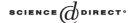


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# Real royal road functions for constant population size \( \frac{1}{2} \)

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#### **Abstract**

Evolutionary and genetic algorithms (EAs and GAs) are quite successful randomized function optimizers. This success is mainly based on the interaction of different operators like selection, mutation, and crossover. Since this interaction is still not well understood, one is interested in the analysis of the single operators. Jansen and Wegener [Proceedings of GECCO'2001, 2001, pp. 375–382] have described so-called real royal road functions where simple steady-state GAs have a polynomial expected optimization time while the success probability of mutation-based EAs is exponentially small even after an exponential number of steps. This success of the GA is based on the crossover operator *and* a population whose size is moderately increasing with the dimension of the search space. Here new real royal road functions are presented where crossover leads to a small optimization time, although the GA works with the smallest possible population size—namely 2.

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# 1. Introduction and history

Genetic algorithms (GAs) and evolution strategies (ESs) have many areas of application. Here we consider the maximization of pseudo-boolean functions  $f_n: \{0,1\}^n \to \mathbb{R}^+_0$ . The success of GAs and ESs depends on the interaction of the different operators, among them the so-called search (or genetic) operators which create new individuals

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from existing ones. A search operator working on one individual is called mutation and a search operator working on two (or more) individuals is called crossover. We only investigate the best-known crossover operators namely one-point crossover and uniform crossover. There have been long debates whether mutation or crossover is "more important". This paper does not contribute to this debate. Our purpose is to investigate when and why crossover is essential.

The problem of premature convergence and the problem to maintain diversity in the population are well known. There are many ideas how to cope with these problems: multi-starts, fitness sharing, niching, distributed GAs, and many more. They all have shown their usefulness in experiments. It has also been possible to analyse highly specialized GAs on some functions. However, the rigorous analysis of GAs is still in its infancy. This motivates the investigation of special properties of single operators (like the takeover time).

The success of GAs is based on the use of populations and the use of crossover operators. Holland [3] has discussed why crossover is a good search operator. This has led to the building-block hypothesis and the schema theory (see also [2]). We are interested in a rigorous analysis and we concentrate on the following parameters:  $T_{A}$  f describes the random number of fitness evaluations until the algorithm A evaluates an optimal search point for f. The expected optimization time  $E(T_{A,f})$  is the expected value of  $T_{A,f}$  and the success probability function  $t \rightarrow \text{Prob}(T_{A,f} \leq t)$  describes the probability of a successful search within a given number of steps. An algorithm is called *efficient* if the expected optimization time is polynomially bounded (with respect to the problem dimension n) or if at least the success probability within a polynomial number of steps converges to 1 (with respect to n). Crossover is essential for a sequence  $f = (f_n)$  of fitness functions if a simple GA (without specialized modules) is efficient while all mutation-based evolutionary algorithms (EAs) are not efficient. Mitchell et al. [8] have looked for such functions and presented the so-called royal road functions  $RR_{n,k}: \{0,1\}^n \to \mathbb{R}_0^+$ . The input string a is partitioned to m blocks of length k each where m = n/k is an integer. Then  $RR_{n,k}(a)$  equals the number of blocks of a containing ones only. This is a nice example function since it seems to be a "royal road" for the building-block hypothesis and the application of one-point crossover. However, Mitchell et al. [9] (see also [7]) have shown that this intuition is wrong. Crossover is not essential when optimizing the royal road functions. Other "GAfriendly" functions like H-IFF (see, e.g., [11]) have been presented but their analysis is not rigorous. Some rigorous analysis has been performed for highly specialized GAs (see, e.g., [1]). Jansen and Wegener [4] have presented so-called real royal road functions for uniform crossover and for one-point crossover and proved rigorously that the expected optimization time of a simple steady-state GA is polynomially bounded while each mutation-based EA needs exponential time until the success probability is not exponentially small. The results hold for populations of size n but not for populations whose size is independent of n.

Hence, the steady-state GA needs a population of moderate size *and* the appropriate crossover operator to be efficient. This raises the question of whether populations without crossover can be essential and the question of whether crossover needs populations whose size grows with n in order to be essential. Jansen and Wegener [5]

have presented functions where mutation-based EAs working with large populations are efficient while all mutation-based EAs with population size 1 are not efficient. Here we answer the second question by presenting *real royal road functions for populations of size* 2.

