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# Enhancing gene expression programming based on space partition and jump for symbolic regression



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### ABSTRACT

When solving a symbolic regression problem, the gene expression programming (GEP) algorithm could fall into a premature convergence which terminates the optimization process too early, and may only reach a poor local optimum. To address the premature convergence problem of GEP, we propose a novel algorithm named SPJ-GEP, which can maintain the GEP population diversity and improve the accuracy of the GEP search by allowing the population to jump ef ciently between segmented subspaces. SPJ-GEP rst divides the space of mathematical expressions into *k* subspaces that are mutually exclusive. It then creates a subspace selection method that combines the multi-armed bandit and the  $\epsilon$ -greedy strategy to choose a jump subspace. In this way, the analysis is made on the population diversity and the range of the number of subspaces. The analysis results show that SPJ-GEP does not signi cantly increase the computational complexity of time and space than classical GEP methods. Besides, an evaluation is conducted on a set of standard SR benchmarks. The evaluation results show that the proposed SPJ-GEP keeps a higher population diversity and has an enhanced accuracy compared with three baseline GEP methods.

## 1. Introduction

Symbolic regression (**SR**) is a regression analysis that discovers a model that best ts a given dataset in the space of mathematical expressions. Unlike machine learning or neural network regression analysis that focuses on optimizing parameters in a prede ned model, SR aims to nd appropriate models and their parameters at the same time. Genetic programming (**GP**) [1] is a commonly used approach in SR to search for the optimal model. GP evolves to change individual structures of the population to generate tted models or computer programs by the three key genetic algorithm (**GA**) operations: selection, crossover, and mutation. To represent a mathematical expression, classical GPs usually describe individual encodings in trees [1 5]. Graph-based GPs, such as graph encoding GP [6,7] and Cartesian genetic programming [8,9], encode individuals into graphs. Linear GPs, such as gene expression programming (GEP) [10 12] and linear GP [13], convert individuals into linear strings.

Since these GPs all utilize GA operations, like genetic algorithms(GA), these GPs are prone to premature convergence [14]. From the perspective of exploration and exploitation [15], the reason for premature convergence is that individuals of a population are similar, hence, they tend to exploit their neighborhood instead of new regions. Therefore, maintaining the

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population diversity is a crucial task in evolutionary algorithms. A diverse population can encourage global exploration and reduce premature convergence [16,17].

In order to preserve the population diversity, the evolutionary computing (EC) community often uses two strategies: 1) parameter control and 2) space partition. The parameter control strategy [18] adjusts parameters of evolutionary algorithms based on population diversity, such as varying population size [19,20], and dynamically adjusting the probability of cross-over [21,22] and mutation [23,24]. The strategy is easy to implement population diversity and does not require additional storage spaces. However, it does not know or remember where individuals are in a search space so that it could produce invalid individuals, such as individuals similar to those of the previous generations.

The space partition strategy [25 30] splits a search space into many subspaces and generates individuals in different subspaces. As individuals in different subspaces have different phenotypes or genotypes, the strategy is easy to control population diversity quantitatively by generating individuals from different subspaces. Meanwhile, the strategy remembers an individual's approximate position in the search space according to the individual's subspace. Although the space partition strategy has been successfully applied in GA, it is not suitable for the SR problem, because the whole search space of SR is so large that maintaining ne-grained subspaces is intractable computationally.

In this paper, we propose a new gene expression programming based on **s**pace **p**artition and **j**ump (named **SPJ-GEP**) to maintain the population diversity. SPJ-GEP has the advantages of the above two strategies: it requires small additional storage space, remembers the position of an individual in the search space, and maintains quantitative population diversity. The SPJ-GEP partitions the space of mathematical expressions into *k* subspaces based on the chromosome coding. Moreover, it initializes individuals in one of the *k* subspaces, as shown in Step 1 in Fig. 1.

Next, SPJ-GEP selects a suitable subspace to search for individuals with better tnesses based on a subspace selection method that combines the multi-armed bandit (**MAB**) [31] and the  $\epsilon$ -greedy strategy [32], as shown in Step 2 and 3 in Fig. 1. This method utilizes MAB to choose one of the subspaces because MAB can balance the exploration by searching other subspaces while maintaining the exploitation of the selected subspace. However, MAB will be invalid when the number of visiting subspaces is higher than a speci c value. To preserve population diversity, the method then switches to the  $\epsilon$ -greedy strategy to choose a subspace according to a proposed time formula. The formula decides when to use the  $\epsilon$ -greedy strategy.

At last, SPJ-GEP uses a new crossover method to make individuals jump from the original subspace to another selected subspace, as shown in Step 4 in Fig. 1. The method makes these newly selected individuals intersect with the best individual in the selected subspace so that they can start searching at the latest local optimal position.

