### **Crossover Techniques in GAs**

#### **Debasis Samanta**

Indian Institute of Technology Kharagpur

dsamanta@iitkgp.ac.in

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### Encoding

- Pitness Evaluation and Selection
- Mating pool
- Crossover
- Mutation
- Inversion
- Onvergence test

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### Encoding

- Pitness evaluation and Selection
- Mating pool
- Crossover

### Mutation

- Inversion
- Convergence test

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#### **Reproduction:**

- Crossover
- Mutation
- Inversion

These genetic operators varies from one encoding scheme to another.

- Binary coded GAs
- Real-coded GAs
- Tree-coded GAs

- A mating pair (each pair consists of two strings) are selected at random. Thus, if the size of mating pool is *N*, then <sup>N</sup>/<sub>2</sub> mating pairs are formed.[Random Mating]
- The pairs are checked, whether they will participate in reproduction or not by tossing a coin, whose probability being *p<sub>c</sub>*. If *p<sub>c</sub>* is head, then the parent will participate in reproduction. Otherwise, they will remain intact in the population.

#### Note :

Generally,  $p_c = 1.0$ , so that almost all the parents can participate in production.

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Once, a pool of mating pair are selected, they undergo through crossover operations.

- In crossover, there is an exchange of properties between two parents and as a result of which two offspring solutions are produced.
- The crossover point(s) (also called k-point(s)) is(are) decided using a random number generator generating integer(s) in between 1 and *L*, where *L* is the length of the chromosome.
- Then we perform exchange of gene values with respect to the k-point(s)

There are many exchange mechanisms and hence crossover strategies.

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### **Crossover Techniques in Binary Coded GA**

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### **Crossover operations in Binary-coded GAs**

- There exists a large number of crossover schemes, few important of them are listed in the following.
  - Single point crossover
  - Two-point crossover
  - Multi-point crossover (also called n-point crossover)
  - Uniform crossover (UX)
  - Half-uniform crossover (HUX)
  - Shuffle crossover
  - Matrix crossover (Tow-dimensional crossover)
  - Three parent crossover

- Here, we select the K-point lying between 1 and L. Let it be k.
- A single crossover point at k on both parent's strings is selected.
- All data beyond that point in either string is swapped between the two parents.
- The resulting strings are the chromosomes of the offsprings produced.

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### Single point crossover: Illustration



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- In this scheme, we select two different crossover points  $k_1$  and  $k_2$  lying between 1 and *L* at random such that  $k_1 \neq k_2$ .
- In the middle parts are swapped between the two strings.
- Iternatively, left and right parts also can be swapped.

### **Two-point crossover: Illustration**



- In case of multi-point crossover, a number of crossover points are selected along the length of the string, at random.
- 2 The bits lying between alternate pairs of sites are then swapped.



- Uniform crossover is a more general version of the multi-point crossover.
- In this scheme, at each bit position of the parent string, we toss a coin (with a certain probability *p<sub>s</sub>*) to determine whether there will be swap of the bits or not.
- The two bits are then swapped or remain unaltered, accordingly.

### **Uniform crossover (UX): Illustration**



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- Here, each gene is created in the offspring by copying the corresponding gene from one or the other parent chosen according to a random generated binary crossover mask of the same length as the chromosome.
- Where there is a 1 in the mask, the gene is copied from the first parent
- Where there is a 0 in the mask, the gene is copied from the second parent.
- The reverse is followed to create another offsprings.

# Uniform crossover with crossover mask: Illustration



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- In the half uniform crossover scheme, exactly half of the non-matching bits are swapped.
  - Calculate the Hamming distance (the number of differing bits) between the given parents.
  - 2 This number is then divided by two.
  - The resulting number is how many of the bits that do not match between the two parents will be swapped but probabilistically.
  - Choose the locations of these half numbers (with some strategies, say coin tossing) and swap them.

### Half-uniform crossover: Illustration



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- A single crossover point is selected. It divides a chromosome into two parts called schema.
- In both parents, genes are shuffled in each schema. Follow some strategy for shuflling bits
- Schemas are exchanged to create offspring (as in single crossover)

### Shuffle crossover: Illustration

Before crossover



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### Matrix crossover

The matrix crossover strategy is expained with the following illustration.



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- In this techniques, three parents are randomly chosen.
- Each bit of the first parent is compared with the bit of the second parent.
- If both are the same, the bit is taken for the offspring.
- Otherwise, the bit from the third parent is taken for the offspring.

