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## Representations for Evolutionary Algorithms

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### Structure of the Tutorial

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- A Short Introduction to Representations
  - Defining Representations
  - Representations, Operators, and Metrics
  - Direct and Indirect Representations
- Design Guidelines for Representations
- Properties of Representations
  - Redundant Representations and Neutral Networks
  - High-Locality Representations
  - Domino Convergence and Genetic Drift

### Scope of the Tutorial

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- Illustrate the influence of representations on the performance of EAS.
- Illustrate the relationship between problem difficulty and used representation/operator.
- Review design guidelines for high-quality representations.
- Focus on some properties of representations
  - Redundant representations and neutral search spaces
  - High-locality representations
  - (Exponentially scaled alleles)

### Defining Representations

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- A representation assigns genotypes to corresponding phenotypes.
- Every search and optimization algorithm needs a representation.
- The representation allows to represent a solution to a specific problem.
- Different representations can be used for the same problem.
- Performance of search algorithm depends on properties of the used representation and how suitable is the representation in the context of the used genetic operators.

- There are many different representations.
- Standard representations are binary, real-valued vectors, messy encodings, tree structures,...
- ... and we assume that everybody has some experience at least with some of them.

Every optimization problem  $f(\mathbf{x})$  can be separated into a genotype-phenotype mapping  $f_g$  and a phenotype-fitness mapping  $f_p$ :

$$f_g(\mathbf{x}_g) : \Phi_g \rightarrow \Phi_p,$$

$$f_p(\mathbf{x}_p) : \Phi_p \rightarrow \mathbb{R},$$

where  $f = f_p \circ f_g = f_p(f_g(\mathbf{x}_g))$ .

A change of  $f_g$  also changes the properties of  $f$ .

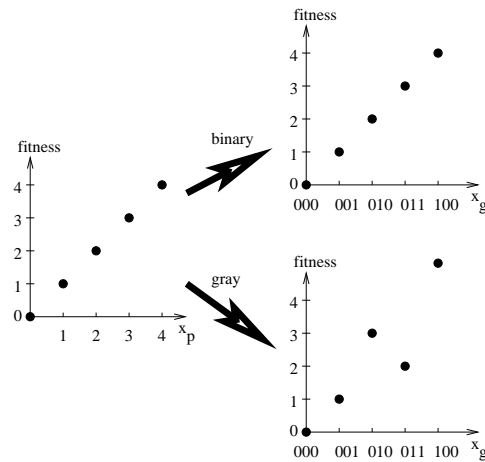
The genetic operators mutation and crossover are applied to  $\mathbf{x}_g$ , whereas the selection process is based on the fitness of  $\mathbf{x}_p$ .

$f_p(\mathbf{x}_p)$  determines the difficulty and complexity of a problem.

$f_g(\mathbf{x}_g)$  is the used representation.

There are  $|\Phi_g|!$  different representations.

- Representations change the character and difficulty of optimization problems.
- For example  $f_p = x_p$ , where  $x_p \in \mathbb{N}$ .
- Different problem depending on the used representations (Gray versus binary).



- Phenotypic problem easy to solve for hill-climber.
- When using bit-flipping GA the Gray-encoded problem is easier to solve than the binary-encoded problem.
- Gray encoding induces less local optima when used on problems of practical relevance (compare Free Lunch theorem (Whitley, 2000)).
- Resulting problem difficulty depends on used search method. If other search methods (e.g. other operators) are used, then problem difficulty is different (compare (Reeves, 2000)).

Representation, metric defined on  $\Phi_g$  and  $\Phi_p$ , and genetic operators depend on each other and are closely related.

- A representation is just a mapping from  $\Phi_g$  to  $\Phi_p$ . It assigns any  $x_g \in \Phi_g$  to an  $x_p \in \Phi_p$ .
- In both search spaces,  $\Phi_g$  and  $\Phi_p$ , a metric is or has to be defined. The metric determines the distances between the individuals and is the basis for measuring similarities between individuals. In general, the metric used for  $\Phi_p$  is defined by the considered problem. The metric used for  $\Phi_g$  is determined by the used search operators.
- Genotypic operators like mutation and crossover are defined based on the used metric.

### Results:

- Metric on  $\Phi_g$  and used operators depend on each other. The one determines the other.
- Representations transform the metric on  $\Phi_g$  to the (problem-dependent) metric on  $\Phi_p$ . (Compare locality, causality, and distance distortion)

### Mutation:

The application of mutation to an individual results in a new individual with similar properties. There is a small distance between offspring and parent.

### Crossover:

Crossover combines the properties of two or more parents in an offspring. The distance between offspring and parent should be smaller than the distance between both parents.

(compare also (Surry and Radcliffe, 1996a) and (Liepins and Vose, 1990))

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Direct Representations

If the genetic operators are applied directly to the phenotypes it is not necessary to specify a representation and the phenotypes are identical with the genotypes:

$$\begin{aligned}f_g(\mathbf{x}_g) &: \Phi_g \rightarrow \Phi_g, \\f_p(\mathbf{x}_p) &: \Phi_g \rightarrow \mathbb{R}.\end{aligned}$$

This means,  $f_g$  is the identity function  $f_g(\mathbf{x}_g) = \mathbf{x}_g$ .

Using direct representations do not necessarily make life easier:

- Design of proper operators is difficult
- How can we apply EDAs?
- Representation issues are not important any more ( $\Phi_g = \Phi_p$  and  $f_g(\mathbf{x}_g) = \mathbf{x}_g$ ).

Representation issues are also relevant to Genetic Programming.

Phenotypes: Programs, logical expressions.

Genotypes: Bitstrings, trees, ...