More precisely, we describe in Section 2 a steady-state GA (called (2+1)GA) working with the smallest possible population size allowing crossover, namely population size 2. This GA is not specialized. It only guarantees that the population contains two different individuals. In Section 3, the real royal road function for uniform crossover and constant population size is presented. It is proven that the (2+1)GA is efficient in the sense, that the success probability after a polynomial number of steps is 1 - o(1), i.e., converging to 1 as  $n\rightarrow\infty$  (Theorem 4). Each mutation-based EA needs exponential time until the success probability is not exponentially small (Proposition 1). In Section 4, the function is changed into a real royal road function for one-point crossover and population size 2. The (2+1)GA needs only a polynomial number of steps to reach a success probability of 1-o(1) (Theorem 6), but the expected optimization time grows exponentially. The reason is that with small probability some bad event happens. This event implies a very large optimization time leading to the large expected optimization time. Therefore, we present in Section 5 a strong real royal road function for one-point crossover and population size 2. For this function the (2+1)GA even has a polynomially bounded expected optimization time (Theorem 11) while mutation-based EAs still need exponential time until the success probability is not exponentially small (Proposition 7). We finish with some conclusions.

# 2. The steady-state (2+1)GA

We describe a simple steady-state GA working on the smallest possible population size allowing crossover namely population size 2.

(2+1)GA

*Initialization*: Randomly choose two different individuals  $x, y \in \{0, 1\}^n$ .

Search: Produce an individual z, more precisely,

- with probability 1/3, z is created by mutate(x),
- with probability 1/3, z is created by mutate(y),
- with probability 1/3, z is created by mutate(crossover(x, y)).

*Selection*: Create the new population  $\mathcal{P}$ .

- If z = x or z = y, then  $\mathscr{P} := \{x, y\}$ .
- Otherwise, let  $a \in \{x, y, z\}$  be randomly chosen among those individuals with the worst f-value. Then  $\mathscr{P} := \{x, y, z\} \{a\}$ .

The reader may wonder why all three possibilities of the search step have probability  $\frac{1}{3}$ . This choice is not essential. Our results hold for all probabilities  $p_1$ ,  $p_2$ ,  $p_3$ , even if they are based on the fitness of x, y, and z as long as they are bounded below by a positive constant  $\varepsilon > 0$ .

We apply the usual mutation operator flipping each bit independently with probability 1/n. In Sections 4 and 5, we apply the usual one-point crossover but create only one child, i.e., choose  $i \in \{1, ..., n-1\}$  randomly and set crossover $(x, y) := (x_1, ..., x_i, y_{i+1}, ..., y_n)$ . Here the order of x and y is chosen randomly. In Section 3, we apply uniform crossover where each bit of the child is chosen from each of the parents with probability  $\frac{1}{2}$ .

## 3. Real royal roads for the (2+1)GA and uniform crossover

Before presenting our new function we emphasize that our purpose is to prove rigorously that the (2+1)GA can outperform all mutation-based EAs. As in many other cases, the first example functions with certain properties are artificial fitness functions designed only to prove the results under considerations.

Our example function has one area of global optima and two local optima of different fitness. It should be difficult to create a globally optimal point by mutation of one of the local optima, but it should be easy to do so by uniform crossover of the two local optima. Hence, the GA has to realize a population consisting of the local optima. The fitness function gives hints to reach the better local optimum first. Since we have always two different individuals, one of them is only close to the local optimum and gets hints to look for the second local optimum. These ideas are now made precise.

To simplify the notation we assume that m := n/6 is an even integer. Let |x| be the length of x,  $||x|| := \text{ONEMax}(x) := x_1 + \cdots + x_n$  denote the number of ones of x, and  $0^k$  a string of k zeros. The Hamming distance H(x, y) equals the number of indices i where  $x_i \neq y_i$ . A path is a sequence  $a_1, \dots, a_p$  such that  $H(a_i, a_{i+1}) = 1$  and the points  $a_i$  are pairwise distinct. The definition of the new real royal road function  $R_n^u$  (u indicates that we use uniform crossover) is based on a path P and a target region T. The path  $P = (a_0, \dots, a_{7m})$  contains 7m + 1 search points: For  $i \le 6m$  let  $a_i := 0^{n-i}1^i$  and for i = 6m + j let  $a_i := 1^{n-j}0^j$ . Let  $R_n^u(a_i) := n + i$  for all  $i \neq 5m$  and  $R_n^{\mathrm{u}}(a_{5m}) := n + 8m$ . This implies that we have two local optima on P, namely  $a^* := a_{5m}$ and  $a^{**} := a_{7m}$ . If the (2+1)GA first finds  $a^*$ , the second individual can search for  $a^{**}$ . Hence, we like to have a good chance of creating an optimal search point by uniform crossover from  $a^*$  and  $a^{**}$ . Let T contain all points  $b1^{4m}c$  where |b|=|c|=m and ||b|| = ||c|| = m/2. Uniform crossover between  $a^* = 0^m 1^{5m}$  and  $a^{**} = 1^{5m} 0^m$  preserves the 4m ones in the middle part. The probability of creating m/2 ones in the prefix equals  $\binom{m}{m/2} 2^{-m} = \Theta(1/m^{1/2})$  (by Stirling's formula). The same holds independently for the suffix. Hence, the probability that uniform crossover on  $\{a^*, a^{**}\}$  creates a target point and mutation does not destroy this property equals  $\Theta(1/m) = \Theta(1/n)$ . Now we give a complete definition of  $R_n^u$  which is illustrated in Fig. 1. Let  $P_1 := (a_0, \dots, a_{5m-1})$ and  $P_2 := (a_{5m+1}, \dots, a_{7m}).$ 

