# Facetwise Modeling of Genetic Algorithms

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# **Run Time Complexity**

- In typical application the total run time of a genetic algorithm is determined by the number of fitness function evaluations.
- Run time of selection algorithm and variation operators can be ignored.
- Number of fitness function evaluations is equal to the number of generations times the population size:

 $\#FitnessFct.Evals = \#Generations \times PopulationSize$ 

### Convergence speed

- Rate at which a population converges is determined by the selection pressure:
  - high selection pressure: fast convergence
  - low selection pressure: slow convergence
- Size of population determines quality of solution found:
  - large population size: more reliable convergence
  - small population size: less reliable convergence
- Trade-off between selection pressure and population size

# Key questions

- How long does a GA with a certain selection pressure runs before it converges?
- What is the minimal population size to ensure reliable convergence?

#### → problem dependent, but:

- We can build analytical models for simple problems,
- Use this as an approximation for some real, complex problems,
- Gives insight in and guidance for designing performant GAs.

### Models

- First, we will build analytical models for the convergence behavior, assuming large enough populations,
- Second, we will build analytical models for the minimal required population size,
- **3** Third, we will test the models on a real, complex problem (map labeling).

# **Selection Intensity**

- To quantify the speed of convergence we need a quantitative measure of selection pressure.
- The selection differential S(t) is the difference between the mean fitness of the parent population at generation t and the population mean fitness at generation t.
- The selection intensity I(t) is the scaled selection differential, obtained by dividing by the standard deviation of the fitness values.
- *I*(*t*) is dimensionless since the standard deviation has the units in which the selection differential is expressed:

$$I(t) = \frac{S(t)}{\sigma(t)} = \frac{\overline{f}(t^s) - \overline{f}(t)}{\sigma(t)}.$$

### Counting Ones fitness function

• Counting Ones, 'fruit fly' of GA theory

$$CO(X) = \sum_{i=1}^{\ell} x_i \qquad x_i \in \{0, 1\}$$

- Probability having 1 at a certain locus: p(t)
- Fitness binomial distributed
- Mean fitness at generation  $t : \bar{f}(t) = l.p(t)$
- Variance at gen.  $t : \sigma_p^2(t) = l.p(t)(1 p(t))$
- Recombination makes no change to population mean fitness
  - $\Rightarrow$  simple, yet accurate convergence models

### **Proportionate selection**

- Probability selecting i (fitness  $f_i$ , proportion  $P_i(t)$ ):  $P_i(t^s) = P_i(t) \frac{f_i}{f(t)}$
- Selection Differential S(t):

$$\overline{f(t^s)} - \overline{f(t)} = \sum_{i=1}^{N} P_i(t^s) f_i - \overline{f(t)}$$

$$= \sum_{i=1}^{N} P_i(t) \frac{f_i^2}{\overline{f(t)}} - \overline{f(t)}$$

$$= \frac{1}{\overline{f(t)}} (\overline{f^2(t)} - (\overline{f(t)})^2)$$

$$= \frac{\sigma^2(t)}{\overline{f(t)}}$$

• Selection intensity  $I(t) = \frac{\sigma(t)}{\overline{f(t)}}$ 

# **Proportionate Selection: Counting Ones**

- mean fitness increase:  $\overline{f(t+1)} \overline{f(t)} = \frac{\sigma^2(t)}{\overline{f(t)}}$
- proportion of optimal alleles p(t)

$$p(t+1) - p(t) = \frac{1}{l}(1 - p(t))$$

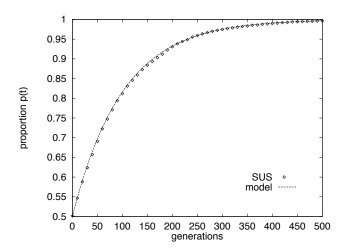
$$\frac{dp(t)}{dt} \approx \frac{1}{l}(1 - p(t))$$

• convergence model (p(0) = 0.5)

$$p(t) = 1 - 0.5e^{-t/l}$$

• convergence speed:  $p(t_{conv}) = 1 - 1/(2\ell)$ 

$$t_{conv} = \ell \ln \left( \ell \right)$$



#### **Truncation Selection**

• Truncating a normal distribution at the top  $\tau\%$  gives fitness increase proportional to the standard deviation:

$$f(t^s) - f(t) = c(\tau) \cdot \sigma(t)$$

- Selection intensity:  $I(\tau) = c(\tau)$
- Values of selection intensity I for truncation selection are constant:

$\tau$	1%	10%	20%	40%	50%	80%
I	2.66	1.76	1.2	0.97	0.8	0.34

### **Truncation Selection**

mean fitness increase

$$\overline{f(t+1)} - \overline{f(t)} = I \ \sigma(t)$$

proportion of optimal alleles p(t)

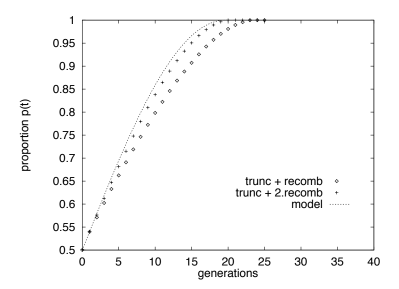
$$p(t+1) - p(t) = \frac{I}{\sqrt{l}} \sqrt{p(t)(1 - p(t))}$$
$$\frac{dp(t)}{dt} \approx \frac{I}{\sqrt{l}} \sqrt{p(t)(1 - p(t))}$$

• convergence model (p(0) = 0.5)

$$p(t) = 0.5(1 + \sin\left(\frac{1}{\sqrt{l}}t\right))$$

• convergence speed ( $p(t_{conv}) = 1$ )

$$t_{conv} = rac{\pi}{2} rac{\sqrt{l}}{l}$$



#### **Tournament Selection**

- Tournament size *s*: the selection intensity *i* is equal to the expected value of the best ranked individual of a sample from *s* individuals taken from the standard normal distribution:
- Can be computed using order statistics:  $I = u_{s:s}$

S	2	3	4	5	6	7
$I=u_{s:s}$	$\frac{1}{\sqrt{\pi}} = 0.56$	0.85	1.03	1.16	1.27	1.35

#### **Tournament Selection**

- Same model as truncation selection, for instance for tournament size s = 2:
- mean fitness increase

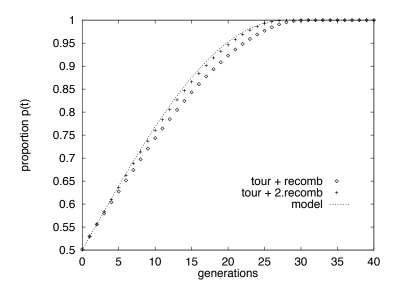
$$\overline{f(t+1)} - \overline{f(t)} = I \ \sigma(t) = \frac{1}{\sqrt{\pi}} \ \sigma(t)$$

• convergence model (p(0) = 0.5)

$$p(t) = 0.5(1 + \sin\left(\frac{t}{\sqrt{\pi l}}\right))$$

• convergence speed ( $p(t_{conv}) = 1$ )

$$t_{conv} = \frac{\pi}{2} \sqrt{\pi l}$$



# Population sizing

- Correct size of the population important:
  - too small: premature convergence to sub-optimal solutions
  - too large: computational inefficient
- We focus on the Counting-Ones problem, but the model can be extended to (slightly) more complex functions
- Key question: how does the optimal population size scales with the complexity of the problem, ie. the length of the string?

### Selection Error

Tournament selection:

```
s_1: 1100011100, fitness = 5

s_2: 0100111101, fitness = 6

\Rightarrow string s_2 is selected!
```

 Competition at the schema level: (order-1 sufficient since we focus on Counting-Ones)

```
partitions ****f *** * and ********f:
schema ****1 **** wins from schema ****0 ****, and
schema ********* 1 wins from schema ********

\Rightarrow correct selection decisions.
```

other partitions: nothing changes.

#### Selection Error

- What is the probability of making a selection error?
- How many selection errors can we afford to make before the optimal bit-value at a <u>cdertain</u> position is completely lost in the population = <u>premature convergence</u>?
- Population sizing is basically a statistical decision making problem.

### Probability selection decision error

• Schemata fitness  $f(H_1: ***1 *** *** **)$  and  $f(H_2: ***0 *** ****)$  binomial distributed  $\rightarrow$  approximating with normal distribution  $\mathbb{N}(\mu, \sigma^2)$ :

$$\mu_{H_1} = 1 + (\ell - 1)p,$$
  $\sigma_{H_1}^2 = (\ell - 1)p(1 - p)$   $\sigma_{H_2}^2 = (\ell - 1)p(1 - p)$ 

(p = probability of having a bit value 1 at any position).