The characteristics of SPJ-GEP indicates that classical GEPs [10 12] are a special case of SPJ-GEP when the number of subspaces k equals 1. On the other hand, if k is large enough that each subspace has only one individual, SPJ-GEP will degenerate into a random selection subspace algorithm. Therefore, k is a critical parameter in SPJ-GEP. In this paper, the range of k is decided by the population diversity and the probability of jump between subspaces. We analyze the complexity of time and space of SPJ-GEP and prove that SPJ-GEP does not signi cantly increase the time and space complexity compared with classical GEPs.

The main contributions in the paper are summarized as follows:

 We propose the SPJ-GEP algorithm, which allows individuals in a population to jump between subspaces according to the MAB and the *ϵ*-greedy strategy. This approach maintains the population diversity.

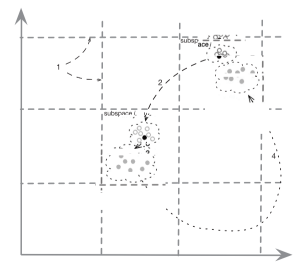
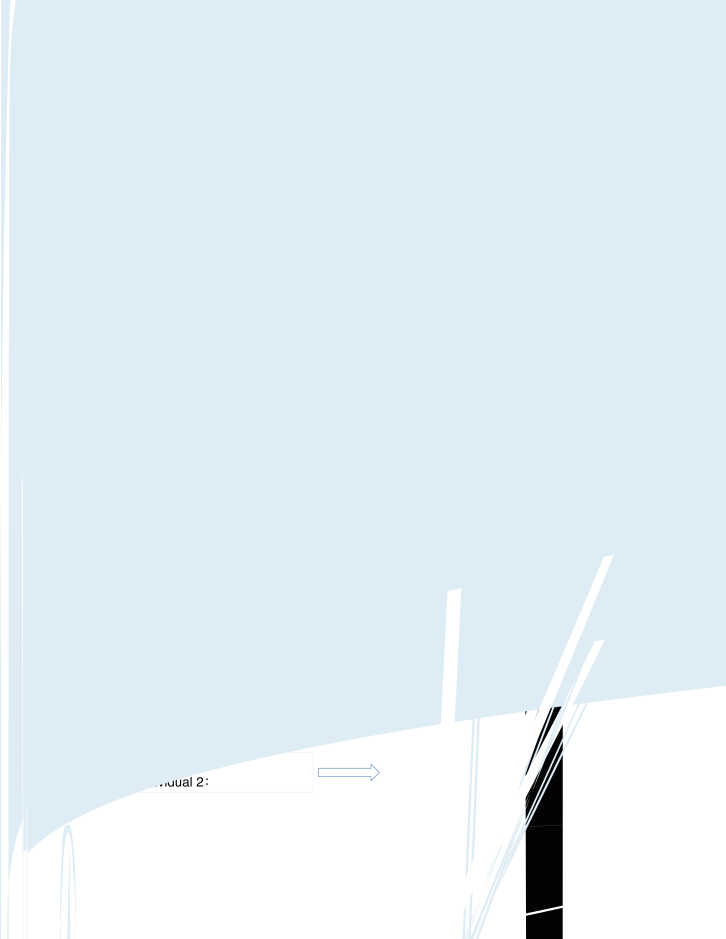


Fig. 1. The SPJ-GEP framework. Circles represent individuals, and dark circles are the best individuals in subspaces. Step 1. space segmentation; Step 2. subspace selection; Step 3. subspace exploitation; Step 4. escape from local optimum by subspace jump.





According to the coding of individuals in GEP, i.e., encoding an individual to a linear structure with the xed-length *l* and the head length *h*, the space of mathematical expression is denoted as  $\Omega_{l,h} = `* \dots ** \dots *'$ . *l* is the length of individuals (the total number of '\*'); *h* is the head length (the number of '<u>\*</u>'); '\*' can be anyone symbol from a symbol set *S* that consists of a function set *F* and a terminal set *T*. In the head, if front '\*'s are replaced by special symbols *s*, it can generate a **subspace**  $\omega_s$ , such as  $\omega_+ = `+ ***'$  in Fig. 3. Therefore, the more special symbols appear in the front of the head, the smaller the subspace's size becomes. For example,  $\omega_{++} = `++ ***'$  is a subspace of  $\omega_+$ , and  $\omega_{++} \subset \omega_+$ .

Based on the above subspace encoding, these subspaces and their relationships can be represented as **a space-partition tree**, where the root node is  $\Omega_{l,h}$ , each of the other nodes is a subspace of  $\Omega_{l,h}$ , and a branch represents a containment relationship between two subspaces, e.g.,  $\omega_{++} \subset \omega_{+}$  as shown in Fig. 3. From the tree, a lot of space-partition sets can be found based on the above two conditions. For example, { $\omega_{+}, \omega_{-}, \omega_{x}$ } and { $\omega_{++}, \omega_{+-}, \omega_{+x}, \omega_{+}, \omega_{x}$ } both are space-partition sets.