$$R_n^{u}(x) := \begin{cases} 15m & \text{if } x \in T, \\ 14m & \text{if } x = a^*, \\ 6m + i & \text{if } x = a_i \in P_1 \cup P_2, \\ 6m - ||x|| & \text{if } x \in R := \{0, 1\}^n - P - T. \end{cases}$$

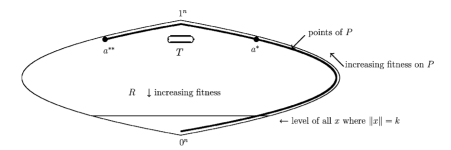


Fig. 1. An illustration of R<sub>n</sub><sup>u</sup>.

**Proposition 1.** Evolution strategies (without crossover) need with probability 1 - o(1) exponentially many steps w.r.t. n to optimize  $R_n^u$ .

We omit the proof of this proposition. The probability to create a target point by mutation from a path point is exponentially small. Hence, one has to search within R for a small target where the fitness function only gives the advice to decrease the number of zeros. This makes it exponentially unlikely to hit T. A complete proof follows the lines of the proof of Proposition 6 of Jansen and Wegener [4].

**Lemma 2.** The probability that the (2+1)GA does not find a search point from  $P_2 \cup T \cup \{a^*\}$  within  $c_1 n^2$  steps is  $2^{-\Omega(n)}$  ( $c_1$  an appropriate constant).

**Proof.** We consider the fitness levels  $L_i$ ,  $1 \le i < 11m$ , containing all search points x where  $R_n^u(x) = i$ . These fitness levels contain all search points outside  $P_2 \cup T \cup \{a^*\}$ . Each of these search points has a better Hamming neighbor. A population  $\mathscr{P} = \{x, y\}$  belongs to  $L_i$  if  $\max\{f(x), f(y)\} = i$ . The probability to leave  $L_i$  is at least the probability of choosing the right individual for the right 1-bit-mutation and, therefore, at least p = 1/(3en). The steps where crossover is chosen can be ignored. They only may increase the success probability. Hence, we have to wait for at most 11m successes in an experiment where each trial has a success probability of at least p. Therefore, the result follows by an application of Chernoff bounds (see, e.g., [10]).  $\square$ 

**Lemma 3.** If the population contains  $a^*$ , the probability that the (2+1)GA does not find an optimal search point, namely a search point from T, within  $c_2n^2$  steps is  $2^{-\Omega(n)}$  ( $c_2$  an appropriate constant).

**Proof.** By the selection procedure, the population will contain  $a^*$  until a search point of T is created. The probability to create some  $a \in P_2$  from  $a^*$  is bounded below by p = 1/(3en). Afterwards,  $\mathscr{P} = \{a^*, a_i\}$ , i > 5m. We now consider the level defined by the index i until  $\mathscr{P} \subseteq \{a^*, a^{**}\} \cup T$ . The probability to increase i is bounded below by p in each step. Finally, the probability to create some target point from the population  $\{a^*, a^{**}\}$  equals by our arguments above  $\Theta(1/n)$ . Moreover, only target points are accepted if  $\mathscr{P} = \{a^*, a^{**}\}$ . Hence, we have to wait for 2m+1 successes in experiments

where each trial has a success probability of  $\Theta(1/n)$ . Therefore, Lemma 3 follows by the same arguments as Lemma 2.  $\square$ 

**Theorem 4.** The success probability that the (2+1)GA with uniform crossover optimizes  $R_n^u$  within  $cn^2$  steps, c an appropriate constant, is 1 - O(1/n).

