Theoretical and empirical runtime analysis of evolutionary algorithms for the PARTITION problem

Bachelor Thesis of

Daniel Lipp

At the Department of Informatics and Mathematics Chair of algorithms for intelligent systems



Advisor: Prof. Dr. Dirk Sudholt

Time Period: 14th May 2023 - 14th August 2023

Statement of Authorship
I hereby declare that this document has been composed by myself and describes my own work, unless otherwise acknowledged in the text.
Passau, August 11, 2023

Abstract

When dealing with NP-hard problems calculating an exact solution will take exponential time for some inputs unless P = NP. One possible way of dealing with the high runtime is abandoning the optimality for a better runtime which is the concept of approximation-algorithms. Randomised Search Heuristics such as Evolutionary Algorithms can be used for this purpose. In this thesis the runtime of multiple Evolutionary Algorithms such as the (1+1) EA, variants of the RLS and the *pmut* mutation operator is analysed theoretically and empirically on PARTITION. Some of the previously known bounds for the (1+1) EA and RLS are improved and expanded to different version of those algorithms. In the end there is an empirical study on the best algorithm variant depending on input type and size revealing the optimal mutation rate is dependent on the input and not always the same for every input of PARTITION.

Deutsche Zusammenfassung

NP-schwere Probleme lassen sich nicht immer in polynomieller Zeit lösen, es sei denn P=NP gilt. Eine Möglichkeit die exponentielle Laufzeit zu vermeiden ist es statt der optimalen Lösung auch schlechtere Lösungen zu akzeptieren, was dem Prinzip der Approximationsalgorithmen entspricht. Randomised Search Heuristics wie Evolutionäre Algorithmen können für diesen Zweck verwendet werden. In dieser Bachelorarbeit wird die Laufzeit mehrerer Evolutionärer Algorithmen wie dem (1+1) EA, Varianten des RLS sowie des pmut Operators theoretisch und empirisch analysiert für PARTITION. Bereits bestehende Ergebnisse werden verschärft für den (1+1) EA und den RLS und erweitert für Varianten der beiden Algorithmen. Am Ende der Arbeit ist eine empirische Analyse, die die verschiedenen Algorithmen auf verschiedenen Eingabetypen und Eingabegrößen vergleicht. Ein der vielen Erkenntnisse ist, dass die optimale Mutationsrate abhängig von der Art der Eingabe ist und nicht global für PARTITION gilt.

Contents

T	Intr	oducti	on	1
2	Pre	limina	ries	3
	2.1	Notati	on	3
	2.2	Backgr	round	3
		2.2.1	Known algorithms for partition	3
		2.2.2	Evolutionary Algorithm	4
		2.2.3	Literature on the RLS and $(1+1)$ EA for PARTITION	5
		2.2.4	Problems for Evolutionary Algorithms	6
	2.3	Higher	mutation rates and heavy tailed mutations	6
3	Rur	$\mathbf{ntime}\ \mathit{A}$	Analysis of different EAs on PARTITION	9
	3.1	Impro	ving bounds on the RLS and the (1+1) EA	9
	3.2		ne Analysis of higher mutation rates	14
	3.3	Binom	ial distributed input	16
4	Ext	erimei	ntal Results	19
	4.1			19
		4.1.1	The Algorithms	19
		4.1.2	Random number generation	20
	4.2	Do bir	nomial inputs have perfect partitions?	21
	4.3		ial distributed values	$\frac{-}{25}$
		4.3.1	RLS Comparison	27
		4.3.2	(1+1) EA Comparison	27
		4.3.3	pmut Comparison	28
		4.3.4	Comparison of the best variants	28
	4.4	Geome	etric distributed values	29
		4.4.1	RLS Comparison	30
		4.4.2	(1+1) EA Comparison	31
		4.4.3	pmut Comparison	31
		4.4.4	Comparison of the best variants	31
	4.5	Unifor	m distributed inputs	32
		4.5.1	RLS Comparison	33
		4.5.2	(1+1) EA Comparison	33
		4.5.3	pmut Comparison	34
		4.5.4	Comparison of the best variants	34
	4.6	powerl	law distributed inputs	35
		4.6.1	RLS Comparison	36
		4.6.2	(1+1) EA Comparison	36
		4.6.3	pmut Comparison	37
		4.6.4	Comparison of the best variants	37

4.7	Eqiuva	lent of linear functions for PARTITION	38
	4.7.1	RLS Comparison	39
	4.7.2	(1+1) EA Comparison	39
	4.7.3	pmut Comparison	40
	4.7.4	Comparison of the best variants	40
4.8	Carste	n Witt's worst case input	42
	4.8.1	RLS Comparison	43
	4.8.2	(1+1) EA Comparison	43
	4.8.3	pmut Comparison	44
	4.8.4	Comparison of the best variants	44
4.9	Multip	le distributions combined	45
	4.9.1	RLS Comparison	45
	4.9.2	(1+1) EA Comparison	46
	4.9.3	pmut Comparison	46
	4.9.4	Comparison of the best variants	46
4.10	Conclu	sion of empirical results	47
Bibliog	raphy		49

1. Introduction

The question of P = NP is still unanswered to this day and solving NP-hard problems for every instance still requires exponential time. To avoid the exponential running time on NP-hard optimisation problems one might use approximation algorithms. Those algorithms do not always return the best possible solution but only a solution with a guaranteed solution quality. For a minimisation problem a (1+0.5)-approximation algorithm will always return a solution that has at most 1.5 times the optimal value. An example of an NP-hard optimisation problem is PARTITION. An instance of this problem is a multiset of n positive numbers $\{w_1,\ldots,w_n\}$ which has to be divided in two subsets with sums that are as close as possible. So a solution of PARTITION is a subset of $I \subset \{1, \ldots, n\}$ which splits the multiset into two subsets. The quality of the solution then is $\max\{\sum_{i\in I} w_i, \sum_{i\notin I} w_i\}$. PARTITION is one of the easiest NP-hard problems and has even been dubbed the easiest NP-hard problem [Hay02]. There are multiple algorithms specifically designed for PARTITION. Some of them return approximations and others always return the best solution. The exact algorithms have a runtime exponential in the input size due to the NP-hardness. Problem specific approximation algorithms are mostly deterministic such as the greedy method, the KK-algorithm or the FPTAS for the subsetsum problem which can be used for partition as well. Another class of non-deterministic algorithms are the so-called Evolutionary Algorithms which mimic the behaviour of evolution. Those algorithms start with a random population of solutions which are then changed with random steps. If the solution is good enough it survives and replaces one of the worst individuals in the population. The EA continues to generate new offspring in an endless loop. In practice the algorithm is given stopping conditions such as reaching a number of iterations or a specific solution quality. This is the principle of an anytime algorithm which can be terminated at anytime and output a valid solution. The longer the waiting time the better the solution might get. The main usage of EA lies in problems without a problem specific algorithms as these mostly outperform the EAs. To better understand their behaviour analysing them on well researched problems might still be beneficial to learn more about this class of algorithms. This thesis researches the runtime of basic EAs such as (1+1) EA and variations of the RLS. The first part is a theoretical analysis with a main focus of lowering bounds that were previously shown. Additionally there are new results for other algorithm variants and also a lemma on different type of inputs. The remainder of this thesis is an empirical analysis of multiple base algorithms with different parameter setting on different kinds of inputs. Here not only the (1+1) EA with different mutation rate and variants of the RLS with different parameter values are researched but also a heavy tailed mutation operator. Apart from typical distributions such as the uniform, geometric and binomial

distribution there are also results for problem specific instances such as an input where one values is as large as all other values combined. In the end the empirical results are condensed into a personal suggestion which algorithm should best be chosen to solve the problem, depending on the input but also in general.

2. Preliminaries

2.1 Notation

A short list of vocabulary used throughout the paper.

EA Evolutionary Algorithm

RSH Randomised Search Heuristic referring to all analysed Evolutionary algorithms

- n The input length of the problem
- w_i The *i*-th object of the input. If not mentioned otherwise the weights are sorted in non-increasing order so: $w_1 \ge w_2 \ge ... \ge w_{n-1} \ge w_n$
- W The sum of all objects: $W = \sum_{i=1}^{n} w_i$
- **bin** When solving Partition a set of numbers is divided into two distinct subsets and in this paper both subsets are referred to as bins
- b_F The fuller bin (the bin with more total weight)
- b_E The emptier bin (the bin with less total weight)
- opt The optimal solution for a given partition instance.
- x A vector $x \in \{0,1\}^n$ describing a solution
- f(x) The fitness function for PARTITION. This means a solution x has a solution quality of $f(x) = \max\{\sum_{i=1}^{n} w_i \cdot x_i, \sum_{i=1}^{n} w_i \cdot (1-x_i)\} = b_F$

2.2 Background

2.2.1 Known algorithms for partition

Multiple methods for generating a solution of PARTITION already exist. Solving the problem with a greedy approach in runtime $\mathcal{O}(n)$ results in an approximation ratio of 3/2 if the elements are not sorted or a ratio of 7/6 if the numbers are sorted [Gra66]. Greedy in this case means putting each element in the currently emptier set while looking at each value exactly once. Another approximation algorithm is the KK-algorithm or also called Largest Differencing Method. With expected time $\mathcal{O}(n \log n)$ it has the same runtime as greedy with sorting and also the same worst case approximation of 7/6. For inputs chosen

uniform random from [0,1] the KK has an expected ratio of $1 + \frac{1}{n^{\Theta(\log n)}}$ in comparison to the greedy algorithm which only reaches an approximation ratio of $1 + \mathcal{O}(\frac{1}{n})$. Instead of putting each element in the currently emptier set the currently largest two values are combined to one value by either subtracting or adding them depending on which results in a better solution. Adding them corresponds to putting the elements in the same set whereas subtracting them means putting them in different sets. There is even a fully polynomial time approximation algorithm (FPTAS) for the subsetsum problem [KMPS03] which can be used for PARTITION by setting the required sum to |W/2|. FPTAS return a solution of at most $(1+\epsilon)$ the optimum in a time that is polynomial both in n and in $\frac{1}{\epsilon}$. There are lots of other approximation-algorithms but also some algorithms that always return the best solution. The Pseudopolynomial time number partitioning algorithm which uses dynamic programming always returns an optimal solution but needs time and space $\mathcal{O}(n^{\frac{m}{2}})$ where m is the largest number in the input [Kor09]. The runtime is only pseudopolynomial because to encode m in the input only $\log_2(m)$ bits are required which causes $m = 2^{\log_2(m)}$ to be exponential in the input size. The Complete Greedy Algorithm (CGA) [Kor98] traverses a binary tree depth first and searches the complete 2^n search space in a greedy way. It functions the same way as the simple greedy algorithm but instead of only looking at only the greedy option at each height it also evaluates the non-greedy option after evaluating the complete subtree of the greedy option. The algorithm continues the depth first search until it either finds a perfect partition or has traversed the whole tree. In the second case it will return the best value found on the way. While the space complexity in only $\mathcal{O}(n)$ the runtime is $\mathcal{O}(2^n)$. Another exact algorithm is the Complete Karmarkar-Karp (CKK) [Kor98]. This algorithm works similar to the complete greedy approach by traversing the binary tree of all solutions. Instead of greedily selecting the next edge here the algorithm behaves like the KK-algorithm described above. It performs better than the CGA for the same reasons as before but also has the same worst case running time as the GCA.

2.2.2 Evolutionary Algorithm

Evolutionary Algorithms mimic the process of evolution and normally behave mostly the same. A run typically looks like this:

- 1. Generate initial population at random
- 2. If stopping condition are met return the currently best solution
- 3. Generate offspring population (e.g. by mutation)
- 4. Evaluate fitness of the offspring
- 5. Select fittest individuals and update population
- 6. Go back to step 2.

For PARTITION a solution $x \in \{0,1\}^n$ separates all numbers into two different sets with $x_i = 0$ meaning w_i is in set 0 whereas $x_i = 1$ meaning w_i is in set 1. So every possible value of x describes a feasible solution but not necessarily a good one. To evaluate the quality of a solution the EA is given a fitness function. The fitness function in this case is $f(x) = \max\{\sum_{i=1}^n w_i \cdot x_i, \sum_{i=1}^n w_i \cdot (1-x_i)\} = b_F$. The goal of the algorithm is to return a solution with minimal fitness. A mutation step in the PARTITION problem will change an algorithm-dependent number of bits from 1 to 0 or vice versa. In this case flipping a bit means putting the element in the other set. A simple implementation of an EA is the so called (1+1) EA (Algorithm 2.1). The first 1 in the brackets refers to size of the population and the second to the amount of mutants created in each iteration of the loop. So it always has only one solution and generates just one new solution in each step. The

Algorithm 2.1: (1+1) EA

```
1 choose x uniform from \{0,1\}^n
2 while x not optimal do
3 x' \leftarrow x
4 flip every bit of x' with probability 1/n
5 if f(x') \leq f(x) then
6 x \leftarrow x'
```

Algorithm 2.2: RLS

```
1 choose x uniform from \{0,1\}^n
2 while x not optimal do
3 x' \leftarrow x
4 flip one uniform random bit of x'
5 if f(x') \leq f(x) then
6 x \leftarrow x'
```

mutation of the current individual is performed by flipping each bit independently with probability 1/n. The amount of flipped bits is binomial distributed with an expected value of $n \cdot \frac{1}{n} = 1$. Another simple EA is the Randomised Local Search (RLS see Algorithm 2.2). Instead of flipping each bit with probability 1/n here one bit is flipped at uniform random. Apart from that the algorithms are the same.