# 3.1.2. Initialization

SPJ-GEP selects a space-partition set by the following strategy. It can easily nd the rst level *ll* where the number of nodes is equal or greater than the number of subspaces (*k*) according to  $ll \ge log_{|S|}^k$ , where |S| is the number of symbols. For example, in Fig. 3, if k = 5 and |S| = 3, then ll = 2. Moreover, the algorithm discovers a space-partition set  $\omega_{++}, \omega_{+-}, \omega_{+x}, \omega_{-+}$ .

$$UCB_{\omega_i} = \frac{1}{f_{\omega_i}^* + 1} + \lambda \sqrt{\frac{2lnt}{n_{\omega_i}}}$$
(2)

where  $\omega_i$  is a subspace,  $f_{\omega_i}^*$  is the tness of the best individual in  $\omega_i$ , t is the number of visiting  $\Omega$  until a particular time, and  $n_{\omega_i}$  is the number of times that the subspace  $\omega_i$  is accessed to. Then, SPJ-GEP selects the best subspace  $\omega_i^*$  that has the maximal *UCB* as its exploration space.

### 3.2.2. Subspace selection based on $\epsilon$ -Greedy method

As visit times increase in a subspace, the size of the condence interval  $(\sqrt{\frac{2lnt}{n_i}}$  in Eq. 2) decrease to zero. That means  $UCB_{\omega_i}$ 

falls back to a greedy method with the subspace value  $(f_{\omega_i}^*)$  and becomes invalid in the balance between exploration and exploitation.

To overcome the above invalidation, SPJ-GEP uses the  $\epsilon$ -greedy method [35] to select a subspace when condence intervals in most of the subspaces tend to be zero. Using Eq. 2 chooses a subspace with the probability  $1 - \epsilon$ ; random chooses a subspace from the above with the probability  $\epsilon$ .

#### 3.2.3. The time of using the $\epsilon$ -Greedy Method

To nd out when  $UCB_{\omega_i}$  (Eq. 2) loses its effect on most subspaces, SPJ-GEP uses Eq. 3 in the following Theorem 1. For example, given  $k = 300, n = 100, G = 100000, \alpha = 0.1$  and  $\beta = 0.8$ , to choose a subspace  $\omega_i$ , it uses Eq. 2 if  $t < -\frac{2 \times 300 \times \ln^7}{0.01^2 \times \ln^{0.6}} \approx 4333918$ , where  $T = n \times G = 100 \times 10000 = 10^7$ ; otherwise, it uses  $\epsilon$ -greedy method.

**Theorem 1.** Let *k* be the number of subspaces, and T be the total number of times of visiting the mathematical expressions space  $\Omega$  after running SPJ-GEP. If the confidence interval  $\sqrt{\frac{2\ln^i}{n_{\omega_i}}} \leq \alpha$ , where  $\alpha \to 0^+$ , UCB<sub> $\omega_i$ </sub> will lose its effect on subspace  $\omega_i$ . Assuming that different  $n_{\omega_i}$  is independent identically distributed (i.i.d.), and each  $n_{\omega_i}$  is an exponential distribution with the parameter  $\lambda$ , the probability that each UCB<sub> $\omega_i$ </sub> loses its effect is greater than  $\beta$  when the number of times that  $\Omega$  is accessed to.

$$t \ge -\frac{2kln^{T}}{\alpha^{2}ln^{\beta}}.$$
(3)

**Proof.** According to the constant *T* and  $\sqrt{\frac{2ln^T}{n_{\omega_i}}} \leq \alpha$ , we have  $n_{\omega_i} \geq \frac{2ln^T}{\alpha^2}$  so that  $UCB_{\omega_i}$  are invalid in  $\omega_i$ . Therefore, if we want to assure that the probability, which each  $UCB_{\omega_i}$  loses its effect, is greater than  $\beta$ , i.e.,  $P(n_{\omega_i} \geq \frac{2ln^T}{\alpha^2}) \geq \beta$ , the following equation

$$\lambda \leqslant -\frac{\alpha^2 \ln^{\beta}}{2 \ln^{T}} \tag{4}$$

must be satis ed, because

$$\begin{split} P(n_{\omega_i} \geq \frac{2\ln^7}{\alpha^2}) &= \int_{\frac{2\ln^7}{\alpha^2}}^{\infty} \lambda e^{-\lambda n_{\omega_i}} dn_{\omega_i} = e^{-\lambda \frac{2\ln^7}{\alpha^2}} \\ &\Rightarrow e^{-\lambda \frac{2\ln^7}{\alpha^2}} \geq \beta. \end{split}$$

As different  $n_{\omega_i}$  is i.i.d. and each  $n_{\omega_i}$  is an exponential distribution with the parameter  $\lambda$ , we get

$$t = E[\sum_{i=1}^{k} n_{\omega_i}] = E[E[\sum_{i=1}^{k} n_{\omega_i} | k]] = E[k]E[n_{\omega_i}] = \frac{k}{\lambda}.$$
(5)

According to formulas 4 and 5, we nally obtain formula 3.