• Distribution of the fitness difference of the best schema and the worst schema  $f(H_1) - f(H_2)$  is also normal distributed:

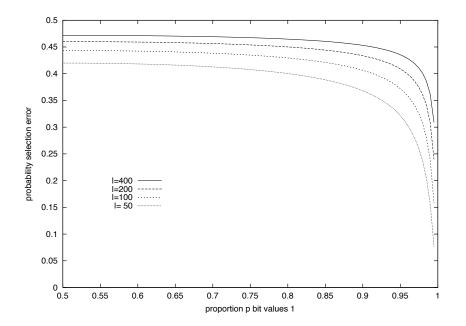
$$\mu_{H_1-H_2} = 1,$$
  $\sigma_{H_1-H_2}^2 = 2(\ell-1)p(1-p)$ 

### Probability selection decision error

 Probability selection error is equal to the probability that the best schema is sampled by a string with fitness less than the sample of the worst schema, which is equal to the probability that the fitness difference of the strings is negative:

$$\begin{array}{lcl} P[SelErr] & = & P(F_{H_1 - H_2} < 0) \\ & = & \Phi(\frac{-1}{\sqrt{2(\ell - 1)p(1 - p)}}) \end{array}$$

- $\Phi(x)$ : Cumulative distribution function of the standard normal distribution.
- $P(X < b) = \Phi(\frac{b-\mu}{\sigma})$



### Probability selection decision error

### Approximation

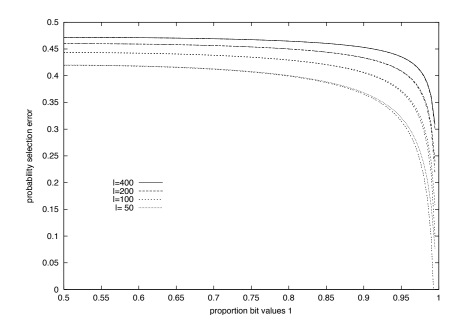
• Approximation by first two terms of power series expansion for the normal distribution:

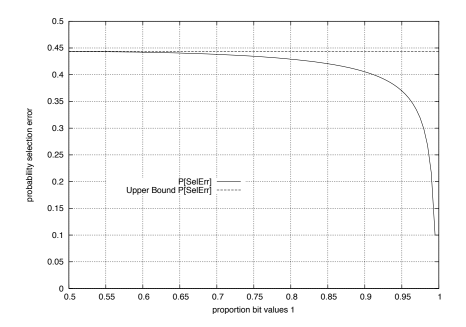
$$P[SelErr] \approx \frac{1}{2} - \frac{1}{2\sqrt{\pi(\ell-1)p(1-p)}}$$

• Selection error is upper bounded by:

$$\left| P[SelErr] \le \frac{1}{2} - \frac{1}{\sqrt{\pi\ell}} \right|$$

this is a conservative estimate of the selection error that ignores the reduction in error probability when the proportion of optimal bit values p(t) increases.





# GA population sizing

### How many selection errors?

- Selection viewed as decision making process within partitions: schemata competition.
- When best schema looses competition we have a selection decision error.
- How many decision errors can we afford to make given a certain population size?
- Answer given by Gambler's ruin model: within each partition a random walk is played.

### Gambler's ruin random walk model

- one-dimensional, discrete space of size N + 1.
- one particle at position  $x \in \{0, ..., N\}$ .
- the particle can move one step to the right with probability p, and one step to the left with probability 1 p.
- when the particle reaches the boundaries (x = 0, or x = N) the random walk ends.
- call  $P_N(x)$  (resp.  $P_0(x)$ ) the probability that the particle is absorbed by the boundary x = N (resp, x = 0) when it is currently at position x



### Gambler's ruin random walk model

• Difference equation:

$$P_N(x) = pP_N(x+1) + (1-p)P_N(x-1)$$

with boundary conditions:  $P_N(N) = 1$ , and  $P_N(0) = 0$ 

• Probability the particle - starting from position  $x_0$  - is absorbed by the x = N boundary:

$$P_N(x_0) = \frac{1 - (\frac{1-p}{p})^{x_0}}{1 - (\frac{1-p}{p})^N}$$

- $P_0(x_0) = 1 P_N(x_0)$
- when p = 1 p = 0.5 we get  $P_N(x_0) = \frac{x_0}{N}$