**Proof.** Applying Lemmas 2 and 3 we are left with the problem whether the (2+1)GA creates a search point from  $P_2$  before a search point from  $T \cup \{a^*\}$  (called bad event). It is sufficient to bound the probability of the bad event by O(1/n). Here we have to cope with the "undesired" effects of uniform crossover. Remember that all search points of  $P_2 \cup T \cup \{a^*\}$  contain 4m ones in their "middle part".

By Chernoff bounds, the probability that the initial population contains a search point with more than  $(\frac{7}{2})m$  ones is exponentially small. As long as no point from  $P \cup T$  is created, we only have search points with at most  $(\frac{7}{2})m$  ones if we start with such strings. Hence, each of the search points has at least m/2 wrong bits in the middle part. The probability to correct a wrong bit by uniform crossover is at most  $\frac{1}{2}$  and the probability to correct it by mutation is 1/n. Hence, it is very likely to create a search point  $a_i$ ,  $i \le (\frac{15}{4})m$ , before creating a point from  $P_2 \cup T \cup \{a^*\} \cup \{a_i \mid (\frac{15}{4})m < i < 5m\}$ (the failure probability is exponentially small). We can repeat these arguments to prove that at some point of time the population contains  $a_i$  and  $a_i$ ,  $0 \le i < j < 4m$ . Uniform crossover applied to  $a_i$  and  $a_j$  creates some  $a_k$ ,  $i \le k \le j$ . Therefore, better points are created by mutation from some  $a_k$ , k < 4m. The probability of creating some point of  $P_2 \cup \{a^*\}$  is exponentially small. Thus, we will obtain in  $O(n^2)$  steps  $a_i$  and  $a_j$ ,  $4m \le i < j < 5m$ , with overwhelming probability. Then better points are created from some  $a_k$ ,  $4m \le k < 5m$ . In this case, there is exactly one (5m - k)-bit mutation to create  $a^*$  and exactly one (5m-k+l)-bit mutation,  $1 \le l \le 2m$ , to create the point  $a_{5m+l}$  from  $P_2$ . The probability of the (5m-k)-bit mutation equals  $q_1 := (1/n)^{5m-k} (1-1/n)^{n-5m+k}$ and the probability of all the (5m-k+l)-bit mutations,  $1 \le l \le 2m$ , altogether is the sum  $q_2$  of all  $(1/n)^{5m-k+l}(1-1/n)^{n-5m+k-l}$ . Since,  $q_2/q_1 = O(1/n)$ , the probability to create a point from  $P_2$  before creating a point from  $T \cup \{a^*\}$  is altogether bounded by O(1/n).

Having this essential result we can play with the definition of  $R_n^u$ . Let  $R_{n,k}^u$  be the variant of  $R_n^u$  where  $a_{5m+1}, \ldots, a_{5m+k}$  belong to the region R. Proposition 1 and Lemma 2 hold also for  $R_{n,k}^u$ . In Lemma 3, we now have to wait for the event to flip the right k+1 bits of  $a^*$  to obtain  $a_{5m+k+1}$ . The expected time for this is bounded above by  $e^{n+1}$ . After  $c_2n^{k+1}\log n$  steps the probability of not creating a point of  $P_2$  is bounded above by

$$\left(1 - \frac{1}{\mathrm{e}n^{k+1}}\right)^{c_2 n^{k+1} \log n} \leqslant \mathrm{e}^{-(c_2/\mathrm{e}) \log n}.$$

This can be made smaller then  $1/n^k$  by choosing  $c_2$  appropriately. However, in the proof of Theorem 4 now  $q_2/q_1 = O(1/n^k)$ , since we need at least (k+1)-bit mutations to create points from  $P_2$ . This leads to the following result.

**Theorem 5.** The success probability that the (2+1)GA with uniform crossover optimizes  $R_{n,k}^u$  within  $cn^{k+1} \log n$  steps, c an appropriate constant, is  $1 - O(1/n^k)$ .