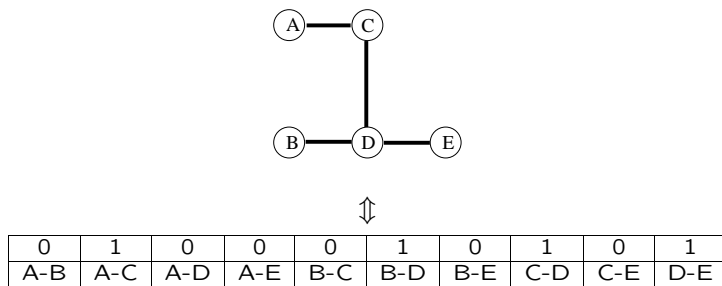
Neglecting proper genotype-phenotype mappings can result in low performance of GP approaches.

## Indirect Representations - Specific Constraints

Example: Tree optimization problems

A tree is a fully connected graph with exactly  $n - 1$  links (for an  $n$  node network). There are no circles in a tree.

A graph can be represented by its characteristic vector.

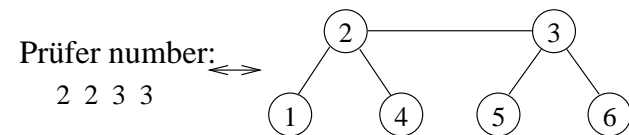


The use of an explicit genotype-phenotype mapping has some benefits:

- Specific constraints can be considered.
- Standardized genetic operators with known behavior and properties can be used.
- An indirect representation is necessary if problem-specific operators are either not available or difficult to design.
- Representation can make problem easier by incorporating problem-specific knowledge.

## Indirect Representations - Specific Constraints (2)

Prüfer numbers are a one-to-one mapping between trees and a sequence of integers. A tree with  $n$  nodes is represented by a string of length  $n - 2$  over an alphabet of  $n$  symbols.



Therefore, using Prüfer numbers allows to consider the constraint that the graph is a tree (For other representations repair operators are necessary).

- When mapping many different types of phenotypes on only a few types of different genotypes (binary, integer, or continuous representations), it is possible to use standardized operators.
- Behavior of EAs for standard representations like binary (simple GAs) or continuous (evolution strategies) representations well understood.
- Mapping phenotypes on binary genotypes allows the use of schemata and effective linkage learning GAs (under the assumption that the problem still remains decomposable and that binary encodings allow a natural encoding of the problem).

- For many real-world problems there are no problem-specific operators available.
- Developing of problem-specific operators is difficult and often additional repair mechanisms must be used to ensure a valid solution.

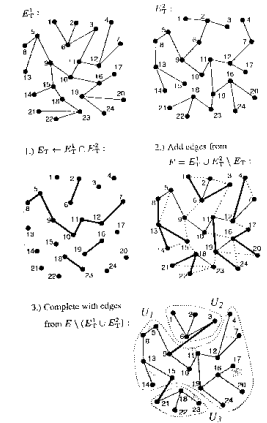
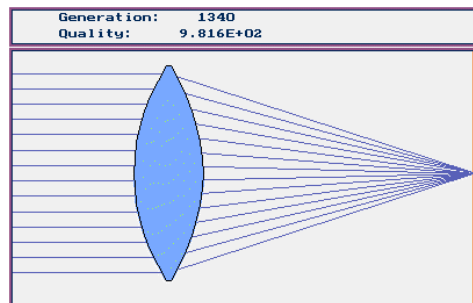


Figure 2: An example for edge crossover ( $d = 3$ ).

(from (Raidl, 2000))

For some types of problems no problem-specific operators exist that can be applied to direct representations.



Incorporating problem-specific knowledge in the representations to increase GA performance:

- Increase the initial supply of solutions that are similar to the optimal solution.
- Use high-locality representations for easy problems.
- Consider specific properties of the optimal solution (e.g. stars and trees).
- Use representations that make a problem easier for a specific optimization method.

- A Short Introduction to Representations
- **Design Guidelines for Representations**
- Properties of Representations
  - Redundant Representations and Neutral Networks
  - High-Locality Representations
  - Domino Convergence and Genetic Drift

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Goldberg's Recommendations (2)

- The recommendations caused a lot of critics (Radcliffe, 1997; Fogel and Stayton, 1994).
- What is a natural representation of a problem? (For example, is using binary representations for encoding real-valued phenotypes a natural representation?)
- Principles mainly aimed at binary representations and crossover-based GAs that process schemata. Not big help for other search methods like evolution strategies or evolutionary programming as these search methods do not process schema.

- Principle of meaningful building blocks: The schemata should be short, of low order, and relatively unrelated to schemata over other fixed positions.
- Principle of minimal alphabets: The alphabet of the encoding should be as small as possible while still allowing a natural representation of solutions (qualified by (Goldberg, 1991))

from (Goldberg, 1989))

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Radcliffe's Recommendations

Representation and operators belong together and can not be separated from each other (Radcliffe, 1992).

Design of representation-independent evolutionary algorithms is possible if the following properties are considered (Surry and Radcliffe, 1996b):

- **Respect:** Offspring produced by recombination are members of all formae to which both their parents belong.
- **Transmission:** Every gene is set to an allele which is taken from one of the parents.
- **Assortment:** Offspring can be formed with any compatible characteristics taken from the parents.
- **Ergodicity:** Iterative use of operators allow to reach any point in the search space.

## Palmer's Recommendations

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- An encoding should be able to represent all possible phenotypes.
- An encoding should be unbiased in the sense that all possible individuals are equally represented in the set of all possible genotypic individuals.
- An encoding should encode no infeasible solutions.
- The decoding of the phenotype from the genotype should be easy.
- An encoding should possess locality. Small changes in the genotype should result in small changes in the phenotype (compare statements about metric).

from (Palmer, 1994))

## Ronald's Recommendations

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- Encodings should be adjusted to a set of genetic operators in a way that the building blocks are preserved from the parents to the offspring (Fox and McMahon, 1991).
- Encodings should minimize nonlinearities in fitness functions (Beasley et al., 1993). This means, representations should make the problem easier.
- Feasible solutions should be preferred.