### Gambler's ruin model (GR) $\rightarrow$ GA

- Position *x* in GR:
  - $\rightarrow$  the number of optimal bit values '1' in the population at a certain partition (position in the string).
- Boundaries x = N (resp. x = 0) in GR:
  → all bit values in the population at the partition are equal to the bit value '1' (resp, '0').
- Absorbing boundary states in GR:
   → population converged to all ones or all zeroes at that partition.
- Probability *p* particle moves one step to the right in GR:
   → probability that the number of optimal bit values 1 in the population at the partition is increased by one = probability correct selection decision.
- Convergence to x = N (resp. x = 0) boundary:
   → Population converges to optimal bit value 1
   (resp. converges to wrong bit value = premature convergence).

- Recall probability selection decision error:  $P[SelErr] \leq \frac{1}{2} \frac{1}{\sqrt{\pi\ell}}$
- Probability convergence to the optimal bit value:

$$\begin{split} P[OptConv] &= \frac{1 - \left(\frac{P[SelErr]}{1 - P[SelErr]}\right)^{N/2}}{1 - \left(\frac{P[SelErr]}{1 - P[SelErr]}\right)^{N}} \\ &\approx 1 - \left(\frac{P[SelErr]}{1 - P[SelErr]}\right)^{N/2} \\ &\approx 1 - \left(\frac{\frac{1}{2} - \frac{1}{\sqrt{\pi\ell}}}{\frac{1}{2} + \frac{1}{\sqrt{\pi\ell}}}\right)^{N/2} \\ &\approx 1 - \left(\frac{\sqrt{\pi\ell} - 2}{\sqrt{\pi\ell} + 2}\right)^{N/2} \end{split}$$

Approximation: denominator approaches 1 much more rapidly as the numerator since P[SelErr] < 1 - P[SelErr]

• Taking the logs:

$$\frac{N}{2} \ln \frac{\sqrt{\pi \ell} - 2}{\sqrt{\pi \ell} + 2} \approx \ln(1 - P[OptConv])$$

• Using the Taylor series approximation:

$$\ln \frac{x-2}{x+2} = \ln(x-2) - \ln(x+2)$$

$$\approx (\ln x - \frac{2}{x} - \frac{2}{x^2}) - (\ln x + \frac{2}{x} - \frac{2}{x^2})$$

$$\approx -\frac{4}{x}$$

we get:

$$\frac{N}{2} \frac{-4}{\sqrt{\pi \ell}} \approx \ln(1 - P[OptConv])$$

• Critical population size:

$$N \approx \ln(1 - P[OptConv]) \frac{\sqrt{\pi \ell}}{-2}$$

The minimal required population size scales as the square root of the problem complexity!

• Probability optimal bit value will be found at certain position:

$$P[OptConv] \approx 1 - e^{\frac{-2N}{\sqrt{\pi\ell}}}$$

# Convergence string length $\ell$

• The number of optimal bits F in the entire string of length  $\ell$  is binomially distributed:

$$P(F = x) = \binom{\ell}{x} P[OptConv]^{x} (1 - P[OptConv])^{\ell - x}$$

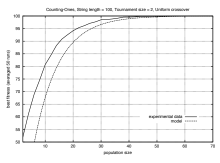
with mean:  $\mu = \ell P[OptConv]$ and variance:  $\sigma^2 = \ell P[OptConv](1 - P[OptConv])$ ,

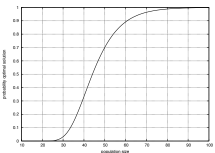
• The probability the optimal string will be reached is:

$$P[OptimalString] = P[OptConv]^{\ell}.$$

# Experimental validation

$$\begin{split} E[Fitness] &= 100~(1-e^{\frac{-2N}{\sqrt{100\pi}}}) \\ P[OptimalString] &= (1-e^{\frac{-2N}{\sqrt{100\pi}}})^{100} \end{split}$$





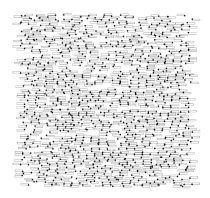
# Map Labeling problem



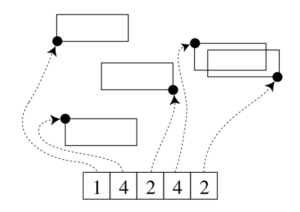
- Place labels next to map features.
- Even basic instances are NP-hard.
- Numerous cartographic rules need to be considered.