2.2.3 Literature on the RLS and (1+1) EA for PARTITION

Carsten Witt proved that the RLS and the (1+1) EA find a $(4/3+\epsilon)$ approximation in expected time $\mathcal{O}(n)$ and a (4/3)-approximation in expected time $\mathcal{O}(n^2)$ [Wit05]. He then introduced an almost worst case input to prove the bound for the approximation ratio is at least almost tight. The input is defined as followed for any $0 < \epsilon < 1/3$ and even n: The input contains two numbers of value $1/3 - \epsilon/4$ and n-2 elements of value $(1/3 + \epsilon/4)$ $\epsilon/2)/(n-2)$. The total volume is normalised to 1. When the two large values are in the same bin, the RSHs are tricked into a local optimum, where only w_1 and w_2 are in the first bin and the remaining elements in the other bin. This results in an almost worst case. To leave this local optimum $\Omega(n)$ bits must be moved in a step separating the two large values. Such a step will never happen for the RLS and only in expected exponential time for the (1+1) EA. This worst case happens with probability $\Omega(1)$. He also proved both RSHs return a $(1+\epsilon)$ -approximation for $\epsilon \geq 4n$ in expected time $\lceil en \ln(4/\epsilon) \rceil$ for the (1+1) EA and $\lceil en \ln(4/\epsilon) \rceil$ for the RLS with probability at least $2^{-(e \log e + e)\lceil 2/\epsilon \rceil \ln(4/\epsilon) - \lceil 2/\epsilon \rceil}$ for the (1+1) EA and at least $2^{-(\log e+1)\lceil 2/\epsilon\rceil \ln(4/\epsilon)-\lceil 2/\epsilon\rceil}$ for the RLS. Afterwards he proved both RHSs reach a solution where the difference between the two bins is at most 1 for uniform distributed inputs on [0,1] after expected time $\mathcal{O}(n^2)$ for the (1+1) EA and $\mathcal{O}(n \log n)$ for the RLS. The difference between the two bis is even bounded by $\mathcal{O}(\log n/n)$ after $\mathcal{O}(n^{c+4}\log n)$ steps with probability at least $1-\mathcal{O}(1-1/n^c)$. This leads to an expected difference of $\mathcal{O}(\log n/n)$ after $\mathcal{O}(n^{c+4}\log n)$ steps. He also analysed exponential distributed inputs with parameter 1. With probability $1 - \mathcal{O}(1/n^c)$ the difference on those inputs is bounded by $\mathcal{O}(\log n)$ after $\mathcal{O}(n^2 \log n)$ steps and even by $\mathcal{O}(\log n/n)$ after $\mathcal{O}(n^{c+4} \log^2 n)$ steps. Additionally he described a polynomial time randomised approximation scheme (PRAS) for the RLS and the (1+1) EA for values of $\epsilon = \Omega(\log \log n / \log n)$. For MAKESPAN-SCHEDULING a list of processing times has to be distributed on a set

of machines while minimising the total time of the fullest machine. With 2 machines this problem is exactly the same as PARTITION. So in a sense MAKESPAN-SCHEDULING is a more general version of PARTITION. This lead to Christian Gunia generalising some results previously shown by C. Witt to MAKESPAN-SCHEDULING on k machines [Gun05]. Solutions for MAKESPAN-SCHEDULING are $x \in \{0, \ldots, k-1\}^n$ and therefore during a mutation x_i is set to a uniform random value from $\{0, \ldots, k-1\} \setminus \{x_i\}$ instead of $1-x_i$. The adapted RSHs reach an approximation ratio of (2k/k+1) in expected time $\mathcal{O}(Wn^{2k-2}/w_n)$. On an instance where every weight is the same the expected optimisation time is bounded by $\mathcal{O}(n \log n)$. He also adapted the almost worse case to the more general problem and proved the RLS does not find a solution better than $(2k/(k+1)-\epsilon)$ in finite time for any $\epsilon > 0$ with constant probability. The second statement for PARTITION on the uniform distributed inputs on [0,1] has an equivalent lemma for MAKESPAN-SCHEDULING on k machines as well.

Another way of dealing with NP-hard problems is identifying a parameter k which defines how hard the problem is to solve. One possible parametrisation of PARTITION is solving whether there is a solution of $f(x) \leq k$. A fixed-parameter tractable problem is a problem that can be solved in time polynomial in the size of the input and g(k) where g is any arbitrary function. Partition falls into this category as it can be decided in time at most $\mathcal{O}(4^k)$ [Fer05]. Andrew M. Sutton and Frank Neumann made a parametrised analysis of PARTITION[SN12]. They parametrised the problem in multiple ways. One of their parametrisation was: given an integer k, is there a solution of at most W/2 + W/k? They showed that a multistart of the (1+1) EA or RLS using runs of length $\lceil en \ln(2k) \rceil$ is a Monte Carlo-fpt algorithm for this parametrised version of PARTITION. They also analysed a parametrisation of the size of the critical path and also the discrepancy (the difference between the two bins).

2.2.4 Problems for Evolutionary Algorithms

There are two well analysed problems for Evolutionary Algorithms which are relevant for this thesis. One of those is OneMax. For OneMax the fitness function $f(x) = \sum_{i=1}^{n} x_i$ has to be maximised. In the optimal solution every bit is set to one. This problem is rather easy as it has only one global optimum and no local optimum. So every step the algorithms makes decreases the Hamming distance to the optimum if the fitness increases. A more general version of this problem are linear functions where every bit is given a weight. Here the fitness function is $f(x) = \sum_{i=1}^{n} w_i \cdot x_i$ which either has to be maximised or minimised. The weights w_i can be any real value. In contrast to OneMax increasing the fitness can increase the Hamming distance to the global optimum if multiple small values switch places with a big value. The runtime for both the RLS and the (1+1) EA is $\Theta(n \log n)$ for both problems and the optimal mutation rate for the (1+1) EA is 1/n which was proven multiple times ([Wit13], [DJL23]). The optimality of 1/n does not hold for every problem which leads to the next section.

2.3 Higher mutation rates and heavy tailed mutations

The RLS and the standard (1+1) EA flip one bit in expectation which is optimal for some problems as seen in the last section. This is not the case for every fitness function. Jump_k has an optimal mutation rate of k/n and a small constant factor deviation from k/n results in an increase of the runtime exponential in $\Omega(k)$ [DLMN17]. The same might hold for PARTITION, because the previously discussed literature only analyses mutation rates with a 1-bit flip in expectation.

One way of creating algorithms with higher mutation rates is adjusting the currently

Algorithm 2.3: (1+1) EA WITH STATIC MUTATION RATE

```
1 choose x uniform from \{0,1\}^n

2 while x not optimal do

3 x' \leftarrow x

4 flip every bit of x' with probability c/n

5 if f(x') \leq f(x) then

6 x \leftarrow x'
```

Algorithm 2.4: RLS_k^B

```
1 choose x uniform from \{0,1\}^n

2 while x not optimal do

3 x' \leftarrow uniform random neighbour of x with Hamming distance \leq k

4 if f(x') \leq f(x) then

5 x \leftarrow x'
```

existing algorithms. For the (1+1) EA this can be done easily. By changing the mutation rate 1/n to c/n for any constant c the algorithm now flips $n \cdot c/n = c$ bits in expectation.

For the RLS it is not that simple, as the RLS chooses a random bit and flips it. Instead of flipping c bits in every step there should be the possibility to flip different amounts of bits in every step. The standard RLS chooses a random neighbour with Hamming distance one. So the variant of the RLS could simply choose neighbours that have a Hamming distance larger than one. The selection should still be uniform random to keep the idea of the RLS intact. One possible way is to choose a random neighbour with Hamming distance $\leq k$. This algorithm will be called RLS_k^B from now on, because it chooses a random neighbour within the Hamming ball with radius k. The amount of neighbours with Hamming distance y is given by $\binom{n}{y}$. For k=4, this results in n neighbours with Hamming distance 1, n(n-1)/2 neighbours with Hamming distance 2, n(n-1)(n-2)/6 for 3 and n(n-1)(n-2)(n-3)/24 for 4. The probability to choose a random neighbour with Hamming distance $y \leq k$ for $k = \mathcal{O}(1)$ is given by

$$P(\text{RLS}_k^B \text{ flips } y \text{ bits}) = \frac{\binom{n}{y}}{\sum_{i=1}^k \binom{n}{i}} = \frac{\Theta(n^y)}{\sum_{i=1}^k \Theta(n^i)} = \frac{\Theta(n^y)}{\Theta(n^k)} = \Theta(n^{y-k}) = \Theta(\frac{1}{n^{k-y}})$$

This variant of the RLS is likely to choose a neighbour with Hamming distance k as the number of neighbours with hamming distance k rises with k for $k \leq n/2$. The probability of flipping only one bit is $\mathcal{O}(\frac{1}{n^{k-1}})$. For some inputs flipping only one bit might be more optimal which is rather unlikely for this variant of the RLS.

An alternative way of changing the RLS is to first choose $y \in \{1, ..., k\}$ uniform random and then choose a neighbour with Hamming distance y uniform random. Here the probability of flipping $y \le k$ bits is given by 1/k, so the algorithm is much more likely to choose to flip only one bit. This variant of the RLS will be referred to as RLS_k^S because it first choses the Hamming sphere and afterwards the neighbour within the selected Hamming sphere.

Both variants of the RLS change at most k bits in each step and therefore only a constant amount of bits. For the (1+1) EA the algorithm will also flip mostly $\mathcal{O}(c)$ bits which is also constant. So neither of the new variants is likely to change up to $\Theta(n)$ bits. Quinzan et al. therefore introduced another mutation operator in [FGQW18] called $pmut_{\beta}$. This operator chooses k from a powerlaw distribution D_n^{β} with exponent β and maximum value

Algorithm 2.5: RLS_k^S

```
1 choose x uniform from \{0,1\}^n
2 while x not optimal do
3 y \leftarrow uniform random value \in \{1,\ldots,k\}
4 x' \leftarrow uniform random neighbour of x with Hamming Distance y
5 if f(x') \leq f(x) then
6 x \leftarrow x'
```

n and then flips k uniform random bits. This algorithm will mostly flip a small number of bits but occasionally up to n bits. Mutation operators like this are called heavy tailed mutations because their tail is not bounded exponentially.

3. Runtime Analysis of different EAs on PARTITION

This chapter focuses on the theoretical analysis of the runtime of Evolutionary Algorithms for PARTITION. The first section is focused on improving previously shown bounds for the (1+1) EA and the RLS. Afterwards there is also an analysis of Evolutionary Algorithms that flip more than one bit in expectation. The last section analyses inputs that follow a binomial distribution.

3.1 Improving bounds on the RLS and the (1+1) EA

As discussed in the Background section C. Witt already showed multiple results for the RLS and the standard (1+1) EA. One of the results is the approximation ratio of at most 4/3 after expected time at most $\mathcal{O}(n^2)$. In this section this bound will be lowered to $\mathcal{O}(n \log n)$ for both algorithms. The special input with $w_1 \geq W/2$ is also analysed for the (1+1) EA with mutation rate c/n. For the analysis of mutation rates c/n the work of C. Witt only has to be slightly adjusted. The first Lemma is a modification of a Lemma in [Wit05].

Lemma 3.1. Let $w_1 \geq W/2$, then for any $\gamma > 1$ and $0 < \delta < 1$, the (1+1) EA with mutation rate c/n with constant $0 < c < \sqrt{n}$ (RLS_k^S) reaches an f-value at most $w_1 + \delta(W - w_1)$ in at most $\lceil \frac{e^c}{c \cdot (1-o(1))} n \ln(\gamma/\delta) \rceil$ $(\lceil kn \ln(\gamma/\delta) \rceil)$ steps with probability at least $1-\gamma^{-1}$. Moreover, the expected number of steps is at most $2\lceil \frac{e^c}{c \cdot (1-o(1))} n \ln(2/\delta) \rceil$ $(2\lceil kn \ln(2/\delta) \rceil)$.

Proof. This Lemma is very similar to C. Witt's Lemma 2 from section '2. Definitions and Proof Methods' in [Wit05]. The proof is mostly the same. He first defines a potential function p(x) = f(x) - f(opt). While p(x) > 0 all steps moving only a small object to the emptier bin are accepted. The expected p-decrease is at least $p_0 \cdot q$ where q is a lower bound on the probability of the algorithm to flip one specific bit. This leads to a next p value of $(1-q)p_0$. Since all steps of both algorithms are independent this argumentation remains valid even if the p value is only an expected value. With q = 1/yn for a constant y > 0 the expected p value after p value after p value is at most

$$p_t \le p_0 (1 - 1/yn)^t = p_0 (1 - 1/yn)^{yn \ln(\gamma/\delta)} \le p_0 \cdot e^{-\frac{1}{y} \cdot yn \ln(\gamma/\delta)} = p_0 (\gamma/\delta)^{-1} = p_0 (\delta/\gamma)$$

Applying Markov's inequality to the non-negative p value implies $p_t \leq p_0 \delta$ with probability $1 - 1/\gamma$. Repeating independent phases of length $\lceil yn \ln(2/\delta) \rceil$ the expected number of

phases is at most 2. Up until here the proof is the same.

Instead of choosing W/2 as the general upper bound for p_0 as in the original lemma here the lower value $W-w_1 \leq W/2$ is chosen because it is more tight for the special case $w_1 \geq W/2$ with $f(opt) = w_1$. The probability of the RLS_k^S to flip one specific bit is $\frac{1}{k} \cdot \frac{1}{n}$ and for (1+1) EA with mutation rate c/n at least

$$\frac{c}{n}(1-\frac{c}{n})^{n-1} \ge \frac{c}{n}(1-\frac{c}{n})^n \ge \frac{c}{n}e^{-c}(1-\frac{c^2}{n}) = \frac{c}{e^cn}(1-o(1))$$

The inequality $(1+x/n)^n \ge e^x(1-x^2/n)$ requires $n \ge 1, |c| \le n$ which both hold. Setting $y = \frac{e^c n}{c \cdot (1-o(1))}$ for the (1+1) EA and y = k for the RLS_k^S concludes the result.

The next Lemma also analyses inputs with $w_1 > W/2$. Its goal is to bound the probability of the (1+1) EA with mutation rate c/n and the $RLS_{k\geq 2}^S$ of flipping the first bit after a certain solution quality has been reached. Inputs with $w_1 > W/2$ are similar to linear functions which strongly suggest a runtime of $\mathcal{O}(n \log n)$ without flips of w_1 . If such a step is too unlikely the algorithms might find an optimal solution before w_1 is flipped and the Hamming distance to the optimum might increase drastically.

Lemma 3.2. For instances with $w_1 > W/2$ the probability of flipping w_1 when $b_E = c \cdot \frac{W-w_1}{2}$ for 1 < c < 2 holds, is at most $\frac{2y(y-1)^2}{n(n-1)(c-1)}$ in a step where any algorithm flips $2 \le y \le n/2$ bits.