## 4. A variant for one-point crossover

In order to obtain a real royal road function for the (2+1)GA and one-point crossover we can consider an appropriate variant of  $R_n^u$ . The probability of creating  $1^n$  by one-point crossover from  $a^{**}=1^{5m}0^m$  and  $a^*=0^m1^{5m}$  is bounded below by a positive constant. The idea is to make  $1^n$  the only target point which cannot be reached easily by mutation from a point on  $P_2$ . For this reason we replace the path between  $a^*$  and  $a^{**}$  by a path which is far away from  $1^n$ . The function  $R_n^1$  is defined in the same way as  $R_n^u$  with two exceptions. The target set T is replaced by  $T:=\{1^n\}$ . The points  $(a_{5m+1},\ldots,a_{7m})$  are now defined by  $a_{5m+2i-1}:=1^i0^{m-i}1^{5m-i+1}0^{i-1}$  and  $a_{5m+2i}:=1^i0^{m-i}1^{5m-i}0^i$ ,  $1 \le i \le m$ . Then the points of  $P_2$  have 5m or 5m+1 ones and are far from  $1^n$  and far from the typical points created in the initialization step. Thus Proposition 1 also holds with the same arguments for  $R_n^1$ .

**Theorem 6.** The success probability that the (2+1)GA with one-point crossover optimizes  $R_n^1$  within  $cn^2$  steps, c an appropriate constant, is 1 - O(1/n).

**Proof.** It is easy to prove the claims of Lemmas 2 and 3 also for  $\mathbb{R}_n^1$ . The old proofs can be used without changes. If the population consists of  $a_i$  and  $a_j$ ,  $0 \le i < j < 4m$ , we can use the arguments of the proof of Theorem 4 to conclude that the probability of creating a point from  $P_2$  before creating a point from  $T \cup \{a^*\}$  is bounded by O(1/n). Let x and y be the search points of the initial population. All search points from  $P_2$  have 4m ones in their middle part. By Chernoff bounds, the probability that x and y have not at least m/2 common zeros in this middle part is exponentially small. In order to obtain a search point from  $P_2$  it is necessary to flip each of these positions at least once. In a phase of cn steps the probability of not flipping a bit at a special position equals  $(1-1/n)^{cn}$  and the probability of flipping it is therefore  $1-(1-1/n)^{cn}$  and thus bounded above by a constant  $\varepsilon < 1$ . Since the bit positions are treated independently by the mutation operator, the probability of flipping m/2 given positions is exponentially small. (These calculations are related to the coupon collector's theorem (see, e.g., [10]).) If this event does not happen we are in the situation of the fitness function 6m - ||x|| = n-ONEMAX(x). The standard analysis of ONEMAX leads to the result that we can expect after cn steps a population of two search points of at most m ones. Each step has a constant probability of decreasing the number of ones in the population. By Chernoff bounds, cn steps for an appropriate c are enough to decrease the number of ones from at most 11m ones to at most 2m ones. The failure probability again is exponentially small. If both search points have at most m ones, crossover can only create a search point with 2m ones and even then the probability of creating 4m ones by mutation is exponentially small. Hence, we create some  $a_i$  and  $a_j$ ,  $0 \le i < j < 4m$ , before some point from  $P_2$  with a probability exponentially close to 1.  $\square$ 

With the same notations and arguments leading to Theorem 5 we can prove a similar theorem for  $R_{n,k}^1$ .

## 5. Real royal roads for the (2+1)GA and one-point crossover

The aim is to define a real royal road function  $R_n^{1*}$  for the (2+1)GA and one-point crossover which even has the property that the expected optimization time of the (2+1)GA is bounded by some polynomial p(n). This implies by Markoff's inequality a success probability of at least  $\frac{1}{2}$  within 2p(n) steps. Since the bound on the expected optimization time holds for *all* initial populations, the success probability within  $O(p(n)\log n)$  steps can be bounded by  $1-O(n^{1/k})$  for each constant k.

The definition of  $R_n^{1*}$  is influenced by the function  $R_n^1$ . We modify  $R_n^1$  in the following way. All  $a_i$ , where i=5m+2j-1 and  $1 \le j \le m$ , i=5m+2 or i=5m+4, now belong to the bad region R. Finally, all other individuals  $a_i$ , i>5m, have the same fitness 13m and define a plateau of constant fitness. The full definition of  $R_n^{1*}$  is

$$R_n^{1*}(x) := \begin{cases} 15m & \text{if } x \in T := \{1^n\}, \\ 14m & \text{if } x = 0^m 1^{5m} =: a_0, \\ 13m & \text{if } x \in \{a_i := 1^i 0^{m-i} 1^{4m} 1^{m-i} 0^i \mid 3 \leqslant i \leqslant m\} =: P, \\ 6m + i & \text{if } x = 0^{n-i} 1^i, \ 0 \leqslant i < 5m, \\ 6m - \|x\| & \text{otherwise.} \end{cases}$$

The following result can be proved like Proposition 1.

