a result the program can make a good balance between the convergence speed (exploitation) and avoiding local minima (exploration).

To demonstrate the reliability of the proposed PSO-DE algorithm clearly, the variable range was set very large. In fact, the measured data can be fitted first with linear least squares [30], so that an approximate solution is obtained. Then the search space of the PSO-DE program can be greatly reduced and the computational efficiency will be improved significantly.

5. Conclusions

This paper presents a reliable method to evaluate the minimum zone form errors of cylinders and cones, which are consistent with the definitions in ISO 1101. A hybrid particle swarm optimization-differential evolution algorithm is proposed, which conducts optimization in two phases. Taking its advantage of fast convergence, particle swarm optimization is firstly implemented from a very large variable space. Then differential evolution is employed to search the optimal region amply, so that local minima can be effectively avoided. To improve the flexibility of the optimization program, the control parameters, e.g. the crossover rate, are trained using a self-adaptive approach. Experimental examples show that this algorithm can achieve higher evaluation accuracy, and obtain better results than literature. It is versatile and can be extended to the minimum zone evaluation of more complex shapes, e.g. ellipsoid, paraboloid and freeform surfaces. Moreover the evaluation criteria can also be minimum circumscribed features or maximum inscribed features, which are suited for different functionality requirements [3, 31]. Therefore, this method could be used for online inspection in high precision manufacturing or form error evaluation on coordinate measuring machines.

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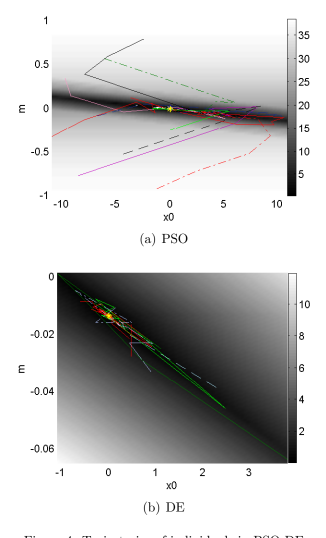


Figure 4: Trajectories of individuals in PSO-DE $\,$