Proof. For a successful flip of w_1 after $b_E \ge \frac{W-w_1}{2}$ holds a total volume of at least $z \ge 2 \cdot (b_E - \frac{W-w_1}{2})$ must be shifted from b_E to b_F . Otherwise the step is rejected because

$$b_F' = b_E + w_1 - z > b_E + w_1 - 2 \cdot (b_E - \frac{W - w_1}{2}) = b_E + w_1 - 2b_E + W - w_1 = W - b_E = b_F$$

which results in an increase of the fitness $(b_F = f(x), b_F' = f(x'))$. Let I be the set of indices of all elements moved from b_E to b_F and $w_{\max} = \max\{w_i | i \in I\}$. $|I| \leq y - 1$ holds because at least w_1 is moved from b_F to b_E . The sum of all elements is at most $w_{\max} \cdot |I| \leq (y-1)w_{\max}$. If $w_{\max} < 2 \cdot (b_E - \frac{W-w_1}{2})/(y-1)$ then $(y-1)w_{\max} < 2 \cdot (b_E - \frac{W-w_1}{2})$ and the step is rejected. Thus at least one of the objects moved from b_E to b_F must have a volume of at least $2 \cdot (b_E - \frac{W-w_1}{2})/(y-1)$. At most $d \leq \frac{b_E}{w_{\max}}$ of these objects can be in b_E if they made up the complete volume of b_E . Simplifying this inequality leads to at most

$$d \leq \frac{b_E}{w_{\max}} \leq \frac{W-w_1}{w_{\max}} \leq \frac{W-w_1}{2(c\frac{W-w_1}{2} - \frac{W-w_1}{2})/(y-1)} = \frac{(W-w_1)(y-1)}{(W-w_1)(c-1)} = \frac{(y-1)}{(c-1)}$$

objects having at least a volume of w_{max} . For a successful flip w_1 and at least one of these d objects must switch bins and the probability for such a step flipping y bits is therefore at most

 $\mathbb{P}(y \text{ bits are flipped}) \cdot \mathbb{P}(\text{the correct } y \text{ bits are flipped})$

$$\leq 1 \cdot \frac{\binom{1}{1} \cdot \binom{\lceil d \rceil}{1} \cdot \binom{n-2}{y-2}}{\binom{n}{y}} = \frac{\lceil d \rceil \frac{(n-2)!}{(n-2-y+2)! \cdot (y-2)!}}{\frac{n!}{(n-y)! \cdot y!}} = \frac{\lceil d \rceil \cdot (n-2)! \cdot (n-y)! \cdot y!}{n! \cdot (n-y)! \cdot (y-2)!} = \frac{\lceil d \rceil y (y-1)}{n(n-1)}$$

$$\leq \frac{y(y-1)}{n(n-1)} \cdot (\frac{y-1}{c-1}+1) = \frac{y(y-1)}{n(n-1)} \cdot (\frac{y-1+c-1}{c-1}) \leq \frac{2y(y-1)^2}{n(n-1)(c-1)}$$

Theorem 3.3 is the last analysis on inputs with $w_1 \geq \frac{W}{2}$. It uses the results of the two previous lemmas and gives an asymptotic bound on the runtime. Instead of giving a runtime for a 4/3-approximation this lemma gives the expected runtime for reaching one of the two optimal solutions. For a non-optimal solution moving one element from the fuller to the emptier bin will always result in an improvement. There are also no local optima which neither of the algorithms is unlikely to leave. So this input is rather easy to solve for the RLS and (1+1) EA and comparable to a linear function or even OneMax if $w_2 = \cdots = w_n$.

Theorem 3.3. If $w_1 \ge \frac{W}{2}$ then the RLS and the (1+1) EA with mutation rate k/n with constant $0 < k < \sqrt{n}$ reach the optimal solution in expected time $\Theta(n \log n)$

Proof. The optimal solution is putting w_1 in one bin and all other elements in the other bin. So the problem is almost identical to OneMax/ZeroMax. A single bit flip of the first bit can only happen, if the emptier bin has a weight of at most $\frac{W-w_1}{2}$. After this flip the weight of the emptier bin is at least $\frac{W-w_1}{2}$ and therefore another single bit flip of w_1 can only happen before another bit is flipped. The run of the RLS can be divided into two phases:

Phase 1: The RLS reaches a search point with $b_E > \frac{W - w_1}{2}$.

Phase 2: The RLS reaches an optimal solution $\Rightarrow w_1$ is in one bin and all other elements are in the other bin.

The expected length of the first phase is at most 2n because the probability of flipping the first bit is $\frac{1}{n}$ and the expected time for such a step then is at most n. After such a step $b_E \geq \frac{W-w_1}{2}$ holds. If the solution is already optimal $b_E = W - w_1 > \frac{W-w_1}{2}$, otherwise there is at least one bit that can be flipped. This bit will be flipped in expected time at most n for the same reason as for w_1 . This leads to a total length of first phase of at most 2n. In the second phase the RLS can no longer flip w_1 as it does not result in an improvement ever again. Therefore the RLS behaves exactly as on OneMax/ZeroMax depending on the value of the first bit and reaches an optimal solution in $\Theta(n \log n)$ resulting in a total runtime of $\Theta(n \log n)$ (Theorem 3 in [Wit14]).

As long as w_1 does not flip the (1+1) EA has to minimize a linear function of n-1 bits which takes $(1+o(1))\frac{e^k}{k}n\ln n$ time (Corollary 4.2 in [Wit13]). The only steps that could hinder the algorithm from optimising the linear function in $\Theta(n\log n)$ would be a flip of the first bit. Such steps invert the optimal solution which could decrease the progress of minimising the linear function. If such a step has an expected time of $\omega(n\log n)$ the linear function is likely to be optimised in expectation before such a step happens.

The probability of the (1+1) EA to flip more than $k + \sqrt{6k \ln(n)} = k + k\sqrt{6\ln(n)/k}$ is limited by Chernoff bounds:

$$\begin{split} & \mathbb{P}((1+1) \text{ EA flips more than } k + k \sqrt{6 \ln(n)/k} \text{ bits}) \\ & \leq \mathbb{P}(X \geq (1 + \sqrt{6 \ln(n)/k}) \cdot k) \leq e^{-k \cdot \sqrt{6 \ln(n)/k}^2/3} = e^{-k \cdot \frac{6 \ln n}{3k}} = n^{-2} \end{split}$$

So the expected time for such a step is at least $n^2 = \omega(n \ln(n))$. Now let's look at steps that flip at most $k + \sqrt{6k \ln(n)}$ bits in a single step. Such a step only successfully flips w_1 if both w_1 is flipped and enough total volume is shifted from b_E to b_F . Due to Lemma 3.1 with $\delta = \frac{1}{n}$ (for n > 1) the solution is at most $w_1 + \delta(W - w_1)$ after expected time

$$2\lceil \frac{e^k}{k(1-o(1))} n \ln(2/\delta) \rceil = 2\lceil \frac{e^k}{k(1-o(1))} n \ln(2/\frac{1}{n}) \rceil = 2\lceil \frac{e^k}{k(1-o(1))} n \ln(2n) \rceil$$
$$= 2\lceil \frac{e^k}{k(1-o(1))} n (\ln(n) + \ln(2)) \rceil \le \frac{2e^k}{k(1-o(1))} n (\ln(n) + 4)$$

The value of b_E is then at least $W - (w_1 + \delta(W - w_1)) = (1 - \delta)(W - w_1) = (1 - \frac{1}{n})(W - w_1)$.

Lemma 3.2 states that the probability of a step flipping with w_1 together with y-1 other bits is at most $\frac{2y(y-1)^2}{n(n-1)(c-1)}$. Applying the bound $y \le k + \sqrt{6k \ln(n)}$ and the value $c = 2(1 - \frac{1}{n})$ this simplifies to

$$\frac{2y(y-1)^2}{n(n-1)(c-1)} \leq \frac{2(k+\sqrt{6k\ln(n)})^3}{n(n-1)(1-\frac{2}{n})} = \frac{2(k+\sqrt{6k\ln(n)})^3}{n(n-1)\frac{n-2}{n}} = \frac{2(k+\sqrt{6k\ln(n)})^3}{(n-1)(n-2)}$$

The probability of one of these steps to happen for any value of y is given by

 $\sum_{y=2}^{k+\sqrt{6k\ln(n)}} \mathbb{P}(y \text{ bits are flipped}) \cdot \mathbb{P}(\text{the correct } y \text{ bits are flipped} | y \text{ bits are flipped})$

$$\leq (k + \sqrt{6k \ln(n)}) \cdot \frac{2(k + \sqrt{6k \ln(n)})^3}{(n-2)(n-1)} = \frac{2(k + \sqrt{6k \ln(n)})^4}{(n-2)(n-1)}$$

$$= \frac{2(o(n^{1/8}))^4}{(n-2)(n-1)} = \frac{o(n^{0.5})}{\mathcal{O}(n^2)} = \mathcal{O}(n^{-1.5})$$

The expected time for any step successfully flipping w_1 is then $\Omega(n^{1.5}) = \omega(n \ln n)$. Let T be the time until the linear function is optimised and p the probability of successfully flipping w_1 . Then the probability that w_1 flips after expected time $\frac{2e^k}{k(1-o(1))}n\ln(n) + 4$ before the linear function is optimised is at most

$$p \cdot E(T) \le \frac{1}{\mathcal{O}(n^{1.5})} \cdot (1 + o(1)) \frac{e^k}{k} n \ln n = \frac{(1 + o(1)) \frac{e^k}{k} \ln n}{\mathcal{O}(n^{0.5})} = \frac{o(n^{0.1})}{\mathcal{O}(n^{0.5})} = o(\frac{1}{n^{0.4}}) = o(1)$$

The probability of w_1 not being flipped after expected time $\frac{2e^k}{k(1-o(1))}n\ln(n)+4$ is $1-o(\frac{1}{n^{0.4}})=1-o(1)$. If such a step happens the fitness does not decrease and the bound on the probability of flipping w_1 still holds. The algorithm will still find the solution in expected time at most $(1+o(1))\frac{e^k}{k}n\ln n$. Since even after a flip all condition are still true the expected time of optimising the linear function after expected time $\frac{2e^k}{k(1-o(1))}n\ln(n)+4$ is given by $\frac{1}{1-o(1)}\cdot(1+o(1))\frac{e^k}{k}n\ln n=\Theta(n\log n)$

The total runtime for the (1+1) EA is

$$\frac{2e^k}{k(1-o(1))} n \ln(n) + 4 + \frac{1+o(1)}{1-o(1)} \cdot \frac{e^k}{k} n \ln n = 4 + \frac{3+o(1)}{1-o(1)} \cdot \frac{e^k}{k} n \ln n = \Theta(n \log n)$$

Theorem 3.3 only proved the asymptotic expected runtime of the (1+1) EA with different mutation rates. It does not proof which mutation rate reaches the solution the fastest in expectation, but it suggests the runtime increases for higher mutation rates. Later in the thesis there is an analysis for this lemma in Section 4.7 which also suggests the optimality of 1/n.

The next few Lemmas and Corollaries help to prove the runtime of $\mathcal{O}(n \log n)$ for the (1+1) EA and the RLS on inputs with $w_1 < W/2$ for reaching an approximation ratio of 4/3. They are mostly rather short with a proof of only a few lines. The only Lemma with a longer proof is Lemma 3.9. It shows the runtime bound for a more restricted type of input and is used to simplify the proof of Lemma 3.10

Lemma 3.4. If $b_F \leq \frac{2}{3} \cdot W$ the approximation ratio is at most $\frac{4}{3}$

Proof.
$$\frac{b_F}{opt} \leq \frac{(2/3) \cdot W}{opt} \leq \frac{(2/3) \cdot W}{(1/2) \cdot W} = \frac{4}{3}$$
, since $opt \geq \frac{W}{2}$

Corollary 3.5. If $w_1 \ge \frac{W}{3}$ and w_1 is in the emptier bin, then the approximation ratio is at most $\frac{4}{3}$

Proof. w_1 is in the emptier bin, so $b_F \leq W - w_1 \leq W - \frac{W}{3} = \frac{2W}{3}$ and with Lemma 3.4 the assumption follows.

Lemma 3.6. Any object of weight v can be moved from b_F to b_E if and only if $b_F - b_E \ge v$

Proof. " \Leftarrow ":

 $b_F - b_E \ge v \Leftrightarrow b_F \ge b_E + v$, so after moving an object with weight v from b_F to b_E , the new weight of b_E is at most the weight of b_F before moving the object, thus the RSH accepts the step.

" \Rightarrow ":

 $b_F - b_E < v \Leftrightarrow b_F < b_E + v$, so moving an object of weight v results in $b_F' = b_E + v > b_F$ which results in the step being rejected.

Corollary 3.7. The RLS is stuck in a local optimum if $b_F - b_E \le w_n$ holds and $b_F > opt$.

Proof. A single bit flip of weight v can only happen if $b_F - b_E \ge v$ (Corollary 3.7). If $b_F - b_E < w_n$ there is no weight which satisfies the condition and therefore no single bit flip is possible. If $b_F - b_E = w_n$ then only objects with weight w_n can be flipped, but those do not change the fitness $(b'_F = b_E + w_n = b_F - w_n + w_n = b_F)$. Since the RLS can only move one bit at a time and only if it results in an improvement, the RLS is stuck.

Corollary 3.8. Every object $\leq \frac{W}{3}$ can be moved from b_F to b_E if $b_F \geq \frac{2W}{3}$

Proof. $b_F \ge \frac{2W}{3} \Rightarrow b_E \le W - \frac{2W}{3} \le \frac{W}{3} \Rightarrow b_F - b_E \ge \frac{2W}{3} - \frac{W}{3} = \frac{W}{3}$ and with Lemma 3.6 the assumption follows.

Lemma 3.9. In expected time $\mathcal{O}(n \log n)$ the weight of the fuller bin can be decreased to $\leq \frac{2W}{3}$ by the RLS and the (1+1) EA if every object besides the biggest in the fuller bin is at most $\frac{W}{3}$ and $w_1 \leq \frac{W}{2}$.

Proof. In expected time $\mathcal{O}(n\log n)$ the RLS can move every object $\leq \frac{W}{3}$ to the emptier bin as long as $b_F \geq \frac{2W}{3}$ due to Corollary 3.8 and Theorem 3.3. So in expected time $\mathcal{O}(n\log n)$ the solution can be shifted to w_1 being in one bin and all other objects in the other bin. The RLS will only stop moving the elements if the condition $b_F \geq \frac{2W}{3}$ is no longer satisfied (Corollary 3.8). If $w_1 \geq \frac{W}{3}$ and every object was moved to the bin without w_1 , then $b_F = \max\{W - w_1, w_1\} = W - w_1 \leq \frac{2W}{3}$, because $w_1 \leq \frac{W}{2}$. So either the RLS moves all objects to the emptier bin or stops moving objects because $b_F < \frac{2W}{3}$ both resulting in $b_F \leq \frac{2W}{3}$. If w_1 is not in the fuller bin, then the result follows by Corollary 3.5. Now assume $w_1 < \frac{W}{3}$. In this case the RLS will move one object per step to the emptier bin. Each object has weight $< \frac{W}{3}$ and therefore one step cannot decrease the weight of the fuller bin from $> \frac{2W}{3}$ to $\leq \frac{W}{3}$. If all objects except one where moved to one bin, the other bin would have a weight of at least $W - w_1 > \frac{2W}{3}$. Therefore the RLS will find a solution with $b_F < \frac{2W}{3}$ before moving all elements from the first to the second bin.