## Ronald's Recommendations (2)

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- The problem should be represented at the correct level of abstraction.
- Encodings should exploit an appropriate genotype-phenotype mapping process if a simple mapping to the phenotype is not possible.
- Isomorphic forms, where the phenotype of an individual is encoded with more than one genotype, should not be used.

from (Ronald, 1997))

## Design Guidelines - Summary

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- Based on observations for specific test problems there are some common, fuzzy ideas about what is a good representation.
- Recommendations too general to be helpful for designing or evaluating representations.
- There is a lack of analytical models describing the influence of representations on EAs.
- To verify (or reject) observations analytical models are necessary.

Representations are redundant if the number of genotypes is larger than the number of phenotypes.

- Using redundant representations  $f_g$  means changing  $f = f_p(f_g)$ . There are additional plateaus in the fitness landscape.
- Redundant representations are more inefficient encodings which use a higher number of alleles but do not increase the amount of encoded information.
- Redundant representations are not an invention of AI researchers but are commonly used in nature.

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## Redundant Representations (2)

There are different opinions regarding the influence of redundant representation on the performance of EAs.

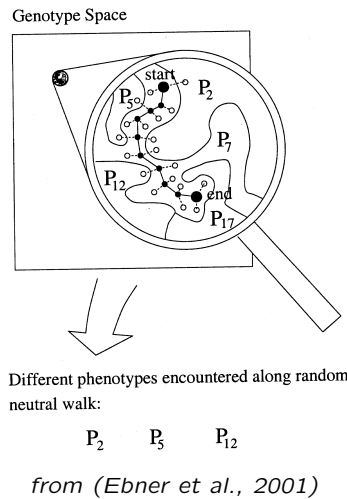
- Redundant representations reduce EA performance due to loss of diversity (Davis, 1989; Eshelman and Schaffer, 1991; Ronald et al., 1995)
- Redundant representations increase EA performance (Gerrits and Hogeweg, 1991; Cohoon et al., 1988; Julstrom, 1999)

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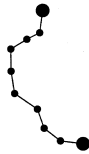
## Redundant Representations (3)

- Large amount of work considers the *neutral theory* (Kimura, 1983). This theory assumes that not natural selection fixing advantageous mutations but the random fixation of neutral mutations is the driving force of molecular evolution.
- Following these ideas redundant representations (neutral networks) are used in EAs with great enthusiasm.
- There is hope that increasing the evolvability of a system also increases the performance of the system (Barnett, 1997; Barnett, 1998; Shipman, 1999; Shipman et al., 2000b; Shackleton et al., 2000; Shipman et al., 2000a; Ebner et al., 2001; Smith et al., 2001c; Smith et al., 2001a; Smith et al., 2001b; Barnett, 2001; Yu and Miller, 2001; Yu and Miller, 2002; Toussaint and Igel, 2002).
- (Knowles and Watson, 2002) showed exemplarily that this is not true!





Neutral Network: Set of genotypes connected by single-point mutations that map to the same phenotype.



In the following slides we present a model that

- explains how redundancy changes the performance of EAs and
- allows quantitative predictions of EA performance (Rothlauf and Goldberg, 2003).

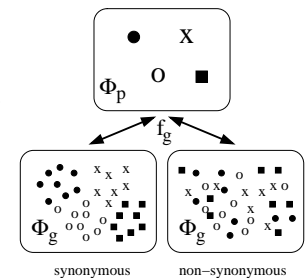
### Benefits of Neutral Networks

- Population can drift along these neutral networks.
- Reducing the chance of being trapped in sub-optimal solutions.
- Population is quickly able to recover after a change has occurred.
- Evolvability of the system increases.

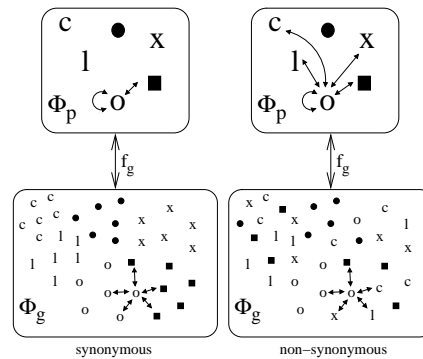
### Synonymously versus Non-synonymously Redundant Representations

When using redundant representations it can be distinguished between:

- Synonymously redundant representations:** All genotypes that encode the same phenotype are similar to each other.
- Non-synonymously redundant representations:** Genotypes that encode the same phenotype are not similar to each other.



- Non-synonymously redundant representations do not allow guided search.
- EA search becomes random.
- Similar effect as low locality representations.



*Effects of small mutation steps*

Synonymously redundant representations can be described using

- order of redundancy  $k_r = \frac{\log(|\Phi_p|)}{\log(|\Phi_g|)}$  and
- over-, resp. underrepresentation  $r$  of the optimal solution due to the problem representation  $f_g$ .

When using the notion of BBs and binary representations:

- $k_r = \frac{k_g}{k_p}$
- $r$ : Number of genotypic BBs of order  $k_g$  that represent the optimal phenotypic BB of order  $k_p$ .

### Example 1:

genotypes $x_g$	$x_p$
00 00, 00 01, 01 00, 01 01	0 0
10 00, 10 01, 11 00, 11 01	1 0
00 10, 01 11, 00 11, 01 11	0 1
10 10, 10 11, 11 10, 11 11	1 1

- $k = 2$  (order of phenotypic BBs)
- $k_r = 2$  (One allele of a phenotype is represented using two alleles of a genotype)
- Uniform redundancy:  $r = 4$  (the best BB (e.g..  $x_p = 11$ ) is represented by four genotypic BBs)

### Example 2:

genotypes $x_g$	$x_p$
000, 001, 010, 100, 101, 110, 011	0
111	1

- $k = 1$  (order of phenotypic BBs)
- $k_r = 3$  (One phenotypic allele is represented using three genotypic alleles)
- Non-uniform redundancy:  $r = 1$  (best BB ( $x_p = 1$ ) is represented by one genotypic BB ( $x_g = 111$ ))

## Population Sizing for GAs

The Gambler's ruin model (Feller, 1957) can be used for modeling the iterated decision making in GAs.