### 3.3. Exploitation with crossover

Suppose that the current population is at the subspace  $\omega_j$ , after SPJ-GEP selects a subspace  $\omega_i$ , it makes all individuals in the population **jump** from the subspace  $\omega_j$  to the subspace  $\omega_i$ . As the aforementioned subspace encoding, codes of these individuals in  $\omega_j$  start with the code of  $\omega_j$ . So, for making them jump, it is necessary to replace their head codes with the code of  $\omega_i$ . For example, given two individuals ' $\pm\pm/$  – ×*xxxxxx*' and ' $\pm\pm$  + × × *xxxxxx*' in the subspace  $\omega_{++}$ , the two individuals will jump into the subspace  $\omega_{+-}$  after they change to ' $\pm-$  – ×*xxxxxx*' and ' $\pm-$  + × × *xxxxxx*' by replacing '++' in their heads with the code '+-' of ' $\omega_{+-}$ '.

Then, it exploits  $\omega_i$  by recombining each of the transferred individuals with the best individual in the subspace  $\omega_i$ . For example, if the best individual is '<u>+-</u> × + + xxxxxx', the above two individuals '<u>+-</u> - ×xxxxxx' and '<u>+-</u> + × ×xxxxx' recom-

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bine with the best individual, respectively. The recombination makes the jumped population start to search from the local optimal space, and speeds up the convergence.

Note that if the selected subspace  $\omega_j$  is equal to  $\omega_i$ , the recombination is the same as the recombination of classical GEPs [10 12] in that any two individuals in the population recombine randomly. So, if the above subspace selection continuously chooses the same subspace, SPJ-GEP will exploit the subspace persistently.

## 4. Analysis of SPJ-GEP

## 4.1. Time and space complexity

Compared with classical GEPs [33,37,38], SPJ-GEP requires additional structures to record the visiting times  $(n_{\omega_i})$  and the best tness  $(f_{\omega_i}^*)$ , as well as extra computation to obtain  $UCB_{\omega_i}$  in each subspace. The additional time and space complexity are related to the number (k) of all subspaces. Suppose the time and space complexities for classical GEPs within g iterations are O(gep) and  $\Theta(gep)$ , respectively. For SPJ-GEP, they are  $O(gep + g \times c \times k) = O(gep + n \times k)$  and  $\Theta(gep + m \times k)$ , where  $n = c \times k, k > 1$ , and g, c, and m are constants. Therefore, if k is within a reasonable range, its time and space complexities are acceptable. In our experimental evaluation, its running time is almost as fast as GEP's because the value of k is generally not very large (detail in Section 4.3).

#### 4.2. Population diversity

To preserve the population diversity, SPJ-GEP always lets the population jump from one subspace to another subspace. Even if an individual in a population immediately jumps back to its original subspace after two jumps, its structure has been signi cantly changed according to the following Lemma 1.

**Lemma 1.** Suppose SPJ-GEP executes single-point crossovers with uniform distribution, and l is the length of an individual encoding without considering its subspace encoding. After the individual has jumped k subspaces, the similarity between the jumped individual, and the original individual is

$$sim(k) = \left(\frac{1+l}{2l}\right)^{\kappa}$$
(6)

**Proof.** Since the crossover point is randomly selected with uniform distribution, after a crossover, the expected length of the original fragment in the new individual is (1 + l)/2. Then, the similarity between the original individual and the jumped individual is (1 + l)/2l after one crossover. Therefore, after *k* crossovers in *k* subspaces, the similarity is  $sim(k) = (\frac{1+l}{2l})^k$ .

For example, if the length of an individual is 20, after jumping only two subspaces, the similarity between original and jumped individuals changes to 0.276. As k increases, sim(k) tends to be zero. Moreover, the two individuals become more different. Therefore, the subspace selection method, which lets the population jump from a subspace to another subspace, diversi es the population so that it helps prevent a local optimum in SPJ-GEP.

## 4.3. The number of subspaces

The number of subspaces k is a critical parameter in SPJ-GEP. If k = 1, it means that there is only one subspace in  $\Omega$ . So, when the subspace is  $\Omega$ , SPJ-GEP degenerates into a standard GEP. If k is large enough that there is only one individual in a subspace, each subspace is an individual. In this case, SPJ-GEP degenerates into the random initialization algorithm, randomly generating an individual (i.e., a subspace) in  $\Omega$ . Therefore, if k is too large or too small, the algorithm performance will be degraded.