### Three parent crossover: Illustration

P1:	1	1	0	1	0	0	0	1
P2:	0	1	1	0	1	0	0	1
P3:	0	1	1	0	1	1	0	1
C1:	0	1	1	0	1	0	0	1

Note: Sometime, the third parent can be taken as the crossover mask.

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#### Non-uniform variation:

It can not combine all possible schemas (i.e. building blocks)

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For example : it can not in general combine instances of

1 1 * * * * * 1

and

* * * * 1 1 * *

to form an instance of

1 1 * * 1 1 * 1.
```

### Positional bias:

The schemas that can be created or destroyed by a crossover depends strongly on the location of the bits in the chromosomes.

### End-point bias:

It is also observed that single-point crossover treats some loci preferentially, that is, the segments exchanged between the two parents always contain the end points of the strings.

### Hamming cliff problem:

A one-bit change can make a large (or a small) jump. A multi-bits can make a small (or a large gap).

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For example, 1000 \implies 0111
(Here, Hamming distance = 4, but distance between phenotype is 1)
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### Similarly, $\textbf{0000} \Longrightarrow \textbf{1000}$

(Here, Hamming distance = 1, but distance between phenotype is 8)

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- To reduce the positional bias and end-point bias, two-point crossover and multi-point crossover schemes have been evolved.
- In contrast, UX and HUX distribute the patterns in parent chromosomes largely resulting too much deflections in the offspring.
- To avoid binary code related problem, gray coding can be used.
- In summary, binary coding is the simplest encoding and its crossover techniques are fastest compared to the crossover techniques in other GA encoding schemes.

### **Crossover Techniques in Real Coded GA**

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Following are the few well known crossover techniques for the real-coded GAs.

- Linear crossover
- Blend crossover
- Binary simulated crossover

• This scheme uses some linear functions of the parent chromosomes to produce the new children.

#### • For example

Suppose  $P_1$  and  $P_2$  are the two parameter's values in two parents, then the corresponding offspring values in chromosomes can be obtained as

$$C_i = \alpha_i P_1 + \beta_i P_2$$

where  $i = 1, 2 \cdots n$  (number of children).  $\alpha_i$  and  $\beta_i$  are some constants.

### Linear crossover: An example

• Example :

Suppose  $P_1 = 15.65$  and  $P_2 = 18.83$ 

$$\alpha_1 = 0.5 = \beta_1$$

$$\alpha_2 = 1.5 \text{ and } \beta_2 = -0.5$$

$$\alpha_3 = -0.5$$
 and  $\beta_3 = 1.5$ 

#### Answer :

$$C_1 = 0.5 \times (P_1 + P_2) = 17.24$$
  
 $C_2 = 1.5 \times P_1 - 0.5 \times P_2 = 14.06$   
 $C_3 = -0.5 \times P_1 + 1.5 \times P_2 = 20.24$ 



#### Advantages

- It is simple to calculate and hence faster in computation
- Can allow to generate a large set of offspring from two parent values
- Ontrols are possible to choose a wide-range of variations

#### Limitations

- **()** Needs to be decided the values of  $\alpha_i$  and  $\beta_i$
- It is difficult for the inexperienced users to decide the right values for α<sub>i</sub> and β<sub>i</sub>
- If  $\alpha_i$  and  $\beta_i$  values are not chosen properly, the solution may stuck into a local optima.

This scheme can be stated as follows.

- Let  $P_1$  and  $P_2$  are the two parameter's values in two parent's chromosomes, such that  $P_1 < P_2$
- Then the blend crossover scheme creates the children solution lying in the range

$$\langle \{P_1 - \alpha (P_2 - P_1)\} \cdots \{P_2 - \alpha (P_2 - P_1)\} \rangle$$

where  $\alpha$  is a constant to be decided so that children solution do not come out of the range of domain of the said parameter.

Solution Another parameter  $\gamma$  has to be identified by utilizing the  $\alpha$  and a random number *r* in the range of (0.0, 1.0) both exclusive like the following:

$$\gamma = (\mathbf{1} + \mathbf{2}\alpha) \, \mathbf{r} - \alpha$$

• The children solutions  $C_1$  and  $C_2$  are determined from the parents as follows,

$$C_1 = (1 - \gamma) P_1 + \gamma P_2$$
  
$$C_2 = (1 - \gamma) P_2 + \gamma P_1$$

### **Blend crossover : An example**

- Example :