**Proposition 7.** Evolution strategies (without crossover) need with a probability exponentially close to 1 exponentially many steps w.r.t. n to optimize  $R_n^{1*}$ .

Finally, we analyse the (2+1)GA with one-point crossover on  $R_n^{1*}$ .

**Lemma 8.** The expected time until the population of the (2+1)GA contains a search point from  $Q := P \cup T \cup \{a_0\}$  is bounded by  $O(n^2)$ . With a probability of  $1 - O(1/n^6)$  this search point is contained in  $T \cup \{a_0\}$ .

**Proof.** As in the proof of Theorem 6 we get an expected time of  $O(n^2)$  for creating a search point from Q. Again we have to create with overwhelming probability the search point from Q by mutation of some  $0^{n-i}1^i$ ,  $0 \le i < 5m$ . The probability that this happens for some  $i \le 4m$  is exponentially small. Otherwise,  $H(0^{n-i}1^i, a_3) = H(0^{n-i}1^i, a_0) + 6$  and we have to compare k-bit mutations with (k + 6)-bit mutations. Hence, we can apply the same arguments as in the proof of Theorem 5.  $\square$ 

Let us first consider what happens if the population contains only individuals of P. Remember that all these individuals have the same fitness.

**Lemma 9.** If the population contains only search points from P, the expected time until the population of the (2+1)GA contains an element from  $T \cup \{a_0\}$  is bounded by  $O(n^9)$ .

	I	II	III	IV	V
$a_i$ $a_j$ $a_i$	$1 \cdots 1$	$1 \cdots 1$	$0\cdots01\cdots1\\0\cdots01\cdots1\\0\cdots01\cdots1$	$0 \cdots 0$	$0 \cdots 0$
length	i	j-i	n-2j	j-i	i

Fig. 2. One-point crossover between  $a_i$  and  $a_j$  resp.  $a_j$  and  $a_i$ .

**Proof.** We only have to consider the situation where the population consists of  $a_i$  and  $a_j$ ,  $3 \le i < j \le m$ . A step is called *essential* if a search point is created, which has a chance of being accepted for the next population, namely a search point from  $Q - \{a_i, a_j\}$ . The probability of producing  $a_{i \pm d}$ ,  $d \ge 1$ ,  $3 \le i \pm d \le m$ , from  $a_i$  by mutation equals  $\Theta(1/n^{2d})$ , since we consider a special 2d-bit mutation. In order to investigate mutate(crossover $(a_i, a_j)$ ) resp. mutate(crossover $(a_j, a_i)$ ) we describe the search points in Fig. 2. If the cut position falls into Region I or Region V (including the borders), we obtain  $a_i$  or  $a_j$ , altogether each of them with the same probability. If the cut position falls into Region III (including the borders) we obtain  $1^i0^{m-i}1^{4m}1^{m-j}0^j$  or  $1^j0^{m-j}1^{4m}1^{m-i}0^i$ , for Region II we obtain some  $1^i0^{s-i}1^{j-s}0^{m-j}1^{4m}1^{m-j}0^j$  or  $1^s0^{m-s}1^{4m}1^{m-i}0^i$ , i < s < j, and for Region IV some  $1^i0^{m-i}1^{4m}1^{m-s}0^s$ , or  $1^j0^{m-j}1^{4m}1^{m-j}0^{j-s}1^{s-i}0^i$ , i < s < j. The situation is almost symmetric if  $3 \le i < j \le m$ . More precisely, the following individuals have the same chance to be created by crossover and a following mutation from the pair  $(a_i, a_j)$ :

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-a_{i-d} and a_{j+d} (as long as i-d \ge 3 and j+d \le m) and -a_{i+d} and a_{j-d} (as long as i+d \le j-d).
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And at least one bit has to flip to obtain an element from  $Q - \{a_i, a_j\}$  from crossover( $a_i$ ,  $a_j$ ) resp. crossover( $a_j, a_i$ ). Hence, the total probability of an essential step is bounded by O(1/n) and  $\Omega(1/n^2)$ . In order to prove the lemma it is sufficient to show a bound of  $O(n^7)$  on the number of essential steps until  $a_0$  or  $1^n$  is produced.

If i = 3, the probability that the next essential step produces  $a_0$  by mutation from  $a_i$  is bounded by  $\Omega(1/n^5)$ . Hence, the expected number of essential steps with a population containing  $a_3$  until  $a_0$  is included in the population is bounded by  $O(n^5)$ . We are done by proving a bound of  $O(n^2)$  on the number of essential steps until the population contains  $a_3$ ,  $a_0$ , or  $1^n$  if we start with  $a_i$  and  $a_j$ ,  $3 \le i < j \le m$ .