A gambler with initial stake  $x_0$  wishes to increase his funds to a total of  $N$  units by making a sequence of bets against a gaming house. Each bet has fixed probability  $p$  of winning ( $q = 1 - p$  of losing), and we wish to know the probability of succeeding (getting  $N$  units) or failing (losing all units).

Following (Harik et al., 1997) the probability that a GA with a population size  $N$  converges after  $t_{conv}$  generations to the correct solution is

$$P_n = \frac{1 - (q/p)^{x_0}}{1 - (q/p)^N}$$

## Population Sizing for GAs (2)

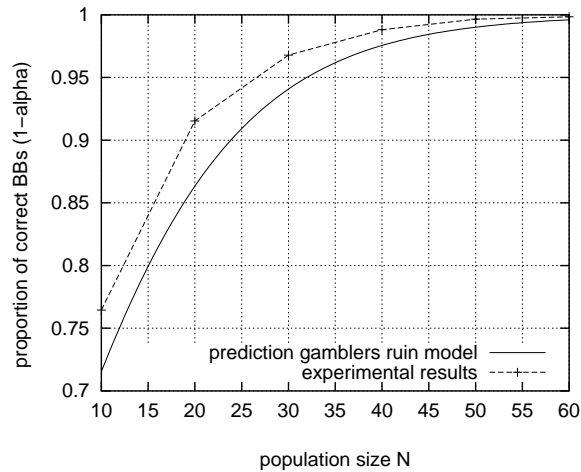
After some calculations we get:

$$N \approx -2^{k-1} \ln(\alpha) \frac{\sigma_{BB} \sqrt{\pi m'}}{d}$$

$N$  is the necessary population size,  $\alpha = 1 - P_n$  the probability  $P_n$  that the optimal BB cannot be found (probability of failure) and  $k$  is the order of the BBs.

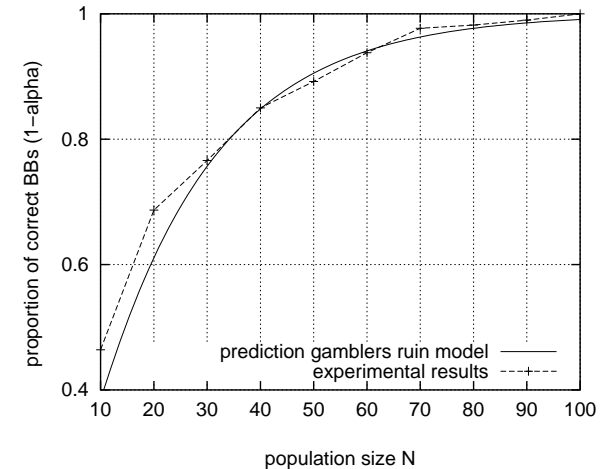
$\sigma_{BB}$  (variance of BBs),  $d$  (fitness difference between best and second best BB),  $m' = m - 1$  (number of BBs) and  $k$  are problem-dependent.

## Population Sizing for GAs (3)



150-bit one-max problem ( $k = 1$ ,  $\sigma_{BB} = 0.25$ ,  $d = 1$  and  $m = 150$ )

## Population Sizing for GAs (4)



Ten concatenated 3-bit deceptive traps ( $k = 3$ ,  $\sigma_{BB} = 1$ ,  $d = 1$  and  $m = 10$ )

Now we have to ask how the redundancy of a representation influences GA performance?

Observation: Redundant representation change the initial supply  $x_0$  of BBs.

For binary problem representation:

$$x_0 = N \frac{r}{2^{k_r}},$$

where  $N$  is the population size.

Conclusions from this model:

- Redundant representations can change the performance of EAs.
- If representations are synonymously redundant:
  - Uniformly redundant representations do not change the performance of EAs!
  - If the optimal BB is overrepresented GA performance increases.
  - If the optimal BB is underrepresented GA performance decreases.
- Redundant representations can not be used systematically if there is no problem-specific knowledge!

When using synonymously redundant representations the existing model can be extended:

$$N \approx -\frac{2^{k_r k-1}}{r} \ln(\alpha) \frac{\sigma_{BB} \sqrt{\pi m'}}{d}$$

The population size  $N$  that is necessary to find the optimal solution with probability  $P_n = 1 - \alpha$  goes with  $O\left(\frac{2^{k_r}}{r}\right)$ .

What must be considered when using redundant representations?

1. How does the used representation change the size of the search space?
2. Is the representation synonymously redundant?
3. Are some solutions overrepresented?

Examining these properties allows the user to increase the performance of EAs!

In the following slides we show how this theory can be used for predicting EA performance when using the trivial voting mapping for binary problems.