According to Lemma 1, the larger *k*, the smaller the similarity. Then, SPJ-GEP needs a smaller similarity to escape from the local optimum. Based on the following Theorem 2, we have the lower bound of *k*. Besides, the larger *k*, the smaller the probability that an individual jumps in its original subspace. The probability must be larger enough that SPJ-GEP can exploit a subspace continuously for a while. Otherwise, SPJ-GEP will always explore a different subspace that breaks the balance between exploration and exploitation. So, based on the following Theorem 3, we have the implicit expression of the upper bound of *k*.

**Theorem 2.** Suppose an individual returns to the original subspace after it jumps k subspaces, in order to guarantee the similarity between the original individual and the new back individual is less than or equal to  $\eta$ , the lower bound of k is  $\frac{\ln^{\eta}}{\ln^{1+1}\ln^{2t}}$ .

**Proof.** According to Eq. 6 in Lemma 1,  $\left(\frac{1+l}{2l}\right)^k \leq \eta$ . Then, we have

$$k \ge \frac{\ln^{\eta}}{\ln^{1+l} - \ln^{2l}} \tag{7}$$

When the running time of SPJ-SEP exceeds a speci c time, SPJ-SEP uses the  $\epsilon$ -greedy strategy to select a subspace according to Theorem 1. Most of the subspaces are selected randomly. So, the jump probability P(i,j) between subspace i and j satis es a long tail distribution. Moreover, the probability that the subspace with the best  $UCB_{\omega_i}$  will be selected again is  $1 - \epsilon$ , and the probability is higher than the probability of selecting other subspace, i.e., P(i, i) > P(i, j). If SPJ-SEP uses the best  $UCB_{\omega_i}$  (MAB) to select a subspace, the same conclusions P(i, j) satis es a long tail distribution and P(i, i) > P(i, j) can be obtained through an analysis similar to the above.

**Theorem 3.** Suppose the jump probability P(i, j) between subspace *i* and *j* satisfies the Zipf distribution [39], in order to guarantee that  $P(i, i) \ge \delta$ , the number of subspace *k* satisfies the following inequality.

$$\frac{1}{\sum_{i=1}^{k} \left(\frac{1}{i}\right)^{\gamma}} \ge \delta$$
(8)

**Proof.** Since  $P(x_{i,j})$  satis es Zipf distribution, whose probability mass function of Zipf is  $f(x) = \frac{1}{x^{\gamma} \sum_{i=1}^{k} (1/i)^{\gamma}}$ , where x = 1, 2, ..., k, k is the number of subspace, and  $\gamma$  is a parameter.

$$p(i,i) = f(1) = \frac{1}{\sum_{i=1}^{k} (1/i)^{\gamma}} \ge \delta$$
(9)

owing to f(1) > f(x) when  $x \neq 1$ .

Table 2 GP Problems.

For example, if  $\gamma = 0$ , the Zipf distribution will degenerate into a uniform distribution. According to Eq. 8,  $\frac{1}{k} \ge \gamma$ . So,  $k \in \left[\frac{\ln^{\eta}}{\ln^{1+1}-\ln^{21}}, \frac{1}{\gamma}\right]$ .

Name Formula Dataset  $x^6 + x^5 + x^4 + x^3 + x^2 + x$ F1 U[-1, 1, 20] F2  $x^4 + x^3 + x^2 + x$ U[-1, 1, 20] U[-1, 1, 20] F3  $x^5 - 2x^3 + x$ F4 U[-1, 1, 20]  $sin(x^2)cos(x) - 1$ F5 U[-1, 1, 20]  $sin(x)+sin(x+x^2)$ F6  $ln(x+1) + ln(x^2+1)$ U[0, 2, 20] F7 2sin(x)cos(y)U[-1, 1, 100] F8 U[-50, 50, 10000] 1.57 + (24.3v)F9  $6.87 + 11\cos(7.23x^3)$ U[-50, 50, 10000] F10 U[-50, 50, 10000]  $2 - 2.1\cos(9.8x)\sin(1.3w)$ F11 0.3xsin(2 - x)E[-1, 1, 0.1] F12 lnx E[1.1] U[0, 1, 100] F13 xy F14  $x^4 - x^3 + \frac{y^2}{2} - y$ U[-3, 3, 20] F15  $\frac{x^3}{5} + \frac{y^3}{2} - y - x$ U[-3, 3, 20] E[0.05, 10, 0.1] F16  $e^{-x}x^3(cosxsinx)(cosxsin^2x-1)$ U[0.3, 4, 100] F17  $\frac{e^{-(x-1)^2}}{1.2+(y-2.5)^2}$ F18 x:E[0.05, 10, 0.1]  $e^{-x}x^3(cosxsinx)(cosxsin^2x-1)$ y:E[0.05, 10.05, 2] \*(y - 5)F19 (x-3)(y-3)+2sin((x-4)(y-4))U[0.05, 6.05, 300] F20  $\tfrac{(x-3)^4+(y-3)^3-(y-3)}{(y-2)^4+10}$ U[0.05, 6.05, 50]

The function sets of  $F_1 - F_7$ ,  $F_8 - F_{10}$ ,  $F_{11} - F_{15}$  and  $F_{16} - F_{20}$  are from Koza [1], Korns [41], Keijzer [42] and Vladislavleva [43], respectively.