The proof for the (1+1) EA is mostly the same. The main difference is the (1+1) EA being able to flip more than one bit in a single step. Such a step could make the emptier bin the fuller bin or increase the number of bits that must be shifted to the emptier bin. But with the results of Theorem 3.3 the proof works exactly the same as for the RLS. The case $w_1 \geq \frac{W}{3}$ does not change only the bin containing w_1 might change. Apart from that there is no difference for the (1+1) EA. The case $w_1 < \frac{W}{3}$ is also rather similar. The (1+1) EA will move elements from the fuller bin to the emptier bin until $b_F < \frac{2W}{3}$ holds. The (1+1) EA can make the emptier bin the fuller bin by moving multiple objects in one step, but this does not hinder it from reaching $b_F < \frac{2W}{3}$. After a step making the previously fuller bin the emptier bin it will continue moving elements until the condition holds.

Lemma 3.10. The RLS and the (1+1) EA reach an approximation ratio of at most $\frac{4}{3}$ in expected time $\mathcal{O}(n \log n)$ if $w_1 < W/2$

Proof. If $w_1 + w_2 > \frac{2W}{3}$ after time $\mathcal{O}(n)$ w_1 and w_2 are separated and will remain separated afterwards (3. Average case analysis, Theorem 1 in [Wit05]). From then on the following holds. If w_1 is in the emptier bin, then the result follows directly by Corollary 3.5. Otherwise all elements in the fuller bin except w_1 have a weight of at most $\frac{1}{3}$ and therefore the result follows by Lemma 3.9 and Lemma 3.4. If $w_1 + w_2 \leq \frac{2W}{3}$ the result follows directly by Lemma 3.9 and Lemma 3.4.

Corollary 3.11. The RLS and the (1+1) EA reach an approximation ratio of at most $\frac{4}{3}$ for every input in expected time $\mathcal{O}(n \log n)$

Proof. This follows directly from Theorem 3.3 and Lemma 3.10. \Box

This concludes the first part of the theoretical analysed aimed at improving previously shown bounds. Not only do the (1+1) EA and the RLS find a 4/3-approximation in expected time $\mathcal{O}(n \log n)$ instead of only $\mathcal{O}(n^2)$ but also do they find the exact solution in expected time $\Theta(n \log n)$ for the special case $w_1 \geq W/2$. The analysis of the special case was also expanded to the (1+1) EA with a mutation rate c/n for a constant c>0.

3.2 Runtime Analysis of higher mutation rates

In this section the analysis is expanded to variants of the RLS. The first analysed input is again the input with $w_1 \ge W/2$.

Corollary 3.12. The expected time until the RLS_k^S and RLS_k^B flips $w_1 \ge W/2$ after they reached a solution with $b_E = c \cdot \frac{W-w_1}{2}$ with 1 < c < 2 is at least $\frac{n(n-1)(c-1)}{2k^2(k-1)^2}$. For constant values of c the expected time is $= \Omega(n^2)$.

Proof. Using Lemma 3.2, the upper bound of $y \leq k$ and the fact that both algorithms flip at most k bits leads to the first part

$$\left(k \cdot \frac{2y(y-1)^2}{n(n-1)(c-1)}\right)^{-1} = \frac{n(n-1)(c-1)}{2ky(y-1)^2} \le \frac{n(n-1)(c-1)}{2k^2(k-1)^2}$$

k is constant and if c is constant too, this leads to expected time $= \Omega(n^2)$ for a flip of the first bit if $b_E = c \cdot \frac{W - w_1}{2}$ with 1 < c < 2 which concludes the second statement.

Lemma 3.13. The RLS_k^S for $2 \le k = O(1)$ reaches the optimal solution on an input with $w_1 \ge W/2$ in expected time $O(n \log n)$

Proof. This proof is similar to proof for the (1+1) EA in Theorem 3.3 and is divided in the same two parts. The first part is proving the RLS_k^S does not flip the first bit in expected time $\mathcal{O}(n\log n)$ after some time T has passed. After time T the algorithm then minimises the linear function in expected time $\mathcal{O}(n\log n)$ before the first bit is flipped and the progress of the linear function might be reset. After expected time $(2\lceil kn\ln(2/0.4)\rceil) = 2\lceil kn\ln(5)\rceil \le 2\lceil 1.61 \cdot kn\rceil \le 4kn$ the RLS_k^S reaches a solution of $f(x) \le w_1 + 0.4(W - w_1)$ (Lemma 3.1). This means $b_E \ge 0.6(W - w_1) = 1.2 \cdot \frac{W - w_1}{2}$. With Lemma 3.12 the expected time of at least $\frac{n(n-1)(c-1)}{k^2(k-1)^2} \frac{n(n-1)}{5k^2(k-1)^2} = \Omega(n^2)$ for a successful flip of w_1 follows. This concludes the first part

The RLS_k^S can be seen as an unbiased unary black box algorithm with a sequence $\mathcal{D}=(p_1,\ldots,p_n)$ where $p_i=1/k$ for $1\leq i\leq k$ and $p_i=0$ otherwise. The mean of this sequence is $\mathcal{X}=\sum_{i=1}^n \mathbb{P}(X=i)i=\sum_{i=1}^k \frac{i}{k}=\frac{1}{k}\cdot\sum_{i=1}^k i=\frac{k+1}{2}$. For any constant value k both the probability $p_1=\frac{1}{k}=\Theta(1)$ and mean $\mathcal{X}=\frac{k+1}{2}=O(1)$ meet the conditions of Theorem 1 in [DJL23] and therefore the RLS_k^S optimises the linear function in expected time $(1+o(1))\frac{1}{p_1}n\ln n=(1+o(1))kn\ln n$. Applying the same arguments as in Theorem 3.3 this results in a probability of not flipping w_1 after expected time 4kn of at least

$$1 - \frac{(1 + o(1))kn\ln n \cdot 5k^2(k-1)^2}{n(n-1)} = 1 - \frac{(1 + o(1))5k^5\ln n}{n-1} = 1 - \frac{o(k^5n^{0.5})}{n-1} = 1 - \frac{1}{o(\sqrt{n})}$$

The expected time of minimising the linear function then is

$$\frac{1}{1 - \frac{1}{o(\sqrt{n})}} \cdot (1 + o(1))kn \ln n = \frac{1}{1 - o(1)} \cdot (1 + o(1))kn \ln n = \Theta(n \log n)$$

The total expected runtime therefore is at most $4kn + \frac{1+o(1)}{1-o(1)} \cdot kn \ln n = \Theta(n \log n)$.

In expectation all RLS_k^S variants have the same asymptotic runtime on inputs with $w_1 \geq W/2$. They also have the same expected asymptotic Runtime as the standard RLS and the (1+1) EA with static mutation rate c/n for constant values of c. Here the same applies as for the (1+1) EA. There proof does not contain the optimality of flipping only 1 bit in expectation, but it strongly suggests it. The main factor deciding the runtime is the runtime on linear functions which is minimal for k = 1. Later in Section 4.7 there is also an experiment for the different values of k for the RLS_k^S . Another algorithms mentioned later is the RLS_k^B which has a much worse runtime as the next lemma suggests.

Lemma 3.14. The expected optimisation time of the RLS_k^B for $2 \le k = O(1)$ on inputs with $w_1 \ge W/2$ is $\Omega(n^k)$ if the solution is not already optimal after the initialisation.

Proof. The solution has only two global optima. The optimum is defined by x_1 because every other x_i must have a value of $1 - x_1$. If the RLS_k^B reaches a search point where the Hamming distance to the optimum is $y \le k$ there are exactly y bits left that must be flipped for an optimal solution. The other solution requires n - y bits to be flipped which needs even more time. The probability of reaching the optimum from a Hamming distance of y is given by

 $\mathbb{P}(RLS_k^B \text{ flips } y \text{ bits}) \cdot \mathbb{P}(RLS_k^B \text{ flips the correct } y \text{ bits } | RLS_k^B \text{ flips } y \text{ bits})$

$$=\Theta(\frac{1}{n^{k-y}})\cdot\frac{\binom{y}{y}}{\binom{n}{y}}=\Theta(\frac{1}{n^{k-y}})\cdot\frac{1}{\frac{n!}{(n-y)!y!}}=\Theta(\frac{1}{n^{k-y}})\cdot\Theta(\frac{y!}{n^y})=\Theta(\frac{y!}{n^k})$$

Because $y \leq k = O(1)$ holds $\Theta(\frac{y!}{n^k}) = \Theta(\frac{1}{n^k})$ also holds. The expected time for such a step to happen is $(\Theta(\frac{1}{n^k}))^{-1} = \Theta(n^k)$. If the algorithm instead successfully flips less than y bits it reaches a search point where the expected optimisation time is still $\Theta(n^k)$. Either way the expected optimisation time is $\Theta(n^k)$ when the algorithm has reached a search point with Hamming distance $1 \leq y \leq k$. For every value of k the RLS_k^B can only flip at most k bits in each step. So it will reach a search point of RLS_k^B if the initial Hamming distance is at least one. This will happen with probability $1 - 1 \cdot (1/2)^{n-1} = 1 - \frac{1}{2^{n-1}}$.

For smaller values of n the actual runtime in might be lower due to constants such as $y! \leq k!$ which were absorbed by the big-O notation. This difference should be noticeable for the smaller values of n but not for higher values. Since the standard RLS is the same as the RLS₁^B the optimal values for the RLS_k^B variants again is at k = 1. This algorithm is also evaluated in the same Section 4.7 as the other algorithms that were analysed for inputs similar to a linear function.

3.3 Binomial distributed input

This section discusses inputs following a binomial distribution $\sim B(m,p)$. Since n is reserved for the size of the input m is used for the distribution instead. At first they seem uninteresting as all values are rather close to the expected value of the distribution. So it seems like the Evolutionary Algorithm only has to find a search point where both bins have an equal amount of values. After investigating this input experimentally this is not true to that extend. Solutions with an equal amount of bits with value 0 and value 1 are indeed close to the optimum, but they are just close and not optimal. There is mostly still a difference of at least one to the optimum. This might cause algorithms like the RLS to get stuck despite the input looking easy at first glance if there are no small elements. If elements can be swapped this input should become solvable again because there are many values slightly bigger or smaller than the expected value. If the algorithm switches those the small difference to the optimum can be closed. The first lemma tries to prove that these input are indeed easy to solve in a sense that this input is very likely to have a perfect partition.

Lemma 3.15. A binomial distributed input $\sim B(m,p)$ has a perfect partition $(b_F - b_E = 0$ for even W and $b_F - b_E = 1$ for uneven W) with high probability if the input size n is large enough.

Proof. Sketch:

- The initial distribution is likely rather close to the optimum
- The difference between the bins is probably not more than 10 expected values
- the large values

Consider a random separation of all values into two sets with equal size if n is even or one set with one value more than the other if n is odd. The sum X of one set is a sum of $\frac{n}{2} \cdot m$ independent Bernoulli trials with probability p. With Chernoff Bounds the following inequality follows:

$$\mathbb{P}(X \ge (\frac{n}{2} + \sqrt{\frac{n}{2}}) \cdot m \cdot p) = \mathbb{P}(X \ge (1 + 2\sqrt{\frac{2}{n}}) \cdot \frac{nmp}{2}) \le e^{-\frac{mnp}{2} \cdot 2\sqrt{\frac{2}{n}}^2/3} = e^{-\frac{2mp}{3}}$$

For $mp \geq 1.5$ the probability is less than $\frac{1}{e}$. Otherwise the input is rather trivial, since the numbers will be concentrated around $mp \leq 1.5$ and most values will be below 10. After moving $\mathcal{O}(\sqrt{\frac{n}{2}}/2)$ objects to the emptier set, the difference between the two sets is at most half the expected value mp of a single value. . . .

Lemma 3.16. With high probability the RLS does not find a perfect partition for an input with distribution $\sim B(m, p)$ if n is large enough, $mp \geq 50$ and $m - mp \geq 50$.

Proof. The RLS will always move one object per step from fuller to the emptier bin. As long as $b_F - b_E > w_i$ holds moving any object of weight at most w_i from b_F to b_E results in a decrease of the fitness $(b'_F = b_E + w_i < b_F - w_i + w_i = b_F)$. If the RLS does not decrease the difference $b_F - b_E$ to at most w_n in any step the solution is never optimal and the RLS therefore must be stuck. So now assume the RLS eventually reaches a search point with $b_F - b_E \leq w_n$. Consider the step which decreases the difference from more than w_n to at most w_n . If this step does not decrease the difference to 0 for even n or 1 for odd n the RLS is stuck due to Corollary 3.7. For a step to decrease $b_F - b_E$ from more than w_n to 0 the RLS must move an object of weight y which is given by $b_F' - b_E' = (b_F - y) - (b_E + y) = 0 \Leftrightarrow b_F - b_E - 2y = 0 \Leftrightarrow y = (b_F - b_E)/2$. Such an element can only exist for even n because only for even n either both bins have an even sum or neither of them. Odd inputs will always have exactly one bin with an even sum and one with an odd sum. For odd n a difference of 1 suffices for a perfect partition and therefore the value of y must be either $\lceil (b_F - b_E)/2 \rceil$ or $\lceil (b_F - b_E)/2 \rceil$. For any binomial distribution the value which occurs the most in expectation is the expected value if $mp \in \mathbb{N}$ or $\lceil mp \rceil$ and |mp| otherwise. Even if the behaviour of the algorithm forces values smaller or bigger than mp in the emptier bin, the number of bigger/smaller values in the fuller bin will still be less in expectation than the amount of mp in both bins. Let's assume that objects with exactly this volume must be selected and that there are two options because $mp \notin \mathbb{N}$. This will give an upper bound on the probability of flipping the right object in the crucial step. The probability of a number to be $\lfloor mp \rfloor$ is $p^{\lfloor mp \rfloor}(1-p)^{m-\lfloor mp \rfloor} \leq 0.5^{50} = \frac{1}{1024^5} < 10^{-15}$ because both $mp \geq 50$ and $m-mp \geq 50$ and $p \leq 0.5$ or $1-p \leq 0.5$. For $\lceil mp \rceil$ the same bound applies with the same arguments. The probability of either of these numbers to be drawn from the binomial distribution is at most $2 \cdot 0.5^{50} = 0.5^{49} \le 10^{-15}$. This leads to $n \cdot 10^{-15}$ values with the right value in expectation. All bits are flipped uniformly at random and therefore the probability of flipping a good bit is at most $\frac{n \cdot 10^{-15}}{|b_F|} \leq \frac{n \cdot 10^{-15}}{n} = 10^{-15}$.