For this purpose, it is sufficient to prove that for a phase of  $cn^2$  essential steps, c an appropriate constant, there is a probability of at least a constant  $\varepsilon > 0$  that we produce  $a_3$ ,  $a_0$ , or  $1^n$ . We ignore the chance of producing  $a_0$  or  $1^n$ . Let  $\{a_{i'}, a_{j'}\}$  be the population created from  $\{a_i, a_j\}$  in an essential step. The *gain* of this step is defined by (i'+j')-(i+j). To produce  $a_{i-d}$  or  $a_{j+d}$ ,  $d \ge 2$ , mutation alone has to flip 2d bits and the mutation following a crossover at least 2d bits. Hence, this happens with

a probability of  $\Theta(1/n^2)$  in an essential step. There is a probability of  $(1-\Theta(1/n^2))^{cn^2} \le \varepsilon'$  where  $\varepsilon' < 1$  is a constant that the phase does not contain such a step. This implies that  $\{a_i, a_j\}$  can create only  $a_k$ ,  $i-1 \le k \le j+1$ . Further, the probability of good steps of length 1  $(a_i \rightarrow a_{i-1}, a_j \rightarrow a_{j-1})$  is the same as the probability of bad steps of length 1  $(a_i \rightarrow a_{i+1}, a_j \rightarrow a_{j+1})$ . This even holds for steps of length d,  $d \ge 2$   $(a_j \rightarrow a_{j-d}, a_i \rightarrow a_{i+d})$ . However, if we produce  $a_j$  from  $a_i$ , we cannot accept the copy of  $a_j$ . This does not disturb the symmetry since with the same probability we produce  $a_i$  from  $a_j$ . If j = m,  $a_{j+1}$  does not exist. This disturbs the symmetry but supports a positive gain.

The steps of length 1 can be considered as independent coin tosses with success probability  $\frac{1}{2}$ . Applying Chernoff bounds,  $\Theta(n^2)$  coin tosses are enough to have a surplus of n wins with a probability of at least a constant  $\varepsilon'' > 0$ . For each fixed point of time the probability that the steps of length d,  $d \ge 2$ , lead to a non-negative gain is by symmetry at least  $\frac{1}{2}$  (it is not exactly  $\frac{1}{2}$ , since the total gain of these steps may be 0). This implies that we create  $a_3$  with a probability of at least  $\varepsilon$  in one phase if c is chosen large enough. (Since the search points  $a_3, \ldots, a_m$  are a plateau of constant fitness of  $R_n^{1*}$ , we have used some ideas contained in the paper of Jansen and Wegener [6].)

**Lemma 10.** If the population contains  $a_0$  and some  $a_j$ ,  $3 \le j \le m$ , the probability for the (2+1)GA that the population contains  $1^n$  or  $a_0$  and  $a_m$  within the next  $cn^4$  steps, c an appropriate constant, is bounded below by some constant  $\varepsilon > 0$ .

**Proof.** We can apply many ideas of the proof of Lemma 9. The probability of creating some  $a_k$ ,  $3 \le k \le m$ , by mutation of  $a_0$  is bounded by  $O(1/n^6)$ . Further, the probability of producing  $a_{j\pm d}$ ,  $d \ge 2$ , by mutation of  $a_j$  is bounded by  $O(1/n^4)$ . Let us investigate the effect of crossover, namely mutate(crossover( $a_0, a_j$ )) and mutate(crossover( $a_j, a_0$ )). We are in the same situation as in Fig. 2 but now i=0. Thus, Regions I and V are empty. If the cut position falls into Region III (including the borders) we obtain  $0^m 1^{4m} 1^{m-j} 0^j$  or  $1^j 0^{m-j} 1^{4m} 1^m$ . To produce  $a_{j\pm d}$ ,  $d \ge 1$ , the first (or the last)  $3 \le j \pm d \le m$  bits of the string and d of the last (or the first) m bits must flip by the mutation following crossover. For each of these cut positions, this leads to a probability of  $O(1/n^4)$  for a successful crossover. If the cut position falls into Region II, we obtain some  $0^m 1^{4m} 1^m 0^{-j} 0^j$  (II.a) or  $1^s 0^{m-s} 1^{4m} 1^m$  (II.b), 0 < s < j, and for Region IV we obtain some  $0^m 1^{4m} 1^m 0^{-s} 0^s$  (IV.a) or  $1^j 0^{m-j} 1^{4m} 1^m 0^{-j} 0^{-s} 1^s$  (IV.b), 0 < s < j. For (II.a) and (IV.b) we distinguish the cases  $s \ge 2$  and s = 1.