## 5. Experiments

# 5.1. Dataset and experimental parameters

In this paper, the dataset consists of 20 SR test problems that are derived from the GP benchmarks [40], as shown in Table 2. The functions and constants of the data set are shown in Table 3. To evaluate the proposed algorithm **SPJ-GEP**, we have created three algorithms SPJ-GEP, SPJ-GEP-ADF, and SPJ-SL-GEP based on the three baseline **GEPs**: GEP [10], GEP-ADF [11], and SL-GEP [12], respectively. The three new algorithms have the same parameters as these GEPs have except for the additional parameters k,  $\alpha$ , and  $\beta$ . The detailed parameters of the above six algorithms are described in Table 4.

We set the number of subspaces *k* a particular value according to the number of nodes on a speci c layer in the spacepartition tree. Since the evaluation consists of six basic benchmarks, and each basic benchmark has different function symbols and terminal symbols, SPJ-GEP, SPJ-GEP-ADF, and SPJ-SL-GEP have different *k* values for different basic benchmarks. In Table 4, the last row shows the range of *k* is [144 1000]. For example, in the Koza basic benchmark, there is a function symbol set  $\{+, -, \times, /, sin, cos, ln(|x|), e^x\}$ , whose length is 8, and a terminal set  $\{x_1, x_2\}$ , whose length is 2. When a space-partition set is obtained by nodes on the 3rd layer in the space-partition tree, the number of the subspaces *k* is  $(8 + 2)^3 = 1000$ .

# 5.2. Verification of subspaces selection method

Looking back at the subspace selection in SPJ-GEP, Inequality 3 is a key to decide when the UCB (Eq. 2) loses its effect on selecting subspaces. To verify the accuracy and the correctness of Inequality 3, we run SPJ-GEP with different values of parameter k (100,200 and 300) on ve test problems. When  $t \ge -\frac{2k\ln^7}{\alpha^2 \ln^6}$ , where  $T = 100000 \times 100$ , such as  $t \ge \frac{2 \times 100 \times \ln^{10^7}}{(0.1)^4 \times \ln^{0.8}} \approx 1444639$ , the algorithm stops. Then, we sum up the number of subspaces where UCB has lost its effect according to the following Inequality

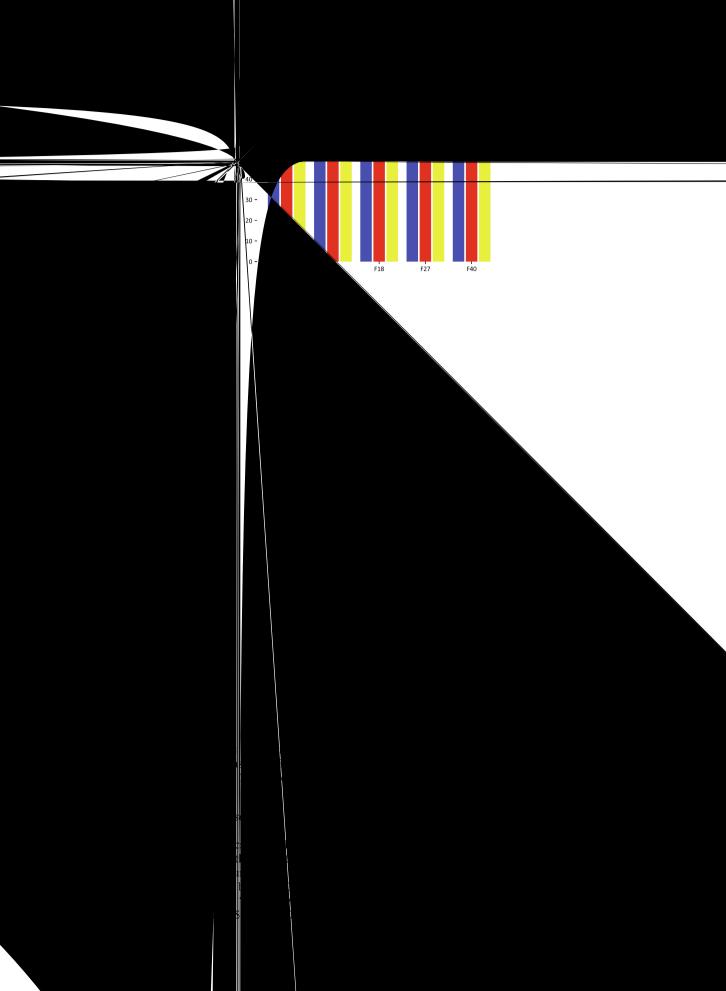
$$\sqrt{\frac{2ln^{T}}{n_{\omega_{i}}}} \leqslant \alpha \tag{10}$$

where  $n_{\omega_i}$  is the number of times that the subspace  $\omega_i$  is accessed to. Finally obtain the probability of subspace convergence (PSC) by Eq. 11

$$PSC = \frac{s}{k} \tag{11}$$

Table 3The Functions and Constants of Data Set.