This means the RLS is very unlikely to find a perfect partition. Given the solution has a perfect partition the RLS will get stuck in a local optimum. \Box

The restriction of $mp \ge 50$ and $m - mp \ge 50$ was not chosen to make the proof work but is also necessary. Without those restrictions the distribution might have many small values and especially elements close to 1. When there are many small values these can be used to fill the small gaps which makes it easy for the RLS to find a perfect partition. On the other hand if p is almost one, then almost every element will be same for smaller values of m. If every element is the same then the RLS must only find a search point with equal amounts of 0s and 1s.

4. Experimental Results

In the following chapter the different variants of the RLS and the (1+1) EA are now analysed empirically for the best algorithm depending on the input. Additionally for most lemmas from the previous chapters there are also tests if they actually hold in practice.

4.1 Code

The complete java code used for all empirical studies is available on GitHub.

4.1.1 The Algorithms

All different variants of the RLS function more or the less the same. They start with an initial random value and then optimise this one value in the loop. The loop can be summarised like this:

- 1. generate a number k of bits to be flipped (algorithm specific)
- 2. flip k random bits
- 3. evaluate fitness of the mutated individual
- 4. replace old value with new value if new value is better
- 5. repeat if not optimal

The (1+1) EA variants behave differently at first glance as they flip each bit independently with probability c/n. This can be seen as n independent Bernoulli trials with probability c/n. The amount of bits that are flipped is therefore binomial distributed and the algorithm can be implemented exactly as the versions of the RLS. The same holds for the pmut operator which generates a number k from a powerlaw distribution and then flips k bits. This leads to only one implementation of a partition solving algorithm which is not only given the input array of numbers but also a generator for the amount of bits to be flipped in each step. The random values for the amount of bits to be flipped are generated according to this table:

Algorithm	Returned value
RLS	1
${\rm RLS}_k^B$	$y \in \{1, \dots, k\}$ with probability $\frac{\binom{n}{y}}{\sum_{i=1}^{k} \binom{n}{i}}$
RLS_k^S	uniform random value $y \in \{1, \dots, k\}$
$(1+1)^{\kappa} EA$	binomial distributed value from $\sim B(n, c/n)$
pmut	powerlaw distributed value from $\sim D_{n/2}^{\beta}$

Algorithm 4.1: GENERIC PARTITION SOLVER

```
1 choose x uniform random from \{0,1\}^n
2 while x not optimal do
3 x' \leftarrow x
4 k \leftarrow \text{kGenerator.generate}()
5 flip k uniform random bits of x'
6 if f(x') \leq f(x) then
7 x \leftarrow x'
```

4.1.2 Random number generation

Java only provides a random number generator for uniform distributed values for any integer interval or random double values $\in [0,1)$. The same holds for the Mersenne Twister with an implementation used from this page. All experiment were executed with both uniform random number generators. The results were rather similar, so only the results for the Fast Mersenne Twister are shown. For this project uniform random numbers do not suffice as for an efficient way of implementing the (1+1) EA or simply for generating a binomial distributed input another random number generator is needed. One of the needed distributions is a binomial distribution. The simplest way to generate a number $\sim B(m,p)$ would be to run a loop m times and add 1 to the generated number if a uniform random value $\in [0,1)$ is less than p. This works perfectly fine and generates numbers according to the distribution. With low values for p this approach is rather inefficient and especially for values of p = 1/m. The expected value in this case is 1 but generating a random number takes time $\mathcal{O}(m)$. Another more efficient way was implemented by StackOverflow user pjs on stackoverflow inspired by Devroyes method introduced in [Dev06]. This method has an expected running time of $\mathcal{O}(mp)$ which is equal to the expected value of the distribution. For the case of p=1/m this runs in expected constant time in comparison to $\mathcal{O}(m)$ for the naive way. This number generation was also used for the implementation of the (1+1) EA instead of running a loop in every step. Algorithm 4.2 uses the second waiting time method which uses the fact that $X \sim B(m, p)$ if X is the smallest integer so that $\sum_{i=1}^{X+1} \frac{E_i}{n-i+1} < -\log(1-p)$ for E_i iid exponential random variables (Lemma 4.5 section X.4 [Dev06]).

Algorithm 4.2: BINOMIAL RANDOM NUMBER GENERATOR

The next generator needed is for geometric distributed values. This generator is only necessary for the generation of geometric distributed inputs but not for the algorithms. The easiest way to generate geometric distributed values is the naive way: generating a uniform random value p' until p' < p holds. The expected running time of this algorithm is

equal to the expected value of the distribution 1/p. So this method is comparably effective to the approach used for binomial random number generation.

Algorithm 4.3: Geometric random number generator

```
1 sum \leftarrow 0
// random() generates a random value \in [0,1)
2 while random() \ge q do
3 \mid sum \leftarrow sum + 1
4 return sum
```

The last generator needed is for powerlaw distributed values. This generator is in contrast to the geometric number generator needed for both the algorithm with the $pmut_{\beta}$ mutation operator and for generating inputs. This implementation is also from stackoverflow. The user gnovice provided the following formula on this page on stackoverflow:

$$x = \left[(b^{n+1} - a^{n+1}) * y + a^{n+1} \right]^{1/(n+1)}$$

a is the lower bound, b the upper bound, n the parameter of the distribution and y the number generated uniform random $\in [0,1)$. The idea behind the formula and the formula itself is explained in a mathworld page. For a powerlaw distribution $P(x) = Cx^n$ for $x \in [a,b]$ normalisation gives

$$\int_{a}^{b} P(x)dx = C \frac{[x^{n+1}]_{a}^{b}}{n+1} = 1 \Leftrightarrow C = \frac{n+1}{b^{n+1} - a^{n+1}}$$

Let Y be a uniformly random distributed variate on [0,1]. Then

$$D(x) = \int_{a}^{x} P(x')dx' = C \int_{a}^{x} x'^{n} dx' = \frac{C}{n+1} (x^{n+1} - a^{n+1}) = \frac{(x^{n+1} - a^{n+1})}{b^{n+1} - a^{n+1}} \equiv y$$

and the variate is given by

$$X = \left[(b^{n+1} - a^{n+1}) * y + a^{n+1} \right]^{1/(n+1)}$$

The values inserted in this formula must be negative. In the original paper for the $pmut_{\beta}$ operator and in the definition normally a powerlaw distribution is $P(x) = Cx^{-n}$ and therefore any positive value for n in this case was negated. Apart from this negation the generator was not changed.

4.2 Do binomial inputs have perfect partitions?

Lemma 3.15 is only valid for larger n. In practice the bound is much smaller depending on the expected value of a single number. Another factor deciding how likely an input is to have a perfect partition is whether n is even or odd. To determine the influence of all factors multiple experiments were conducted. The goal of the first experiment was to determine the influence of the array size to the input having a perfect partition and the fact if n is even or odd. So for every possible combination of $p \in \{0.1, 0.2, \dots, 0.8, 0.9\}, m \in \{10, 100, 1000, 10^4, 10^5\}$ and $n \in \{2, 3, 4, \dots, 19, 20\}$ 1000 randomly generated inputs of size n were tested for a perfect partition. Due to the small values for n it was possible to brute force the results in a short amount of time. The results are visualised in figure 4.1 to figure 4.9.

On the x-axis is the size of the input and on the y-axis the percentage of inputs that had a perfect partition. The different graphs in each figure resemble the different values of p

80% 70% 60% 50% 40% 30% 10 11 12 Input size

Figure 4.1: Percentage of Binomial inputs with perfect partitions for m=10

used for generating the inputs. The graph for p = 0.1 resembles the percentage of inputs that had a perfect partition with values generated from the distribution $\sim B(m, 0.1)$ with m being dependent on the figure. For figure 4.1 m has the value 10.

Figure 4.1 is a bit overloaded with information and the zigzag makes it hard to gain any benefit from the graphs. That's why for $m \in \{10, 100, 1000\}$ there is one figure for the even input sizes of n and one for the odd. Figures which show only results for either even or odd values of n have dotted graphs, because the values in between the points do not exist. The dotted lines are only in the figure for a better visualisation of the trend and not meant for interpretation apart from the marked values. For $n \ge 10,000$ all values for the odd input sizes are 0 %, so there is no point in showing the data in a separate figure.

Figure 4.2: Percentage of Binomial inputs with perfect partitions for m = 10 for even n

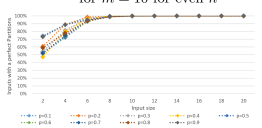


Figure 4.3: Percentage of Binomial inputs with perfect partitions for m = 10 for odd n

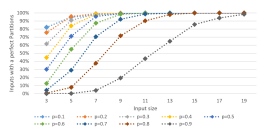


Figure 4.4: Percentage of Binomial inputs with perfect partitions for m = 100 for even n

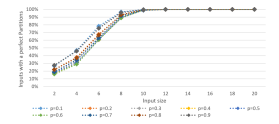
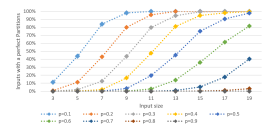


Figure 4.5: Percentage of Binomial inputs with perfect partitions for m = 100 for odd n



It is easy to see that for small inputs sizes it is relevant if n is even or odd for higher expected values as all curves in figure 4.9 oscillate between 0% and 100% for $n \ge 14$. For odd inputs the probability of a perfect partition decreases much more drastically with mas for even inputs because the expected value of a single number increases with m. If all values are much higher the small differences between the values can no longer even out the fact of one set having more elements than the other. The oscillation therefore increases with increasing m. For n=20 all 1000 inputs had a perfect partition for every combination

Figure 4.6: Percentage of Binomial inputs with perfect partitions for m = 1000 for even n

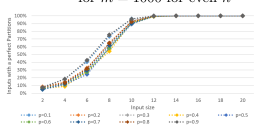


Figure 4.8: Percentage of Binomial inputs with perfect partitions for m = 10,000

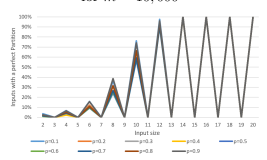


Figure 4.7: Percentage of Binomial inputs with perfect partitions for m = 1000 for odd n

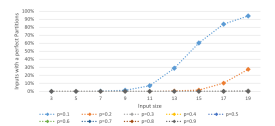
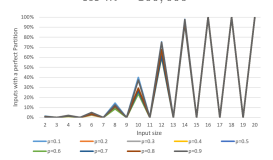


Figure 4.9: Percentage of Binomial inputs with perfect partitions for m = 100,000



of p and m but for n=19 only combinations where $mp \leq 300$ holds had at least one input with a perfect partition. For expected values of up to 10^5 it seems to be almost granted that an input of length 20 has a perfect partition if it is binomial distributed. Even for only 12 binomial generated values more than 50% of the inputs had a perfect partition (see figure 4.9). Another visible effect is the decreasing percentage with rising p. This may be a direct result of the value chosen for p but can also be an indirect result as the value for p changes the expected value for a constant p. The expected value may have an influence on the number of perfect partitions because it influences the highest value of the input. For uniform distributed inputs Borgs et. al. showed that the coefficient of number of bits needed to encode the max value/p has a huge impact on the number of perfect partitions [BCP01]. For a coefficient p 1 it tends to 0. This was only proven for the uniform distributed input, but it might also hold for a binomial distributed input. This leads to the second experiment.

In the second experiment the inputs were generated a bit differently. Here the goal was to keep the expected value fixed for any combination of p and n and set the value of m to e/p for all $e \in \{10, 20, 30, 40, 50, 100, 200, 500, 1000, 2000, 5000, 10000, 50000\}$ so that $E(X) = mp = e/p \cdot p = e$. With this setup the influence of the expected value is almost isolated from the other parameters. The probability is still linked to p as p also influences the variance mp(1-p).

Figure 4.10 to figure 4.13 again show the percentage of perfect inputs with different settings of m, p, n. The x-axis is the expected value mp of a single number of the input. The different graphs show the percentage for different input sizes. It seems as if the value of p has a much smaller influence than the expected value. For a fixed expected value and a fixed input size a higher value for p seems to only slightly increase the percentage of inputs with a perfect partition. The expected value influences the percentage significantly more. For p = 0.1, n = 14 the value decreases from 100% at E(X) = 10 to below 20% at

E(X) = 50000 (figure 4.10). For p = 0.9 the percentage only drops below 50% but still decreases by a factor of 2 (figure 4.13).

Figure 4.10: Percentage of Binomial inputs with perfect partitions for p=0.1

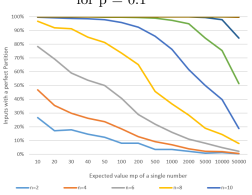


Figure 4.11: Percentage of Binomial inputs with perfect partitions for p=0.2

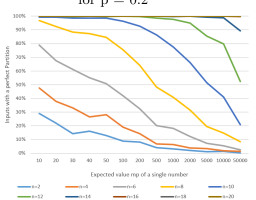


Figure 4.12: Percentage of Binomial inputs with perfect partitions

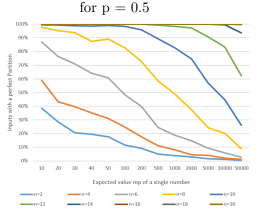
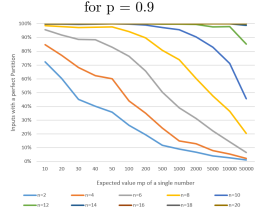


Figure 4.13: Percentage of Binomial inputs with perfect partitions



The last experiment showed that for n=20, 1000/1000 inputs had a perfect partition. This raised the question of how the amount of perfect partition changes with changing values for m, p, n. Figure 4.14 to figure 4.17 show the amount of perfect partitions a binomial distribution $\sim B(m,p)$ has. For these figures 10,000 random binomial inputs with the given values for m and p were generated. Each input was then tested for the number of perfect partitions it has. The used method was again brute force to ensure correctness which was only possible due to the small input sizes. After all runs the average values were combined in the given figures. The value of p is dependent on the picture and each value of $m \in \{10, 100, 1000, 10000\}$ has its own graph within the figure. The x-axis is the size of the input and the y-axis the number of perfect partitions the input has. Notice that all graphs have a y-axis with a logarithmic scale. Since the graphs are all linear the actual values rise exponentially. The number of perfect partitions is mostly multiplied by a factor between 3 and 4 when the input size increases by 2.

The higher the value of m the closer the curves of p and 1-p get. For m=1000 and m=10000 the values are almost the same for every input size. For p=0.1, m=10 an input with expected values of 1 seems to much more likely to have a perfect partition than an input with expected value $10 \cdot 0.9 = 9$. With growing m this has less impact.