Case 1:  $s \ge 2$ . To obtain  $a_{j\pm d}$  at least four bits have to flip in the mutation following crossover:

- the first 2 resp. last 2 ones, since  $s \ge 2$ ,
- at least one other *special* bit of the first resp. last m positions, since  $d \ge 1$ ,
- at least  $d \ge 1$  special bits of the last resp. first m positions.

This again leads for each of these cut positions to a probability of  $O(1/n^4)$  for a successful mutate–crossover step.

Case 2: s = 1. Here we can guarantee only three flipping bits at selected positions. We can use the same arguments but have to take into account that s = 1. However,

the case s = 1 refers to only one cut position of crossover. This leads to a probability of  $O(1/n^3)$  for a successful mutate–crossover step.

For (II.b) and (IV.a) the mutation following a crossover has to flip k + |s - k| bits to generate  $a_k$ . This leads for  $s \ne 3$  to a probability of  $O(1/n^4)$  and for s = 3 to a probability of  $O(1/n^3)$  for a successful mutate–crossover step. Therefore, since each cut position has the same probability of 1/(n-1), altogether we get a probability of

$$\frac{(n-3)\cdot O(1/n^4) + 2\cdot O(1/n^3)}{n-1} = O(1/n^4)$$

for creating some  $a_k$ . Hence, the probability of a successful crossover or a mutation flipping 4 or even more bits is bounded by  $O(1/n^4)$ . The probability of including  $a_m$  in the population can be analysed in the same way as the probability of including  $a_3$  in the proof of Lemma 9.  $\square$ 

**Theorem 11.** The expected time until the (2+1)GA has optimized  $R_n^{1*}$  is bounded above by  $O(n^6)$ .

Proof. By Lemmas 8 and 9, an expected number of

$$(1 - O(1/n^6))O(n^2) + O(1/n^6) \max(O(n^2), O(n^9)) = O(n^3)$$

steps are enough to obtain a population containing  $a_0$  or  $1^n$ . In the second case we are done. Otherwise, the population will contain  $a_0$  until it has optimized  $R_n^{1*}$ . If the second individual of the population does not equal some  $a_j$ ,  $3 \le j \le m$ , a 6-bit mutation will create one from  $a_0$ , what needs an expected time of  $O(n^6)$ . By Lemma 10 the expected time until the population then contains  $a_0 = 0^m 1^{5m}$  and  $a_m = 1^{5m} 0^m$  is bounded by  $O(n^4)$ . Afterwards, the probability of creating  $1^n$  is at least 1/(10e) (choose crossover, its cut position s in the middle, namely  $s \in \{m, \ldots, 5m\}$  and do not mutate any bit). Hence, with a probability of 1 - o(1) we produce  $1^n$  before  $a_m$  is replaced by some  $a_j$ ,  $3 \le j \le m - 1$ . In this case, we can repeat the arguments. The expected number of these phases is 1 + o(1).  $\square$ 

We also have defined a real royal road function for uniform crossover where also the expected time of the (2+1)GA is polynomially bounded. The construction of the function and the analysis of the (2+1)GA are more complicated than the results presented here. The reason is the following. If uniform crossover is applied to two search points a and b with a small Hamming distance, the same result can be obtained with a not too small probability by mutating a or b. If the Hamming distance is large, each point which can be created by uniform crossover from a and b has an exponentially small probability of being created. This differs from one-point crossover where each possible search point has a probability of at least 1/(n-1) to be created. This implies that uniform crossover can be useful only if there are many good search points "between a and b". This again makes it more difficult to control the "undesired effects" of uniform crossover—in particular, in small populations.

#### 6. Conclusions

The question whether crossover without special methods to ensure the diversity of the population and without a population whose size grows with the dimension of the search space can improve a mutation-based EA significantly has been solved. Fitness functions have been presented where a simple GA with population size 2 is efficient while mutation-based EAs need with overwhelming probability exponentially many steps. Efficiency is defined as a success probability of 1-o(1) within a polynomial number of steps or even as a polynomial expected optimization time. The most important types of crossover, namely uniform crossover and one-point crossover, have been investigated. These are the first results to prove for some examples that crossover can be essential even for populations of size 2. Nevertheless, in most cases of application, it is useful to have larger populations and some method to preserve the diversity in the population.

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