Name	Functions	Constants(ERC)
Koza Korns	$+, -, \times, /, sin, cos, e^n, ln( n ) +, -, \times, /, sin, cos, e^n, ln( n ) n^2, n^3, \overline{n}$	None Random nite 64bit

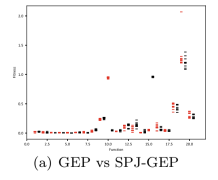


than GEP-ADF. Although SL-GEP and SPJ-SL-GEP nd the same nun average tness value than SL-GEP.

In Table 6, the results of the Wilcoxon Signed-Rank Test[44] at According to the comparative data, SPJ-GEP, SPJ-GEP-ADF, and SPJ GEP, GEP-ADF, and SL-GEP, respectively.

All best tnesses found by the above six algorithms are shown problems, the range of tness obtained by SPJ-GEP, SPJ-GEP-ADF, by GEP, GEP-ADF, or SL-GEP. These results show that SPJ-GEP can algorithms.

However, SPJ-GEP is not superior to GEP, GEP-ADF or SL-GEP i performance of GEP, GEP-ADF or SL-GEP is better than SPJ-GEP. Fo of GEP tness value both are smaller than those of SPJ-GEP tnes GEP) tness value is smaller than that of SPJ-GEP-ADF (SPJ-SL-G the mathematical expression space is split into many subspaces,



rrect results, SPJ-SL-GEP can still obtain a smaller

It considers both the average tnesses and RMSE. still obtain a better performance than the original

s in Fig. 5. The results indicate that, in most of the .-GEP is smaller than the range of tness obtained more accurate performance than the baseline GEP

e tested problems. For a few special problems, the ple, as shown in Fig. 6(a), the average and the range ; similarly, in Fig. 6(h), the range of GEP-ADF (SLat is because, from the subspace's view, although ze of each subspace is still huge so that GEP, GEP- -

0.05-

0.04-

0.03

0.02-

0.01-

0.00-

ADF, and SL-GEP may only search in one or several subspaces. Moreover, the crossover and the mutation of SPJ-GEP (SPJ-GEP-ADF or SPJ-SL-GEP) are the same as those of GEP (GEP-ADF or SL-GEP) in a subspace. Those make it possible for the baseline GEPs to nd more accurate results than these SPJ-GEP algorithms if the subspace that they search for is the target subspace in which the correct results are located.

## 5.4. Comparison of population diversity

Since the crossover and the mutation of SPJ-GEP are the same as those of the baseline GEPs in a subspace, what makes SPJ-GEP obtain more accurate results than these GEPs in most of the problems? It is attributed to the SPJ-GEP's ability to jump between subspaces, which maintains population diversity. To observe the population diversity, the different degree (dg) of individuals is dened by the following equation.

$$dg = \frac{D}{N} \tag{12}$$

where *N* is the number of individuals in a population, and *D* is the number of different tness values of individuals. For example, if 100 individuals generate 70 different tness values, the different degree is 0.7. Therefore, *dg* can represent the population diversity, and *dg* is computed at every 100 generations in the above algorithms. By collecting *dg* values of the above six algorithms that run in the test problem – F11, the population diversity changes of these algorithms are shown in Fig. 7.

As shown in Fig. 7, GEP, GEP-ADF, and SL-GEP have a lower level of the population diversity, while SPJ-GEP. SPJ-GEP-ADF and SPJ-SL-GEP have a higher level of the population diversity. Besides, the average amplitude of blue curves that represent these SPJ-GEPs is higher than that of black curves that represent the three baseline GEPs. Although crossover and mutation contribute to maintaining population diversity in the early stages of these GEPs running, they tend to make little differences between individuals in the later stages. However, the jump always makes signi cant differences between individuals in SPJ-GEPs, according to Lemma 1.

#### 5.5. Convergence comparison

Fig. 8(a) and (b) illustrate that SPJ-GEPs (SPJ-GEP, SPJ-GEP-ADF, and SPJ-SL-GEP) can obtain better results than the baseline GEPs(GEP, GEP-ADF, and SL-GEP). As the number of iterations increases, the baseline GEPs gradually fall into local optimum space. Meanwhile, owing to the low population diversity, they cannot escape from the local optimum space with a high probability. However, SPJ-GEPs can easily escape from a local optimum space because the jump between subspaces transfers individuals to different subspaces and maintains a high population diversity. Therefore, when the baseline GEPs present premature convergence, SPJ-GEPs still nd new better results even in the later stage of 100,000 generations. That demonstrates that SPJ-GEPs can always explore new subspaces with a high probability.