Figure 4.14: Amount of perfect partitions for p = 0.1

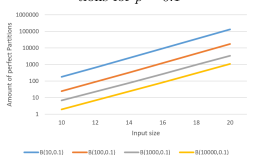


Figure 4.15: Amount of perfect partitions for p = 0.9

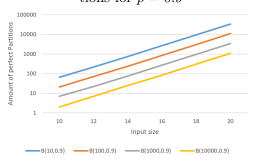


Figure 4.16: Amount of perfect partitions for p = 0.2

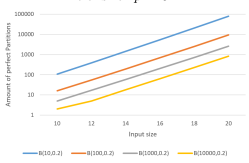
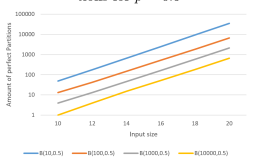


Figure 4.17: Amount of perfect partitions for p = 0.5



So the binomial input should be easy to solve due to the exponential number of perfect partitions. It might be harder for the smaller values of n as there are only a few perfect partitions. Due to the small number of total possibilities it should still be easy to solve for the small values of n as long as the RSH is not stuck in a local optimum. The number of iterations might be high in terms of the big-O notation but should still be small in the absolute value.

4.3 Binomial distributed values

In the following subsections the performance of the different algorithms is tested for different kinds of inputs. The exact distributions of the input are explained separately in each subsection. The procedure for each comparison is always the same. A random input is generated according to the distribution and then solved by every algorithm. All algorithms had the same two stopping conditions. The first was reaching a perfect partition and the second was taking more than $10 \cdot n \ln(n)$ steps. For the lower values of n the step limit of 100,000 was used instead. For n = 20 giving the algorithm only 600 steps is rather small. In some cases the smaller inputs are even more difficult to solve. Most modern computer should be able to handle 100,000 iterations in a short amount of time anyway. So the minimum step limit of 100,000 seemed reasonable. If either of these conditions was met, the algorithm returned its current best solution. This step is repeated 10,000 times. The results are presented in a table containing multiple statistics for each algorithm over all 10,000 runs. The data is explained in the table below.

column name	meaning
algo type	type of algorithm (RLS, RLS_k^B , RLS_k^S , (1+1) EA or pmut)
algo param	parameter of the algorithm or '-' if it is the standard variant
avg mut/change	average #bits flipped for iterations leading to an improvement
avg mut/step	average #bits flipped for any iteration
total avg count	average #iterations for all runs
avg eval count	average #iterations of runs returning an optimal solution
max eval count	maximum #iterations of runs returning an optimal solution
min eval count	minimum #iterations of runs returning an optimal solution
fail ratio	ratio of unsuccessful runs to all runs
avg fail dif	average value of $f(x) - f(opt)$ for non-optimal solutions
p-value	the p-value of a Mann-Whitney-U-Test between two columns

Sometimes every algorithm managed to find an optimum in each run. To avoid redundancy and shorten the thesis the rows 'total avg count' and 'avg fail dif' are not shown in the table in those cases. The p-value between two columns refers to the p-value of a Mann-Whitney-U-Test between the algorithms in the columns left and right to the value. H_0 is: the algorithm on the left is not faster than the algorithm on the right. H_1 in this case then is: the algorithm once the left is better than algorithm on the right. So if the p-value is below 0.05 the algorithm on the left is statistically significant faster.

Firstly the different variants of the RLS are compared with values of $k \in \{2, 3, 4\}$, then the performance of the (1+1) EA with static mutation rate c/n with $c \in \{1, 2, 3, 5, 10, 50, 100\}$ and lastly the performance of the $pmut_{\beta}$ mutation operator with the parameter $\beta \in$ $\{1.25, 1.5, \ldots, 2.75, 3.0, 3.25\}$. Afterwards there is a comparison for multiple input sizes of the best algorithms because the best algorithm is often dependent on the size of the input. Normally there are three tables for each input. The first states how often the algorithms did not find an optimal solution for the different input sizes ('fails in 10000 runs' in top left cell). The second gives their average performance for the successful runs ('avg' in top left cell) and the last the performance for all runs ('total avg' in top left cell). The last two tables differ in the unsuccessful runs. Often the algorithm is stuck in a local optimum it won't leave in reasonable time or even never for variants of the RLS. In these cases the step limit is the deciding factor on how big the penality for this run is. So neither of the two average values alone is enough to give a complete insight on the performance. Sometimes a variant of the RLS is much faster than the other algorithms for a specific input but is also the only algorithm to get stuck in a local optimum. This creates the possibility to start the RLS variant with a low step limit and switch to the (1+1) EA if the RLS variant does not return an optimal solution. Giving both tables for the different average values might help with this decision.

The first analysed inputs are inputs following a binomial distribution $\sim B(m,p)$ as those inputs have been researched in the previous subsection. The results showed that the expected value of a single number is the main driver for the amount of perfect partitions the input has. The results also suggested the inputs tend to have more perfect partitions if the expected value is lower. The more perfect partitions an input has relative to the number of all possible partitions, the more likely the different RSHs are to find one of those. Therefore researching inputs with higher expected values seems more interesting but generating higher values takes more time with a random number generator that needs $\mathcal{O}(mp)$ time. To keep the time for generating one set of numbers reasonable the values chosen for all tests are m=10000, p=0.1, n=10000 with the expected value for a single number being mp=1000. Figure 4.18 shows a random binomial distributed input of length n=10000. For this input type almost every time all elements were sharply concentrated around the expected value with all values being at 1000 ± 200 (for figure 4.18 even closer at

 1000 ± 121). So after reaching a difference to the optimum of below (1000 - 200)/2 = 400 the algorithm can no longer achieve an improvement by flipping a single bit (Corollary 3.7)



Figure 4.18: Distribution of a random binomial input

4.3.1 RLS Comparison

algo type	RLS^B_b	RLS^B_b	RLS_s^S	RLS_s^S	RLS_s^S	RLS^B_b	RLS
algo param	b=2	b=4	s=2	s=4	s=3	b=3	-
avg mut/change	2.000	4.000	1.602	2.563	2.000	2.735	1.000
avg mut/step	2.000	4.000	1.500	2.500	2.000	3.000	1.000
total avg count	295	409	454	538	636	488,329	919,832
avg eval count	295	409	454	538	636	380,018	103
max eval count	1,900	3,717	$3,\!562$	4,300	6,057	$920,\!419$	199
min eval count	4	2	5	4	3	9	9
fail ratio	0.000	0.000	0.000	0.000	0.000	0.200	0.999
avg fail dif	-	-	-	-	-	1	246
p-value	0.0	0.00	0.00	0.0	0.0	0.0	000

The RLS_2^B seems to perform the best as it mostly switches two elements which works great for binomial distributed inputs. The same algorithm with b=4 performs a bit worse but still good as switching 4 elements can be beneficial as well. The variant of RLS³ on the other hand does not reach the optimal solution in 20% of the inputs with an average difference of 1. It also needs 1000 times more iterations to find an optimum on average compared to the best algorithm RLS_2^B . The RLS_s^S variants behave mostly the same with s=2 being the best, followed by s=4 and s=3. In this case the variant of s=3 is by far not as bad as for the RLS_3^B because the probability of flipping 2 bits is 1/3 as compared to $\mathcal{O}(n^{-1})$ for the RLS^B₃. The RLS^S_k all seem to be a good option for binomial inputs with values of $k \in \{2,3,4\}$. The standard RLS on the other hand performs by far the worst as it only moves one element per step. It only managed to reach the optimal solution 13 times for 10,000 different inputs. The average number of iterations for those inputs was only 103 so the RLS likely had a good initialisation with a few lucky steps leading directly to the optimum. For all other cases the average difference between the bins was 246 which is close to the median of the values from 0 to (1000 - 100)/2 = 450. This is likely due to the RLS being unable to improve the solution once the current solution has a difference below half of the lowest value (Corollary 3.7).

4.3.2 (1+1) EA Comparison

For the (1+1) EA the best static mutation rate seems to be 3/n. The probability of flipping 2 or 4 bits as n goes to infinity for mutation rate 1/n approaches $13/24e \approx 0.199$, for 2/n approaches $8/3e^2 \approx 0.361$, for 3/n approaches $63/8e^3 \approx 0.392$, for 4/n approaches $56/3e^4 \approx 0.342$ and for 5/n approaches $77/2e^5 \approx 0.259$. So the highest probability has

c=3, followed by c=4 and c=2 then c=5 and lastly c=1. For higher values of c the probability decreases further as the expected number of flipped bits is c for mutation rate c/n.

algo type	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM
algo param	3/n	4/n	2/n	5/n	-	10/n	50/n	100/n
avg mut/change	3.103	3.953	2.339	4.861	1.695	9.727	49.587	99.529
avg mut/step	3.000	4.000	2.000	4.999	1.000	10.000	50.001	100.001
avg eval count	594	642	645	731	1,080	1,370	7,052	13,624
max eval count	6,084	$5,\!368$	$6,\!151$	8,083	8,767	$18,\!297$	$113,\!206$	$155,\!424$
min eval count	3	3	0	3	7	7	3	5
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
p-value	0.0	0.0	0.0 0.0	001 0.0	0.0 0.0	0.0	0.0 0.0	000

The static mutation rate 3/n seems to perform the best with both 4/n and 2/n being a close second place. The next best values are 5/n and the standard 1/n. From then on the number of iterations rises monotonically with rising mutation rate. The higher mutation rates perform significantly worse but the still find a solution within the limit as opposed to the standard RLS.

4.3.3 pmut Comparison

algo type	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut
algo param	2.00	2.25	2.50	1.75	2.75	3.00	3.25	1.50	1.25
avg mut/change	6.667	3.890	2.815	15.093	2.289	2.009	1.833	38.481	95.474
avg mut/step	8.413	4.356	2.900	22.396	2.270	1.935	1.729	70.698	224.871
avg eval count	610	612	629	641	645	677	717	721	974
max eval count	6,462	5,408	7,478	5,681	6,321	8,262	6,415	8,077	11,014
min eval count	3	0	1	3	7	8	3	7	3
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
p-value	0.18	821 0.00	0.1	.088 0.0	004 0	.0000 0	0.0000	0.0000 0.0	0000

For the $pmut_{\beta}$ mutation operator the choice of β seems to be much more insignificant than for the RLS or (1+1) EA. Here all values perform comparably good with only the value of $\beta = 1.25$ having a clear performance difference compared to next best value. All values of β reach an optimal solution in every case. The worst variant of the $pmut_{\beta}$ operator still performs much better than the worst value for the (1+1) EA and even better than the worst RLS variant. There is no clear best algorithm because there is no statistically significant difference between the best to values.

4.3.4 Comparison of the best variants

Looking at the previous tables for this setting of m = 10000, p = 0.1, n = 10000 the RLS₂^B performs better than the (1+1) EA and $pmut_{\beta}$ mutation for all values of c/n and β by a factor of at least 2. This is likely from the fact that this version of the RLS flips almost only two bits which seems to be close to optimal for this kind of input. There are many values close to the expected value which can be switched to make small adjustments to the fitness value. To further investigate which algorithm performs best on all binomial distributed inputs now a comparison with different input lengths follows. The parameters of the distribution were not changed.

The following table shows the number of runs in which the algorithms did not find an optimal solution within $\max\{100,000;10\cdot n\ln(n)\}$ steps for different input sizes of n. For $n \leq 1382$ the step limit is 100,000 and for the bigger values of n it is $10\cdot n\ln(n)$.

fails in 10000 runs	20	50	100	500	1000	5000	10000	50000
RLS_2^B	2158	241	28	0	0	0	0	0
RLS_4^B	0	3	14	0	0	0	0	0
RLS_2^S	2101	9	0	0	0	0	0	0
(1+1) EA (3/n)	0	0	0	0	0	0	0	0
(1+1) EA (4/n)	0	0	0	0	0	0	0	0
pmut (2.0)	0	0	0	0	0	0	0	0
pmut (2.25)	0	0	0	0	0	0	0	0

The (1+1) EA and $pmut_{\beta}$ always reach an optimal solution but the RLS variants do not. The RLS variants that can only flip two bits per step perform significantly worse for very small inputs. They are probably more likely to get stuck in a local optimum where a step flipping 4 bits or more would be necessary. So the RLS₂^B does perform better for larger inputs but is much more likely to get stuck in a local optimum for smaller n values. The next table contains the average number of iterations the algorithm needed to find an optimal solution for all runs where the algorithms managed to find an optimal solution. Here it still looks like the RLS₂^B finds the solution with the lowest amount of steps, because the cases where the algorithm is stuck in a local optima are not contained in this table.

avg	20	50	100	500	1000	5000	10000	50000
RLS_2^B	158	227	317	237	244	271	291	394
RLS_4^B	332	335	639	377	375	394	407	474
RLS_2^S	266	381	362	387	403	428	455	551
(1+1) EA (3/n)	602	498	502	549	555	587	604	672
(1+1) EA $(4/n)$	525	547	557	595	611	634	641	716
pmut (2.0)	881	515	522	563	578	602	611	670
pmut (2.25)	1058	528	513	552	566	596	606	685

The next table contains the overall average amount of iterations for every run. So runs where no optimal result was found add 100,000 to the sum of all iterations for small values of n.

total avg	20	50	100	500	1000	5000	10000	50000
RLS_2^B	21704	2631	596	237	244	271	291	394
RLS_4^B	332	365	778	377	375	394	407	474
RLS_2^S	21220	470	362	387	403	428	455	551
(1+1) EA (3/n)	602	498	502	549	555	587	604	672
(1+1) EA (4/n)	525	547	557	595	611	634	641	716
pmut (2.0)	881	515	522	563	578	602	611	670
pmut (2.25)	1058	528	513	552	566	596	606	685

Here the RLS₂^B is only the best algorithm for values of $n \ge 500$. Below this bound choosing the (1+1) EA with static mutation rate 3/n is a safer choice as the (1+1) EA reaches an optimal solution for every input (in this experiment).

4.4 Geometric distributed values

For the geometric distribution the chosen default value is p=0.001. This results in an expected value of 1000 which is the same as for the binomial distribution in the last subsection. This should make the results more comparable. The maximum value is theoretically not limited but for the implementation in Java the maximum value was set to the maximum value of a long value $=2^{63}-1=9,223,372,036,854,775,807$. Without this maximum the value might overflow and instead be negative with high absolute value. Figure 4.19 shows a random geometric distributed input. The span of all values is way higher than for the binomial distribution, although they have same expected value. Here

the values are not in the interval [800, 1200] but rather between 0 and 9000. The theoretical limitation of the values being at most $2^{63} - 1$ seems to not have an influence on the results. The geometric distribution does not only have low values close or equal to 1 but also has mostly values that are very small. This should lead to 1-bit flips being effective as the small values can remove the small differences. Because there are so many small values moving only one bit might be better than switching two elements.