## Trivial Voting Mapping (2)

### Examples:

genotypes $x_g$	$x_p$
000, 001, 010, 100	0
110, 101, 011, 111	1

- $k = 1$
- $k_r = 3$
- $u = 2$

genotypes $x_g$	$x_p$
000	0
001, 010, 100, 110, 101, 011, 111	1

- $k = 1$
- $k_r = 3$
- $u = 1$

## Trivial Voting Mapping

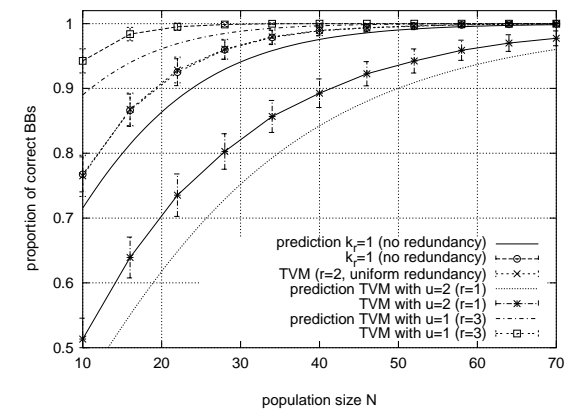
- The trivial voting mapping (TVM) assigns binary phenotypes to binary genotypes.
- One bit of the phenotype is represented by  $k_r$  genotypic bits.
- In general, a phenotypic bit is 0 if less than  $u$  genotypic bits are zero. If more than  $u$  genotypic bits are 1 then the phenotypic bit is 1.
- For  $u = k_r/2$  the value of the phenotypic bit is determined by the majority of the genotypic bits (majority vote)

In general:

$$x_i^p = \begin{cases} 0 & \text{if } \sum_{j=0}^{k_r-1} x_{k_r i+j}^g < u \\ 1 & \text{if } \sum_{j=0}^{k_r-1} x_{k_r i+j}^g \geq u, \end{cases}$$

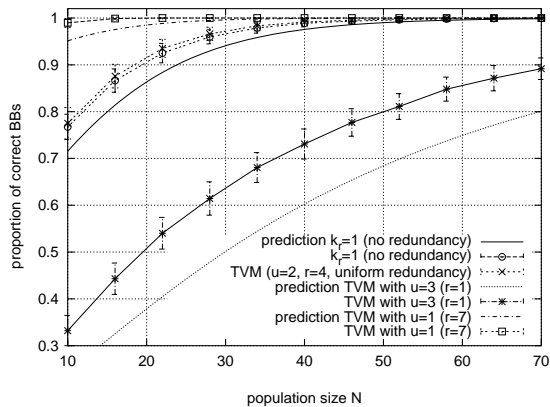
where  $u \in \{1, \dots, k_r\}$ .

## Trivial Voting Mapping (3)



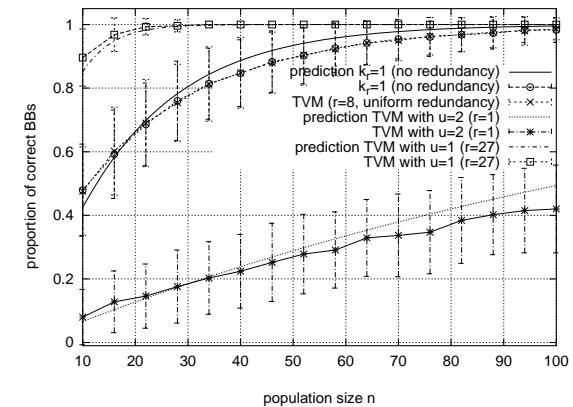
Experimental and theoretical results of the proportion of correct BBs on a 150-bit one-max problem using the trivial voting mapping for  $k_r = 2$ .

## Trivial Voting Mapping (4)



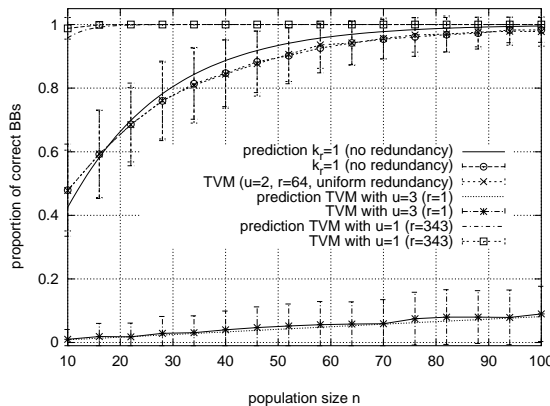
Experimental and theoretical results of the proportion of correct BBs on a 150-bit one-max problem using the trivial voting mapping for  $k_r = 3$ .

## Trivial Voting Mapping (5)



Experimental and theoretical results of the proportion of correct BBs for ten concatenated 3-bit deceptive traps and  $k_r = 2$ .

## Trivial Voting Mapping (6)



Experimental and theoretical results of the proportion of correct BBs for ten concatenated 3-bit deceptive traps and  $k_r = 3$ .

## Redundant Representations - Summary

- There are theoretical models that allow us to predict the expected GA performance when using redundant representations ( $N = O(2^{k_r/r})$ ).
- There are guidelines for the design of redundant representations:
  - Do not use non-synonymously redundant representations!
  - If you use redundant representations you have to investigate:
    - \* How does the representation change the size of the search space?
    - \* Are solutions similar to the optimal solution overrepresented?
  - If there is no knowledge about the optimal solution use a uniformly redundant representation.

- Representations (genotype-phenotype mappings) can change the structure of the neighborhood and the structure of the fitness landscapes.
- Each neighbor can be reached directly by a move (mutation, crossover, etc.). Therefore, the neighborhood structure depends on the used operator and the used metric.
- The set of neighbors can be different for the genotypes and phenotypes.
- The distance between two individuals is determined by the number of moves between both individuals.

## Locality (2)

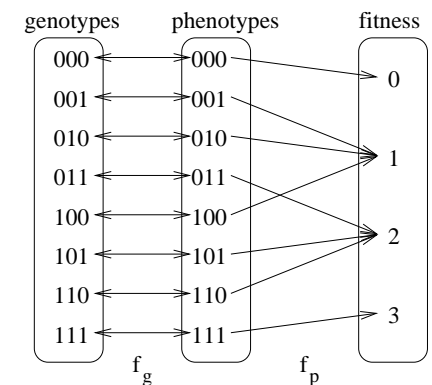
**The locality of a representation describes how well neighboring phenotypes correspond to neighboring genotypes.** Therefore, the locality of a representation is high, if neighboring genotypes correspond to neighboring phenotypes.