However, owing to the jump between subspaces and the above subspace selection method, it is dif cult for SPJ-GEPs to exploit a local space continuously in a period. Therefore, the convergence speed of SPJ-GEPs is slower than that of these base-line GEPs in the early stage of 100,000 generations. For example, in Fig. 8(a), at approximately 40,500 generations, GEP nds the best result with the tness 0.025, while SPJ-GEP nds that with 0.078.

# 6. Related work

Similar to our idea of space partition that maintains the population diversity in this study, a method named NrGA was proposed to use a binary space partitioning (BSP) tree. NrGA recursively subdivides space into two and stores individual visiting information, and the method is integrated with GA so that individual revisits are completely eliminated [25 27]. Although NrGA can maintain the population diversity by visiting the BSP tree, it needs a lot of additional time and space to maintain the tree. Especially for the huge space of mathematical expressions in SR, maintaining the BSP tree will become an impossible mission. However, in the paper, SPJ-GEP does not construct a space-partition tree but utilizes its abstract structure to obtain a space-partition set. Moreover, SPJ-GEP maintains population diversity by letting individuals jump between subspaces instead of eliminating revisiting individuals.

Many distributed evolutionary algorithms (dEA) use a similar idea of space partition to distribute individuals of the population to different subspaces (multiple processors or computing nodes) [28]. These dEAs can be divided into island [29] [45],

cellular [46] [30], hierarchical [47], and pool models [48]. These dEAs can increase the population diversity because their subpopulations run independently in different subspaces, and transfer their best individuals by a migration strategy. As they are distributed algorithms, they pay more attention to the communication cost and scalability and pay no attention to the subspace selection method, so that they could waste a lot of computation time on invalid subspaces.

Different from the above methods of partitioning global space, Tsutsui [49] and Huang [50] proposed two methods of local spaces that are created by the convergence status of the present population respectively. Tsutsui [49] proposed the forking GA (fGA) which divides the search space for each population into subspaces depending on the convergence status of the population and the solutions obtained so far. Then two types of fGAs (genotypic fGA and phenotypic fGA) are created to maintain population diversity by de ning two searching subspaces of each sub-populations, respectively. One is the salient schema which de nes subspaces by phenotype parameters of the present population. The other is the neighborhood hypercube which de nes the local subspaces around the current best individual in the phenotypic feature space. The forking GA can avoid the premature convergence of populations because the searching method enables the population to exploit different local subspaces. Although Huang [50] proposed a differential evolution (DE) method based on the three spaces: local space, opposition space, and global space, the three spaces are all local spaces because the global space refers to the space near the best individual in a population, and its opposition space. So, the method can accelerate convergence but cannot avoid falling into a local optimum.

## 7. Conclusion and future work

In the paper, we propose a novel algorithm, SPJ-GEP, to deal with the SR problem. Using the new approach that partitions the space of mathematical expressions into subspaces, SPJ-GEP guides the population effectively jump among these subspaces with a subspace selection method. SPJ-GEP maintains the population diversity while keeping the balance between subspace exploration and exploitation. Therefore, the proposed SPJ-GEP has the following advantages. SPJ-GEP can be easily embedded in other GEPs because its three key components space partition, subspace selection, and crossover, are compatible with other GEPs. As shown in the evaluation analysis, SPJ-GEP does not signi cantly increase the time and space complexity compared with classical GEPs. SPJ-GEP can overcome the problem of premature convergence and avoid falling into a local optimum.

Although SPJ-GEP surpasses the tested baseline GEPs on most benchmarks, it has two weaknesses that prevent it from quickly nding better results than the baseline GEPs on a few benchmarks. One is that SPJ-GEP does not de ne how to choose the best space-partition set from the space-partition tree. After all, the quality of the set directly affects the search results of SPJ-GEP. The other weakness is that the selected subspace may not be the subspace where the optimal result is located. In this case, the jump to these selected subspaces could result in an unreasonable search. In the future, we will address these weaknesses by quantifying these subspaces.

### **CRediT** authorship contribution statement

**Qiang Lu:** Conceptualization, Methodology, Validation, Writing - original draft, Writing - review & editing. **Shuo Zhou:** Formal analysis, Data curation, Software, Visualization. **Fan Tao:** Visualization, Validation. **Jake Luo:** Writing - review & editing. **Zhiguang Wang:** Supervision.

#### **Declaration of Competing Interest**

The authors declare that they have no known competing nancial interests or personal relationships that could have appeared to induce the work reported in this paper.

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