# Basic map-labeling problem

- Set of points in the plane.
- Each point has rectangular fixed sized label at 4 possible positions.
- Find a labeling with maximum number of non-overlapping labels.

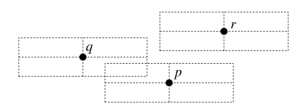


# Encoding



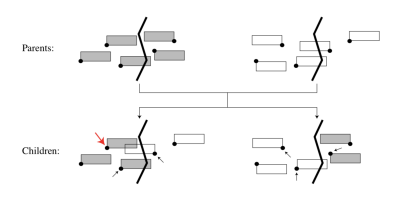
# Rival Groups

- Two points are rivals if their labels can overlap.
- A point together with its rivals is called a rival group.



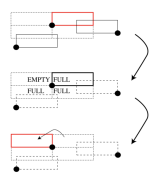
# Crossover on Rival Groups

- Crossover is done by repeatedly choosing rival groups.
- Crossover is complementary: half of a parent is copied to a child and the other half is copied from the other parent.

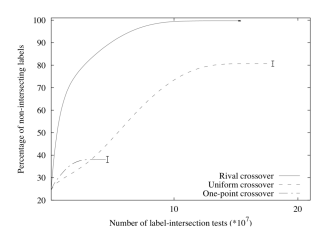


### Geometric Local Search: slot filling

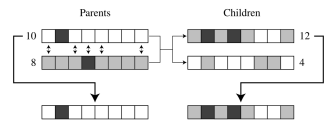
• After crossover a geometrically local optimizer is applied to points which may have a conflict.



### Rival Crossover



### **Elitist Recombination**



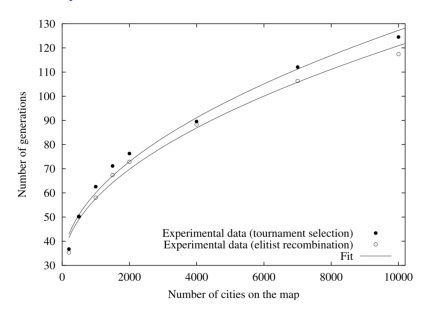
Best two of family replace parents.

# Scalability

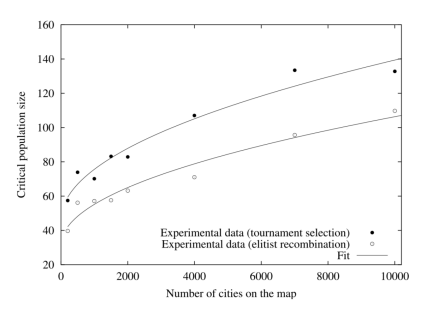
- $Cost(Eval) = O(\ell)$ : each city can be checked in constant time.
- $PopSize = O(\sqrt{\ell})$ : If gambler's ruin model is applicable.
- *Generations* =  $O(\sqrt{\ell})$ : If convergence model is applicable.
- $RunTime = O(\ell^2)$

 $RunTime = Cost(Eval) \times PopSize \times Generations$ 

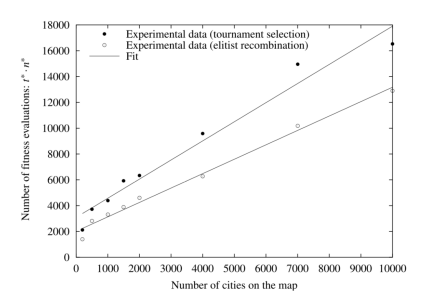
### Scalability Number of Generations



# Scalabiilty Minimal Population Size



### Scalability Number of Fitness Evaluations



# Modeling applicable?

### Assumptions of models are satisfied:

- Fitness function can be kept simple (uniformly scaled, semi-separable, and additively decomposable).
- Crossover is linkage-respecting and mixes well.
- Disruption is minimized by the geometrically local optimizer.

#### Bottom line

Theoretical insights can be used to design efficient genetic algorithms for real-world problems.