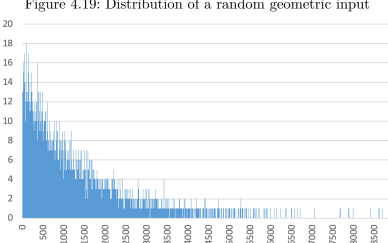


Figure 4.19: Distribution of a random geometric input

4.4.1 RLS Comparison

algo type	RLS_s^S	RLS_s^S	RLS_s^S	RLS^B_b	RLS	RLS^B_b	RLS_b^B
algo param	s=2	s=3	s=4	b=2	-	b=3	b=4
avg mut/change	1.483	1.958	2.428	2.000	1.000	3.000	4.000
avg mut/step	1.500	2.000	2.500	2.000	1.000	3.000	4.000
total avg count	2,218	2,597	2,924	3,130	3,816	4,062	4,853
avg eval count	2,218	2,597	2,924	3,130	1,886	4,062	$4,\!853$
max eval count	34,066	$38,\!333$	$48,\!275$	$43,\!116$	$68,\!298$	$46,\!410$	$51,\!427$
min eval count	0	0	1	7	4	4	0
fail ratio	0.000	0.000	0.000	0.000	0.002	0.000	0.000
avg fail dif	-	-	-	-	1	-	-
p-value	0.0	0.00	0.0	0.0	0.0	0.0	000

For these inputs the variants of the RLS perform differently to the binomial input. The only similarity is the RLS being the only algorithm that did not find an optimal solution for every input. If the RLS did find an optimal solution in those 21 cases it instead might be the best RLS variant. The other algorithms are ranked by their probability of flipping only one bit. This means at first the three ${\rm RLS}_s^S$ variants from 2 to 3 to 4 and then the same for the RLS^B_b variants. So it does seem like moving mostly one element at once is better for the geometric input in comparison to two elements for the binomial distribution. In the 21 cases where the RLS did not find an optimal solution it was most likely stuck in a local optimum where no small value was left.

4.4.2	(1+1)	$\mathbf{E}\mathbf{A}$	Com	parison
-------	-------	------------------------	-----	---------

algo type	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM
algo param	2/n	-	3/n	4/n	5/n	10/n	50/n	100/n
avg mut/change	2.246	1.551	3.048	3.936	4.861	9.822	49.750	99.707
avg mut/step	2.000	1.000	3.000	4.000	5.000	10.000	50.000	100.001
avg eval count	3,097	3,505	3,518	4,009	4,807	7,758	18,457	25,993
max eval count	$39,\!490$	$60,\!533$	39,048	$47,\!881$	56,204	$91,\!305$	$173,\!851$	$354,\!479$
min eval count	10	0	6	5	3	5	9	3
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
p-value	0.0	000 0.2	110 0.0	0.0	0.0	0.0	0.0	000

The results for the (1+1) EA are similar to the results of the RLS. From mutation rate 2/n on the runtime increases with rising mutation rate. The only part that does not fit into the theory of 1 bit flips being superior is the mutation rate 2/n performing better than the standard 1/n. There is also no statistically significant difference between 1/n and 3/n. All variants reach an optimal solution within the given limit for the number of iterations.

4.4.3 pmut Comparison

algo type	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut
algo param	3.25	3.00	2.75	2.50	2.25	2.00	1.75	1.50	1.25
avg mut/change	1.689	1.862	2.162	2.681	3.834	6.885	15.848	41.832	104.749
avg mut/step	1.729	1.934	2.270	2.905	4.367	8.500	22.219	70.666	224.556
avg eval count	2,207	2,268	2,376	2,480	2,586	2,813	3,196	3,898	5,399
max eval count	40,626	45,193	39,295	41,558	40,339	46,110	43,223	61,121	60,767
min eval count	0	5	9	5	7	2	3	1	10
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
p-value	0.0	714 0.00	0.0	0.00	0.0	0.00	0.00	0.0	0000

The results for the $pmut_{\beta}$ operator are even more clear than for the (1+1) EA. With decreasing values for β the amount of flipped bits per step increases. The performance decreases as well with decreasing values for β which fits into the theory of one bit flips being better for geometric distributed inputs. The number of repetitions of the algorithm might simply be too small to make the small difference in the performance between the two values visible. The difference in the performance for the $pmut_{\beta}$ operator is not as drastic as for the (1+1) EA. Only $\beta = 1.5$ and $\beta = 1.25$ perform much worse the next best value.

4.4.4 Comparison of the best variants

The setup for the evaluation of lower values for n is the same as for binomial distributed inputs. The first table lists the number of runs where the different algorithms did not find the optimal solution within the time limit.

fails in 10000 runs	20	50	100	500	1000	5000	10000	50000
RLS	9805	9457	8975	5994	3941	325	24	0
RLS_2^S	8936	5357	1396	0	0	0	0	0
(1+1) EA (1/n)	4192	1022	327	12	6	0	0	0
(1+1) EA (2/n)	1165	53	6	0	0	0	0	0
pmut (3.25)	2285	481	146	0	0	0	0	0
pmut (3.0)	1712	334	94	0	0	0	0	0

For small inputs the geometric distributed input seems to have inputs without a perfect partition because there were many iterations where neither of the algorithms found an optimal solution within the time limit. It is still likely to have a perfect partition even for the small values in comparison to other distributions which follow afterwards. Many algorithms especially the variants of the RLS seem to be likely to get stuck in a local

optimum. The (1+1) EA finds an optimum in most of the runs, so the geometric distributed inputs also seem to be likely to have a perfect partition for small values. They definitely are harder to solve for smaller input sizes than the binomial inputs, but they still have a perfect partition most times. The next table visualises the average number of iterations the algorithms needed for finding an optimal solution if the algorithm managed to do so.

avg	20	50	100	500	1000	5000	10000	50000
RLS	32	79	153	579	950	1859	1922	1797
RLS_2^S	391	2124	5005	4218	3530	2362	2160	2229
(1+1) EA (1/n)	22471	18343	12834	8342	6511	3815	3458	3371
(1+1) EA (2/n)	16360	9243	6452	4503	4020	3171	3141	3133
pmut (3.25)	23440	15929	9658	5644	4406	2434	2162	2172
pmut (3.0)	21901	14696	9186	5222	4150	2510	2208	2213

The variants of the (1+1) EA and of the pmut algorithm seem to take about 20,000 iterations for n=20 if they manage to find the optimal solution. They also perform better and better the bigger the input gets. This is probability caused by the many additional small values that can be used for smaller adjustments to the fitness. Also a really high value does not have as much of an effect, because there are possibly other larger values which cancel each other out, if they are in different bins. The standard (1+1) EA does not only find a perfect partition less often than the (1+1) EA with $p_m = 2/n$, it also needs more iterations on average if it does. So the (1+1) EA with $p_m = 2/n$ performs indeed better for every input size. The last table again lists the total average number of steps.

total avg	20	50	100	500	1000	5000	10000	50000
RLS	98050	94574	89765	60172	39985	15639	4128	1797
RLS_2^S	89401	54556	18266	4218	3530	2362	2160	2229
(1+1) EA (1/n)	54971	26688	15684	8452	6567	3815	3458	3371
(1+1) EA (2/n)	26104	9724	6508	4503	4020	3171	3141	3133
pmut (3.25)	40934	19972	10977	5644	4406	2434	2162	2172
pmut (3.0)	35272	17545	10040	5222	4150	2510	2208	2213

The RLS is only an option if the input is large enough $(n \ge 50,000)$. For smaller input sizes especially for $n \le 1000$ choosing the (1+1) EA with mutation rate 2/n seems like the best choice. For larger values this (1+1) EA does not find an optimal solution the fastest but is still fast enough to be a viable option. Another rather save option is $pmut_{3.25}$. This algorithm performs worse for $n \le 1000$ but is still good in comparison to the other algorithms. For n > 1000 $pmut_{3.25}$ starts to outperform the best version of the (1+1) EA and almost all other researched algorithms.

4.5 Uniform distributed inputs

For the uniform distribution the default values were 1 for the lower bound and 50000 for the upper bound (exclusive). The range was limited to 50000 to reduce the time the algorithms needs to find an optimal solution. The higher the values are with too few values the more likely the input is to not have a perfect partition [BCP01]. This will cause the algorithms to always reach the limit for the number of iterations which drastically increases the time needed for the experiment. The length of the input was 50000.

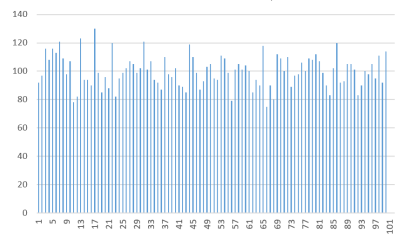


Figure 4.20: Distribution of a random uniform input (10000 values between 1 and 100)

4.5.1 RLS Comparison

algo type	RLS_b^B	RLS_s^S	RLS_s^S	RLS_s^S	RLS^B_b	RLS^B_b	RLS
algo param	b=2	s=3	s=4	s=2	b=3	b=4	-
avg mut/change	2.000	1.991	2.481	1.501	3.000	4.000	1.000
avg mut/step	2.000	2.000	2.500	1.500	3.000	4.000	1.000
total avg count	73,797	93,552	95,746	96,698	101,538	112,431	2,134,880
avg eval count	73,797	$93,\!552$	95,746	96,698	$101,\!538$	$112,\!431$	$45,\!419$
max eval count	$727,\!262$	1,216,841	1,769,247	1,954,892	1,211,069	$1,\!532,\!872$	404,899
min eval count	23	30	40	36	15	27	38
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.390
avg fail dif	-	-	-	-	-	-	1
p-value	0.0	000 0.03	219 0.00	0.00	0.00	0.00	000

The picture for the RLS variants on this type of input is not clear. There in no obvious tendency for all variants. The only obvious thing is the RLS being the worst RLS variant again. Every variant reaches the optimal solution in every case except for the RLS which only manages for 61 % of the inputs. All other variants reach an optimal solution for every input. The RLS_2^B seems to be the best variant for these kinds of inputs. The next best variants are the three RLS_s^S variants which differ only slightly between each other.

4.5.2 (1+1) EA Comparison

algo type	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM
algo param	4/n	3/n	2/n	5/n	10/n	-	50/n	100/n
avg mut/change	4.012	3.096	2.292	4.945	9.887	1.580	49.788	99.729
avg mut/step	4.000	3.000	2.000	5.000	10.000	1.000	50.000	100.000
total avg count	107,309	111,459	115,595	115,597	166,608	188,301	369,943	551,189
avg eval count	$107,\!309$	$111,\!459$	$115,\!595$	$115,\!597$	166,608	188,301	369,943	$548,\!386$
max eval count	943,738	1,166,443	1,099,884	1,368,459	$1,\!579,\!593$	2,516,372	4,093,515	$4,\!552,\!116$
min eval count	182	173	22	228	107	245	365	157
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.001
avg fail dif	-	-	-	-	-	-	-	1
p-value	0.2	651 0.40	99 0.01	.37 0.00	0.02	83 0.00	0.00	000

The (1+1) EA seems to perform better with a lower mutation rate. $p_m = 4/n$ reaches an optimal solution the fastest but there is no statistically significant difference between 4/n

and 3/n and also not between 3/n and 2/n. To both sides the average speed of convergence decreases with decreasing/increasing mutation rate. For the uniform distributed input all variants of the (1+1) EA reach an optimal solution within the step limit except for 100/n which does not find an optimal solution in about 0.1 % of the inputs.

4.5.3 pmut Comparison

	algo type	pmut	pmut	pmut	pmut	$_{ m pmut}$	pmut	$_{ m pmut}$	$_{ m pmut}$	pmut
	algo param	2.00	2.25	1.75	2.75	2.50	3.00	3.25	1.50	1.25
	avg mut/change	7.943	4.039	24.963	2.220	3.079	1.905	1.708	99.853	314.277
	avg mut/step	10.095	4.562	34.759	2.274	2.931	1.934	1.729	158.122	719.683
	avg eval count	106,010	109,947	110,591	110,601	113,455	116,213	117,593	137,223	167,556
	max eval count	1,468,480	1,490,905	$1,\!104,\!274$	$1,\!588,\!457$	1,571,705	1,675,031	1,269,567	$1,\!473,\!465$	1,814,702
	min eval count	67	137	223	257	109	128	316	50	379
_	fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
_	n-value	0.2	914 0.0	441 0.0	196 0.2	822 O 4	535 0.4	001 0.0	000 0.0	000

The optimal value for β seems to be somewhere around 2.0 to 2.25 as there is no statistically significant difference between those two. The neighbouring values in both directions need more time the greater the difference to 2.0 A major increase of the runtime happens at $\beta = 1.5$ and even worse at $\beta = 1.25$, yet even the worst parameter manages to find an optimal solution in every run.

4.5.4 Comparison of the best variants

For the uniform distributed input the best variant of the RLS once again seems to perform the best. But by looking at the smaller values this does not hold in general.

fails in 10000 runs	20	50	100	500	1000	5000	10000	50000
RLS_2^B	9968	9698	8967	3362	2771	69	0	0
RLS_3^S	9733	7570	6196	4307	4242	385	22	0
RLS_4^S	8975	6341	5279	4228	4036	338	14	0
(1+1) EA (4/n)	6731	4915	4441	3969	3920	361	7	0
(1+1) EA (3/n)	7743	5714	5067	4172	4064	358	15	0
pmut (2.0)	8111	6859	6253	5004	4760	582	44	0
pmut (2.25)	8321	7214	6548	5062	4878	680	56	0

The RLS variants are the most likely to get stuck in a local optimum for $n \leq 100$. The (1+1) EA variants also often do not find an optimal solution, but this happens less frequently. The more values the input has the more likely it is for any of the algorithms to find a perfect partition. Between n = 100 and n = 500 the performance of the RLS^B drastically increases and for $n \geq 500$ this variant of the RLS stays the best variant for the remaining input sizes. Uniform distributed inputs seem to be much less likely to have a perfect partition for the small input sizes which can be explained by the coefficient of Borgs et. al. [BCP01].

avg	20	50	100	500	1000	5000	10000	50000
RLS_2^B	260	1875	6572	35798	37550	72436	75307	75126
RLS_3^S	4314	34518	36531	40001	40001	96379	107429	94308
RLS_4^S	19231	39005	39755	40011	40693	94726	105722	96892
(1+1) EA (4/n)	36909	40016	40210	41477	41432	97262	110025	108893
(1+1) EA (3/n)	36532	39907	39927	40507	40280	95037	108079	104834
pmut (2.0)	40468	40768	39431	42058	40498	106805	125448	109674
pmut (2.25)	39735	40412	38898	42199	40798	107024	126929	108445

The amount of steps needed to find an optimal solution seems to be nearly constant for every algorithm as the number of steps does not strictly increase with n but sometimes even decreases for $n \leq 1000$. This is caused by the number of steps the algorithm was given. For $n \leq 1000$ the time limit was 100,000 and for the bigger values it was $10n \ln(n)$.