  - $P_1 = 15.65$  and  $P_2 = 18.83$
  - lpha= 0.5 and  $\gamma=$  0.6



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- This scheme is based on the probability distribution of generated children solution from the given parents.
- A spread factor α is used to represent the spread of the children solutions with respect to that of the parents, as given below.

$$\alpha = \left| \frac{C_1 - C_2}{P_1 - P_2} \right|$$

Here  $P_1$  and  $P_2$  are represent the parent points and  $C_1$  and  $C_2$  are two children solutions.

Three different cases may occurs:

• Case 1:  $\alpha$  < 1 (Contracting Crossover)

The spread of children is less than the parents.

• Case 2:  $\alpha > 1$  (Expanding Crossover)

The spread of children is more than the parents.

• Case 3:  $\alpha = 1$  (Stationary Crossover)

The spread of children is same as that of the parents.

#### **Probability Distribution:**

• Case 1: For Contracting Crossover

$$C(\alpha) = 0.5(q+1)\alpha^2$$

• Case 2: For Expanding Crossover

$$E(\alpha) = 0.5(q+1)\frac{1}{\alpha^{q+2}}$$

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### Simulated binary crossover in Real-coded GAs

Following steps are used to create two children solutions  $C_1$  and  $C_2$  from the parents  $P_1$  and  $P_2$ .

• Create a random number  $r \in \{0.0 \cdots 1.0\}$ 

.

2 Determine  $\alpha'$  such that

$$\int_0^{lpha'} {m {\cal C}}(lpha) {m d} lpha = {m r}, \, {
m if} \; {m r} < {
m 0.5}$$

and

$$\int_{1}^{\alpha'} E(\alpha) d\alpha = r$$
, if  $r > 0.5$ 

**③** Using the value of  $\alpha'$  obtain two children solution as follows

• 
$$C_1 = 0.5 [(P_1 + P_2) - \alpha' |P_2 - P_1|]$$

•  $C_2 = 0.5[(P_1 + P_2) + \alpha' |P_2 - P_1|]$ 

### Simulated binary crossover in Real-coded GAs

#### Example:

- $P_1 = 15.65$
- $P_2 = 18.83$

 $\alpha'={\bf 1.0772}$ 

#### Assuming expanding crossover with r > 0.5



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### Advantages and limitations

#### Advantages

- We can generate a large number of offspring from two parents.
- 2 More explorations with diverse offspring.
- Results are accurate and usually terminated with global optima.
- Termination with a less number of iterations.
- Crossover techniques are independent of the length of the chromosome.

#### Limitations

- Computationally expensive compared to binary crossover.
- If proper values of parameters involved in the crossover techniques are not chosen judiciously, then it may lead to premature convergence with not necessarily optimum solutions.

### **Crossover Techniques in Order GAs**

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### Crossover techniques in order GA

- Any binary crossover techniques are not applicable to Order coded GAs
  - Example



Here, the offspring are not valid chromosomes

 Since, sequence of gene values are important, Real-coded crossover techniques, which are to produce real number from two given real numbers are also not applicable to Order-coded GAs.

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### Crossover techniques in order GA

Some important crossover techniques in Order-coded GAs are:

- Single-point order crossover
- 2 Two-point order crossover
- Partially mapped crossover (PMX)
- Position based crossover
- Precedence-preservation crossover (PPX)
- Edge recombination crossover

Assumptions: For all crossover techniques, we assume the following:

- Let *L* be the length of the chromosome.
- *P*<sub>1</sub> and *P*<sub>2</sub> are two parents (are selected from the mating pool).
- $C_1$  and  $C_2$  denote offspring (initially empty).

Given two parents  $P_1$  and  $P_2$  with chromosome length, say *L*. **Steps:** 

- Sandomly generate a crossover point K such that (1 < K < L).
- Copy the left schema of  $P_1$  into  $C_1$  (initially empty) and left schema of  $P_2$  into  $C_2$  (also initially empty).
- So For the schema in the right side of  $C_1$ , copy the gene value from  $P_2$  in the same order as they appear but not already present in the left schema.
- Subscription  $\Theta$  Repeat the same procedure to complete  $C_2$  from  $P_1$ .

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#### Example :



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It is similar to the single-point order crossover, but with two k-points.