Locality, causality, and distance distortion describe how well the metric on  $\Phi_p$  fits to the metric on  $\Phi_g$ . If they fit well the locality is high.

Representations  $f_g$  that change the distances between corresponding genotypes and phenotypes modify the difficulty of the problem ( $\text{difficulty}(f) \neq \text{difficulty}(f_p)$ ).

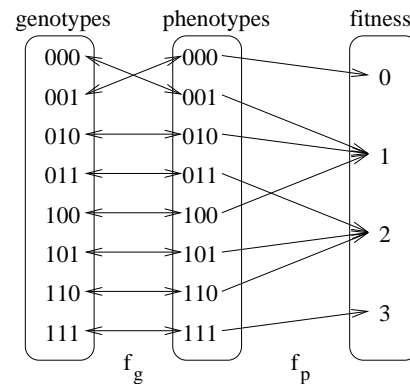
## Locality - An Example

- Both, genotypes and phenotypes are binary.
- We use the bit-flipping operator as a move (Hamming distance).
- One-max problem.
- All building blocks (regarding genotypes and phenotypes) are of size  $k = 1$ . Therefore, problem is easy for selectore-combinative GAs.

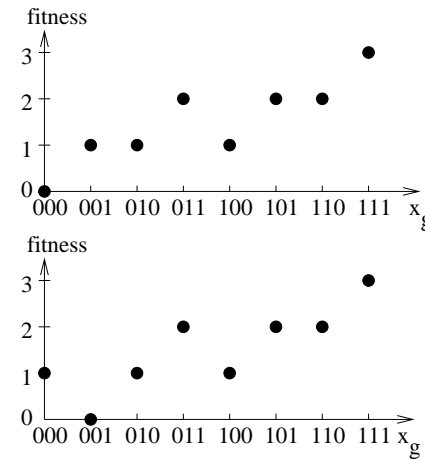


## Locality - An Example (2)

- A representation with lower locality.
- The neighborhood structure changes.
- Not all genotypic building blocks are of size 1. Although,  $f_p$  remains unchanged,  $f$  becomes more difficult.



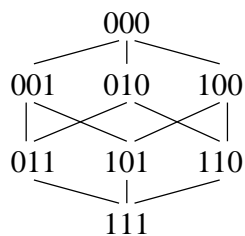
## Locality - An Example (3)



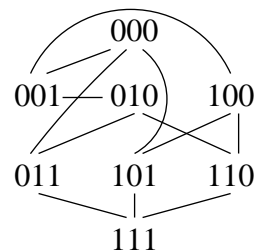
- High-locality representation.
- Problem easy for selectorecombinative GAs.
- Different fitness for genotypes 000 and 001.
- Problem more difficult for selectorecombinative GAs.
- Neighborhood not preserved by representation.

## Locality - An Example (4)

Neighborhood structure of the genotypes:



Resulting neighborhood structure of the phenotypes:



## Comparing Representations

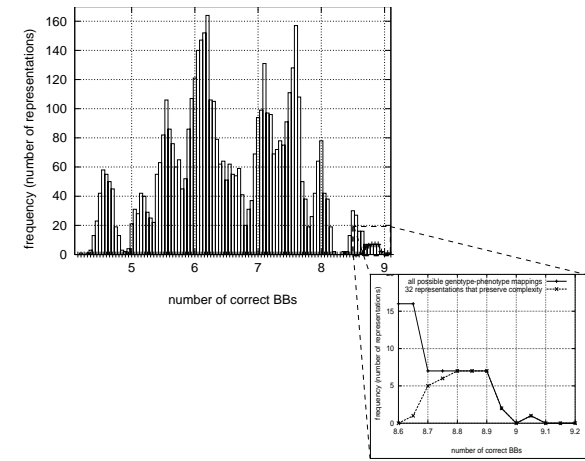
- We compare the performance of selectorecombinative GAs over all different representations for the one-max problem.
- When focusing on binary bitstrings and assigning  $l$ -bit genotypes to  $l$ -bit phenotypes, there are  $2^l$  different representations.
- For  $l = 3$  there are 8 different genotypes, resp. phenotypes, and  $8! = 40320$  different representations.
- 36 different representations result in the same overall problem  $f$  (for the one-max problem).



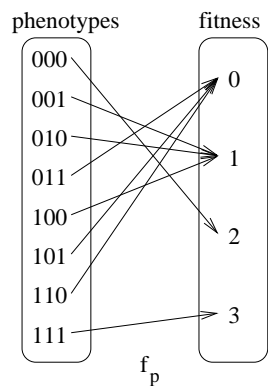
## Comparing Representations (2)

- To reduce problem complexity,  $x_g = 111$  is always assigned to  $x_p = 111$ . Therefore, there are  $7! = 5040$  different representations.
- We concatenate ten 3-bit problems and use a GA with tournament selection of size 2, uniform crossover, and  $N = 16$ .

## Comparing Representations (3)

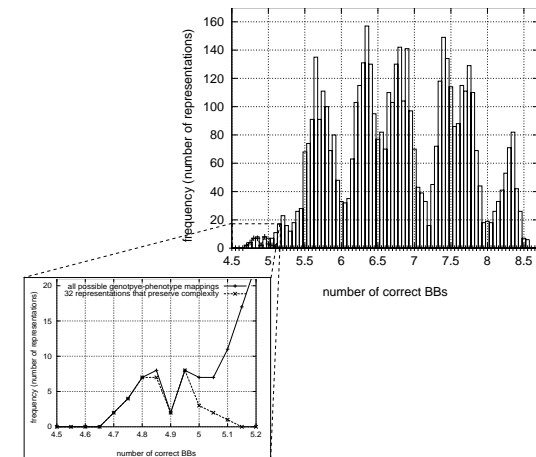


## Comparing Representations (4)



- We compare the performance of selectorecombinative GAs over all different representations for the deceptive trap problem.
- To reduce problem complexity,  $x_g = 111$  is always assigned to  $x_p = 111$ . Therefore, there are  $7! = 5040$  different representations.
- We concatenate ten 3-bit problems and use a GA with tournament selection of size, uniform crossover, and  $N = 16$ .