#### Steps:

- Randomly generate two crossover points  $K_1$  and  $K_2$ . 1 <  $K_1$ ,  $K_2 < L$
- 2 The schema in middle of  $P_1$  and  $P_2$  are copied into  $C_1$  and  $C_2$  (initially both are empty), respectively in the same location as well as in the same order.
- So The remaining schema in  $C_1$  and  $C_2$  are copied from  $P_2$  and  $P_1$  respectively, so that an element already selected in child solution does not appear again.

Example :



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Let the parent chromosomes be  $P_1$  and  $P_2$  and the length of chromosomes be *L*.

#### Steps:

- (a) Create a vector *V* of length *L* randomly filled with elements from the set {1,2}.
- (b) This vector defines the order in which genes are successfully drawn from  $P_1$  and  $P_2$  as follows.
  - We scan the vector *V* from left to right.
  - 2 Let the current position in the vector V be i (where  $i = 1, 2, \dots, L$ ).
  - 3 Let *j* (where  $j = 1, 2, \dots, L$ ) and *k* (where  $j = 1, 2, \dots, L$ ) denotes the *j*<sup>th</sup> and *k*<sup>th</sup> gene of *P*<sub>1</sub> and *P*<sub>2</sub>, respectively. Initially j = k = 1.

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#### If i<sup>th</sup> value is 1 then

Delete  $j^{th}$  gene value from  $P_1$  and as well as from  $P_2$  and append it to the offspring (which is initially empty).

Else

Delete  $k^{th}$  gene value from  $P_2$  and as well as from  $P_1$  and append it to the offspring.

Repeat Step 2 until both P<sub>1</sub> and P<sub>2</sub> are empty and the offspring contains all gene values.

### Precedence preservation order crossover : Example

#### Example :



**Note :** We can create another offspring following the alternative rule for 1 and 2.

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#### Steps :

- Choose *n* crossover points  $K_1, K_2 \cdots K_n$  such that  $n \ll L$ , the length of chromosome.
- The gene values at  $K_1^{th}, K_2^{th} \cdots K_n^{th}$  positions in  $P_1$  are directly copied into offspring  $C_1$  (Keeping their position information intact).
- Solution The remaining gene values in  $C_1$  will be obtained from  $P_2$  in the same order as they appear there except they are already not copied from  $P_1$ .
- We can reverse the role of  $P_1$  and  $P_2$  to get another offspring  $C_2$ .

### Position-based order crossover : Example

Let su consider three *k*-points namely  $K_1$ ,  $K_2$  and  $k_3$  in this example.



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- This crossover technique is used to solve TSP problem when the cities are not completely connected to each other.
- In this technique, an edge table which contains the adjacency information (but not the order).
- In the other words, edge table provides connectivity information.

### Edge recombination order crossover: Illustration

#### Example

- Let us consider a problem instance of a TSP with 9 cities.
- Assume any two chromosome  $P_1$  and  $P_2$  for the mating.

$$\begin{array}{c} \mathsf{P1:} & 1 \longrightarrow 2 \longrightarrow 4 \longrightarrow 6 \longrightarrow 8 \longrightarrow 7 \longrightarrow 5 \longrightarrow 3 \\ \mathsf{P2:} & 4 \longrightarrow 3 \longrightarrow 5 \longrightarrow 7 \longrightarrow 8 \longrightarrow 6 \longrightarrow 2 \longrightarrow 1 \end{array}$$

#### Connectivity graph:



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### Edge recombination order crossover: Illustration

#### Edge table for the connectivity graph:



City	Connectivity							
1		2	4	3				
2		1	4	7	6			
3		1	4	5				
4		1	2	3	6			
5		3	7	8				
6		2	4	8				
7		2	5	8				
8		5	6	7				

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### Edge recombination order crossover: Illustration

#### Steps:

Let the child chromosome be  $C_1$  (initially empty).

- **(**) Start the child tour with the starting city of  $P_1$ . Let this city be *X*.
- Append city X to C.
- Oblight Delete all occurrences of X from the connectivity list of all cities (right-hand column).
- From city X choose the next city say Y, which is in the list of minimum (or any one, if there is no choice) connectivity links.
- So Make X = Y [i.e. new city Y becomes city X].
- Repeat Steps 2-5 until the tour is complete.



## **Any questions??**

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