## Comparing Representations (5)



- When using high locality representations, genotypic neighbors correspond to phenotypic neighbors.
- High locality representations do not change the structure and difficulty of the problem.
  - Easy problems remain easy.
  - Difficult problems remain difficult.
- Locality depends on the used distance metrics which depend on the used operators.

## Exponentially Scaled Alleles

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The alleles of a genotype can be of different importance for the construction of the phenotype.

In many real-world problems it is unclear if the genotypic alleles are uniformly or non-uniformly scaled.

A GA solves the most important alleles first and continues with lower salient alleles (domino convergence)

Genotypic alleles that have little influence on the phenotype are randomly fixed due to genetic drift.

- 
- A Short Introduction to Representations
  - Design Guidelines for Representations
  - Properties of Representations
    - Redundant Representations and Neutral Networks
    - High-Locality Representations
    - **Domino Convergence and Genetic Drift**

## Domino Convergence

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The contribution of the genotypic alleles to the construction of the phenotype can be either uniformly or non-uniformly scaled.

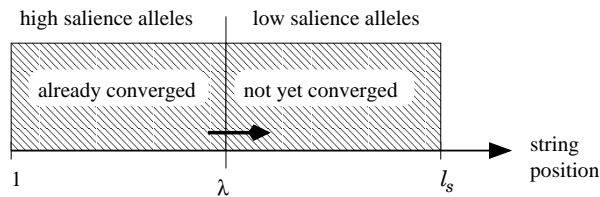
- Uniformly scaled representations:
  - Unary encoding, Gray encoding
  - All alleles are solved implicitly in parallel.
- Exponentially scaled representations:
  - Binary encoding
  - The alleles are solved step by step and domino convergence occurs.

## Domino Convergence (2)

The BinInt problem:  $f(x) = \sum_{i=0}^{l-1} x_i 2^{l-i-1}$  can be decomposed in

- the exponentially scaled representation  $f_g(x_g) = \sum_{i=0}^{l-1} 2^i x_{g,i}$ ,
- and the problem  $f_p(x_p) = x_p$ .

When using GAs and non-uniformly scaled representations domino convergence occurs.



## Domino Convergence (3)

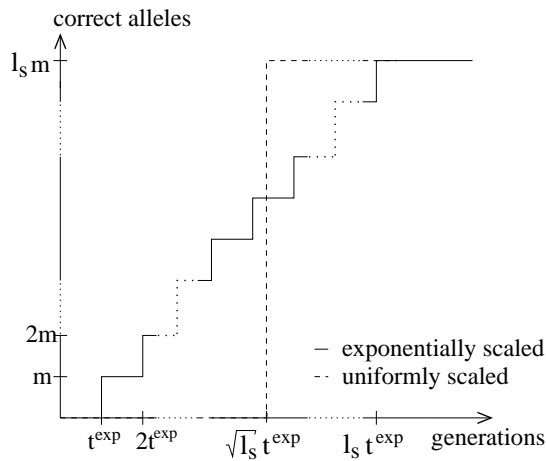
Domino convergence changes the dynamics of selectorecombinative GAs.

Time complexity (neglecting genetic drift):

Uniformly scaled alleles		Exponentially scaled alleles	
const sel. int.	prop. sel.	const sel. int.	prop. sel.
$O(\sqrt{l})$	$O(l \ln(l))$	$O(l)$	$O(2^l)$

Exponentially scaled representations result in longer GA runs!

## Domino Convergence (4)

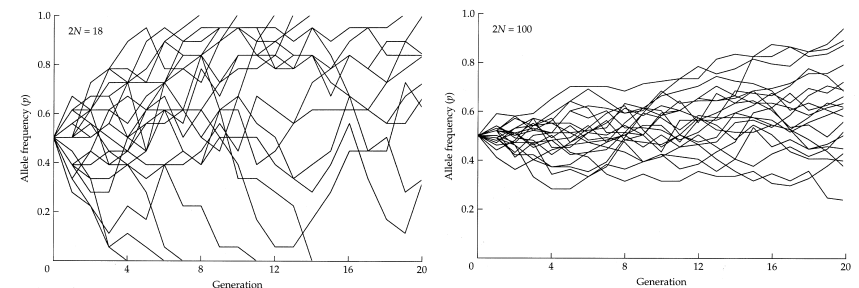


Comparison of time complexity using constant selection intensity:

$t^{exp}$ : time for solving one exponentially scaled allele  
 $m$ : number of exponentially scaled building blocks  
 $l_s$ : length of one exponentially scaled building block

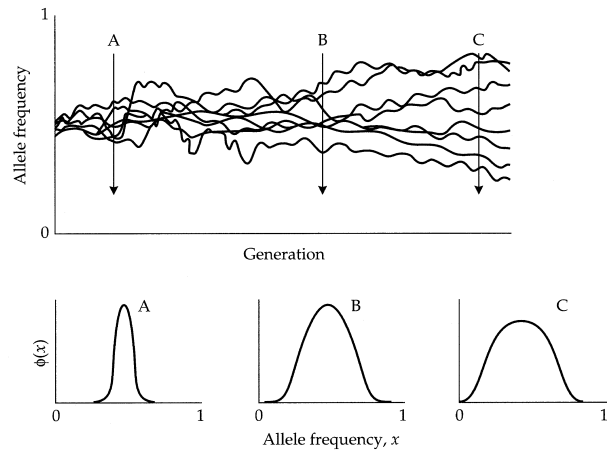
## Genetic Drift

If there is no selection pressure, genetic drift occurs. The random process of sampling individuals can result in a population with only one type of allele.



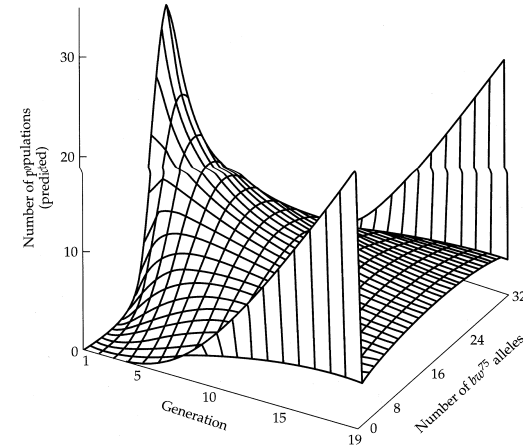
from: (Hartl and Clark, 1997, p. 271)

## Genetic Drift (2)



from: (Hartl and Clark, 1997, p. 274)

## Genetic Drift (3)

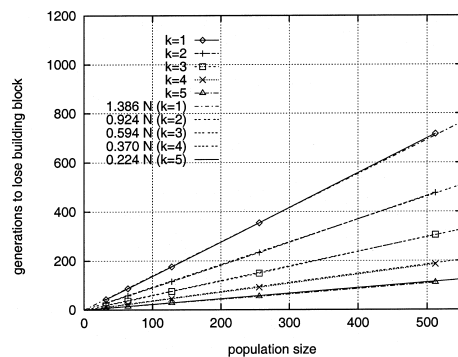


from: (Hartl and Clark, 1997, p. 281)

## Genetic Drift (4)

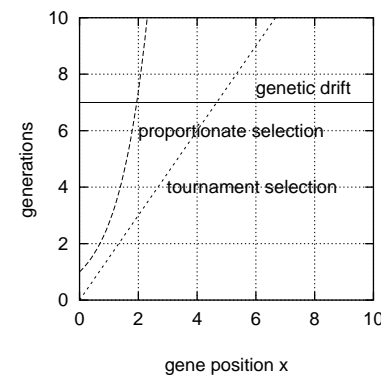
The drift time – using random sampling with replacement – in GAs is proportional to the population size  $N$ :

$t_{drift} = cN$ ,  
where  $c$  depends on the initial proportion.



from: (Lobo et al., 2000)

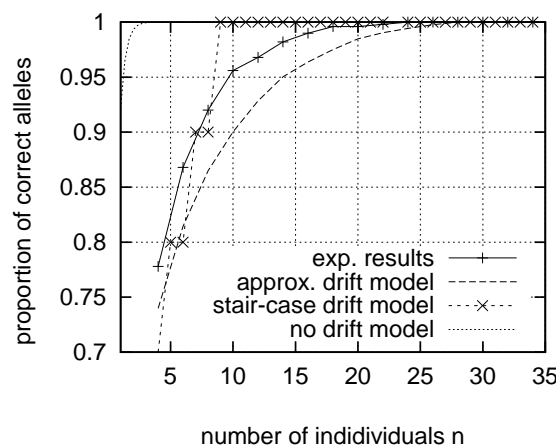
## Genetic Drift and Domino Convergence



from: (Thierens et al., 1998)

Combining domino convergence and drift models:

- Drift models predict a constant generations upper boundary.
- Lower salient alleles are fixed randomly due to genetic drift.

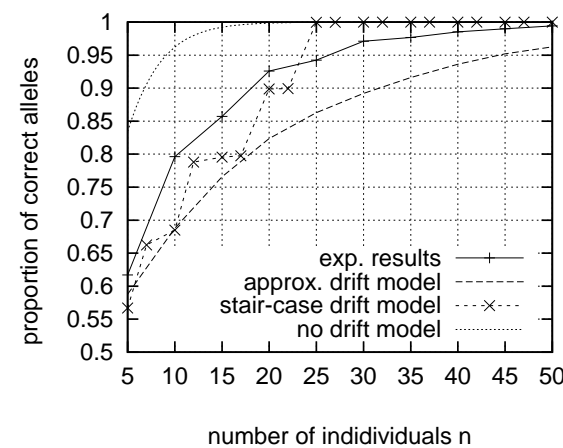


$m = 1, l_s = 5$

Simple GA with uniform crossover, no mutation, and tournament selection of size 2 without replacement.

Approx. drift model: Population sizing model based on the drift model of (Kimura, 1964).

Stair-case drift model: Solving time for one exp. scaled allele  $t^{exp}$  stays constant. Unsolved alleles remain in their initial state for  $t < t_{drift}$ .



$m = 10, l_s = 5$

Simple GA with uniform crossover, no mutation, and tournament selection of size 2 without replacement.

Approx. drift model: Population sizing model based on the drift model of (Kimura, 1964).

Stair-case drift model: Solving time for one exp. scaled allele  $t^{exp}$  stays constant. Unsolved alleles remain in their initial state for  $t < t_{drift}$ .

## Exponentially scaled Representations - Summary

- Representations using exponentially scaled alleles change the dynamics of selectorecombinative GAs.
  - Exponentially scaled representations allow to find rough approximations after short time.
  - Uniformly scaled representations allow to find the best solution in shorter overall time.
- Due to genetic drift GAs using exponentially scaled representations need a larger population size.

## Take home message

- **Representations are important!**
- Representations change the difficulty of a problem!
- Distinguish carefully between genotypes and phenotypes!
- Representations that are non-synonymously redundant are no good idea.
- Synonymously redundant representations can help you if you have problem-specific knowledge!
- Representations should have high locality if you want to solve **easy** problems.
- (Scaling of alleles changes dynamics of search. Non-uniformly scaled alleles are fast, but inaccurate.)

Thanks for your attention and patience!

Further reading:

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Second edition available in a few weeks!

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