Interestingly enough the average runtime decreases from n = 10000 to n = 50000 for most algorithms despite the bigger input size.

total avg	20	50	100	500	1000	5000	10000	50000
RLS_2^B	99680	97036	90348	57383	54855	74874	75307	75126
RLS_3^S	97445	84088	75856	65842	65452	109064	109219	94308
RLS_4^S	91721	77682	71558	65374	64629	105918	106864	96892
(1+1) EA (4/n)	79375	69498	66763	64704	64390	109125	110592	108893
(1+1) EA (3/n)	85675	74244	70366	65327	64550	106881	109299	104834
pmut (2.0)	88754	81395	77305	71052	68821	125374	128948	109674
pmut (2.25)	89881	83398	78907	71457	69676	128705	131376	108445

My general advice would be choosing the RLS₂^B for $n \ge 500$ and the (1+1) EA with $p_m = 4/n$ otherwise.

4.6 powerlaw distributed inputs

This distribution has mostly small values, but occasionally it also generates bigger values. The lower the parameter the higher the values get and also the amount of big values increases. For a parameter of $\beta = 2.75$ and a maximum value of 10,000 the distribution looks like in Figure 4.21. All values are rather small and less than 100, also half of the values are one. So this input seems rather easy for $\beta = 2.75$.

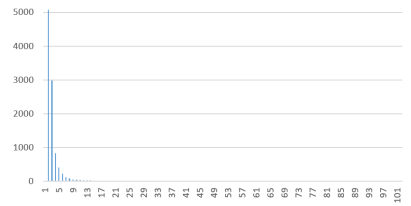


Figure 4.21: Distribution of a random powerlaw input with $\beta = 2.75$

For a value of $\beta=1.25$ the distribution looks a bit different. There are less small values close to one and instead also big values even over 1000. Figure 4.22 is cropped to get a more clear view for the smaller values. The higher values mostly occurred 0 to 2 times. The highest value 9948 occurred only once. Researching inputs like this should be more interesting which is why $\beta=1.25$ was chosen for the experiment. To give a better view on this type of input there is also a table for $\beta=2.75$ at the evaluation of the (1+1) EA. The results for the other algorithms were mostly the same, but these are not shown here for better readability. The size of the input in this case is 20,000.

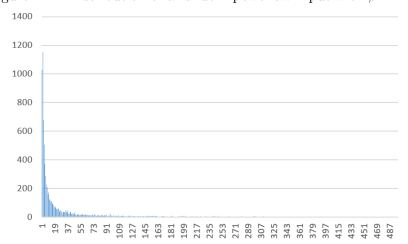


Figure 4.22: Distribution of a random powerlaw input with $\beta = 1.25$

4.6.1 RLS Comparison

algo type	RLS_s^S	RLS_b^B	RLS_s^S	RLS^B_b	RLS_b^B	RLS_s^S	RLS
algo param	s=4	b=3	s=3	b=2	b=4	s=2	-
avg mut/change	2.458	3.000	1.975	2.000	4.000	1.488	1.000
avg mut/step	2.501	3.000	2.000	2.000	4.000	1.500	1.000
avg eval count	302	330	342	359	384	417	577
max eval count	1,214	1,616	1,414	1,380	$2,\!574$	2,077	2,400
min eval count	4	3	7	10	5	15	16
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000
p-value	0.0	0.0	155 0.0	0.0	002 0.0	0.00	000

The input is even easier to solve for the RLS variants than the binomial distributed inputs. There is no clear tendency and all algorithms have a rather equal runtime. All algorithm manage to find an optimal solution in every run.

4.6.2 (1+1) EA Comparison

The first table shows the results for parameter $\beta = 2.75$

algo type	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA
algo param	50/n	100/n	10/n	5/n	4/n	3/n	2/n	-
avg mut/change	49.922	99.873	10.009	5.053	4.105	3.156	2.280	1.534
avg mut/step	49.989	100.017	9.999	4.999	4.005	3.003	2.000	0.999
avg eval count	84	103	111	157	184	208	273	461
max eval count	1,281	1,488	1,946	3,030	3,043	3,283	4,744	7,036
min eval count	0	1	3	1	0	0	2	0
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000

For $\beta=2.75$ the results are much different from $\beta=1.25$. Until $p_m \leq 50/n$ the speed of convergence increases but at $p_m=100/n$ the speed decreases again. The optimal value seems to be somewhere around $p_m=50/n$. The (1+1) EA variants are generally faster than all RLS variants when comparing the maximum number of iterations. For mutation rates $3/n \leq p_m \leq 100/n$ the (1+1) EA is also faster on average. The next table shows the results for a powerlaw distribution with $\beta=1.25$.

algo type	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM
algo param	4/n	3/n	5/n	2/n	-	10/n	50/n	100/n
avg mut/change	3.955	3.074	4.860	2.267	1.562	9.618	49.506	99.633
avg mut/step	4.003	2.999	4.999	1.999	1.000	10.001	50.000	100.000
avg eval count	284	295	311	366	640	1,190	47,165	87,514
max eval count	1,224	1,214	1,734	1,510	2,723	14,032	419,043	$1,\!123,\!496$
min eval count	15	8	15	3	0	15	13	1
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
p-value	0.0	0.0 0.0	016 0.0	0.0 0.0	0.0	0.0 0.0	0.0 0.0	000

With this setting the optimal value is shifted to somewhere around $p_m = 4/n$. The higher mutation rates perform drastically slower with $p_m = 100/n$ being 500 times slower than the optimal value. So the parameter of the distribution does change the optimal parameter for the Evolutionary Algorithm solving the input. The same was true for the RLS and also for pmut.

4.6.3 pmut Comparison

algo type	pmut								
algo param	1.50	1.75	1.25	2.00	2.25	2.50	2.75	3.00	3.25
avg mut/change	48.278	17.438	124.596	7.062	3.895	2.708	2.173	1.876	1.692
avg mut/step	99.756	26.795	370.116	9.098	4.440	2.921	2.275	1.932	1.728
avg eval count	181	193	219	225	265	301	337	366	388
max eval count	691	750	1,118	762	1,088	1,173	1,488	1,534	1,755
min eval count	8	11	5	12	13	11	11	8	13
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
n-value	0.0	000 0.0	000 0.0	000 000	000 0.0	000 0.0	000 000	000 0.0	000

The optimal value here seems to be somewhere around $\beta = 1.5$ instead of $\beta = 1.25$ for inputs from $\sim D_{20.000}^{2.75}$. So the optimal value is only lightly smaller in comparison to the (1+1) EA where the optimal value almost change from one side of the spectrum to the other.

4.6.4 Comparison of the best variants

The RLS₄^S performs equally good as the (1+1) EA variant with $p_m = 4/n$, but both are slower than $pmut_{1.5}$. Choosing any algorithm should solve the input quite fast if the parameter of the algorithm is somewhat close to the optimum.

fails in 10000 runs	20	50	100	500	1000	5000	10000	50000
RLS_4^S	7873	3943	685	0	0	0	0	0
RLS_3^B	7985	4058	706	0	0	0	0	0
(1+1) EA (4/n)	7801	3661	456	0	0	0	0	0
(1+1) EA (3/n)	7805	3719	522	0	0	0	0	0
pmut (1.5)	7800	3603	372	0	0	0	0	0
pmut (1.75)	7803	3608	371	0	0	0	0	0

The RLS is once again the algorithm that is the most likely to be stuck in a local optimum. Compared to the other algorithms it is not as drastic as for the binomial input for example. Only for n < 500 the algorithms do not find a global optimum in every run. The setting of the parameter and the choice of the algorithm almost doesn't affect the amount of runs without an optimal result. This type of input is probably easy to solve if it has a perfect partition. The two stopping conditions where a step limit and finding a perfect partition or a partition with difference of one between the two bin for uneven n. So in 78% of the runs the algorithms might have found an optimal solution, but the stopping conditions did not trigger as the found solutions were no perfect partitions.

avg	20	50	100	500	1000	5000	10000	50000
$\overline{\mathrm{RLS}_4^S}$	663	1909	1310	137	147	201	243	421
RLS_3^B	283	1158	1270	187	197	242	278	429
(1+1) EA (4/n)	613	1150	842	173	179	214	243	365
(1+1) EA (3/n)	768	1573	991	151	158	206	242	397
pmut (1.5)	890	440	185	141	145	160	170	200
pmut (1.75)	868	590	246	128	134	154	169	233

Looking at the time the algorithms needed on average the runs that hit the step limit could have possibly been no failed runs. The easiest are inputs with size n = 500. For smaller values of n the algorithms sometimes fail and even in a good run they need more iterations to find an optimal solution. Due to the increasing size of the input the algorithms need more time for the bigger values.

total avg	20	50	100	500	1000	5000	10000	50000
RLS_4^S	78871	40586	8071	137	147	201	243	421
RLS_3^B	79907	41268	8240	187	197	242	278	429
(1+1) EA (4/n)	78144	37339	5364	173	179	214	243	365
(1+1) EA (3/n)	78218	38178	6159	151	158	206	242	397
pmut (1.5)	78196	36312	3898	141	145	160	170	200
pmut (1.75)	78220	36457	3947	128	134	154	169	233

 $pmut_{1.75}$ and $pmut_{1.5}$ are not only the best variant for the bigger values of n but also for smaller inputs as well. They are the least likely to be stuck in a local optimum, and are also the fastest if they reach a global optimum.

4.7 Equivalent of linear functions for PARTITION

The input of this section is more or less equivalent to linear functions. All values except the first follow any distribution whereas the first value is the sum of all other values. The optimal solution is therefore the 100...00 or the 011...11 string. So the input is almost identical to a linear function with positive weights that is maximised/minimised depending on the value of the first bit.

For linear functions the mutation rate of 1/n was proven to be optimal for the (1+1) EA [Wit13]. This should also hold for this input. The RLS variants should also perform worse than the standard RLS. The higher the value for β the better the $pmut_{\beta}$ mutation should perform. Flips of the first bits could decrease the runtime, depending on how often they happen. By doing some testing with various algorithm variants of the RLS and the (1+1) EA it looked like the last bit was only flipped at most once for every input. There was only one case where it was flipped twice, but it was never flipped more than twice per run. The average number of flips was also mostly closer to zero than to one.

The experiments where conducted with the variant where the smaller values are all one. So for every run of each algorithm the input was $[n-1,1,1,\ldots,1,1]$. An input like this takes less time for every algorithm, but the results are mostly the same.

4.7.1	RLS	Com	parison

algo type	RLS	RLS_s^S	RLS_s^S	RLS_s^S	RLS_b^B	RLS_b^B	RLS_b^B
algo param	-	s=2	s=3	s=4	b=3	b=2	b=4
avg mut/change	1.000	1.181	1.688	1.865	3.000	1.998	3.998
avg mut/step	1.000	1.500	2.000	2.500	3.000	2.000	4.000
total avg count	91,068	168,429	235,016	309,492	921,030	921,030	921,030
avg eval count	91,068	$168,\!429$	$235,\!016$	$309,\!492$	-	-	-
max eval count	175,757	$357,\!695$	545,716	793,900	-	-	-
min eval count	$62,\!400$	110,193	$153,\!263$	$197,\!812$	-	-	-
fail ratio	0.000	0.000	0.000	0.000	1.000	1.000	1.000
avg fail dif	-	-	-	-	36	53	263
p-value	0.0	0.0	0.0	0.0	000 1.0	000 1.0	000

As expected the standard RLS reaches an optimal solution the fastest. It also reaches an optimal value for every instance. The RLS_s^S variants need more iterations to find an optimal solution. By looking at the average values more closely it seems like the average number of steps for the ${\rm RLS}_s^S$ is roughly $25,000+70,000s\pm5,000$. The standard RLS is equivalent to RLS_k^S or RLS_k^B with k=1. So the value of k=1 seems to be optimal for the RLS variants too. The RLS_b^B variants on the other hand do not reach any of the two optimal solutions in any run. This is most likely caused by their very low possibility of flipping only one bit in a single step. They would eventually reach the optimal solution as well, but this would take much longer than for the RLS. The probability of flipping only one bit in a step is $\mathcal{O}(n^{1-k})$ which results in a single bit flip every $\mathcal{O}(n^{k-1})$ steps in expectation. Because the fitness can only improve for OneMax making steps flipping more bits does not harm the fitness. The bound for OneMax is $\mathcal{O}(n \log n)$ and with the previous result the expected number of steps is bounded by $\mathcal{O}(n \cdot \mathcal{O}(n^{k-1}) \cdot \log(n \cdot \mathcal{O}(n^{k-1}))) = \mathcal{O}(n^{k-1+1} \cdot (k-1+1) \cdot \log(n)) = \mathcal{O}(kn^k \cdot \log(n))$ This problem is not equivalent to OneMax, as a flip of the bit with the highest value inverts the fitness function to ZeroMax but the result might still hold as the bound for the standard RLS for this input is the same as for the RLS on OneMax. There is no statistically significant difference between the RLS_b^B variants because all variants reached the step limit in each run. Only the amount of steps was used to for the significance test it looks like all variants are the same. If the test was executed with the differences to the optimum there would be a statistically significant difference between each algorithm.

4.7.2 (1+1) EA Comparison

algo type	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM
algo param	-	2/n	3/n	4/n	5/n	10/n
avg mut/change	1.272	1.751	2.334	2.967	3.638	7.874
avg mut/step	1.000	2.000	3.000	4.000	5.000	10.000
total avg count	231,317	292,626	494,410	867,801	921,030	921,030
avg eval count	$231,\!317$	$292,\!626$	$493,\!983$	$815,\!543$	-	-
max eval count	$522,\!203$	609,129	$913,\!330$	920,992	-	-
min eval count	$153,\!161$	189,643	287,830	$576,\!429$	-	-
fail ratio	0.000	0.000	0.001	0.495	1.000	1.000
avg fail dif	-	-	1	1	17	569
p-value	0.0	0.00	0.0	0.00	000 1.0	000

For this input the same as for OneMax holds. The static mutation rate $p_m = 1/n$ is the optimal value and the performance of the (1+1) EA decreases with rising mutation rate. Only for $p_m \leq 2/n$ the (1+1) EA managed to find one of the two optimal solutions in $10 \cdot n \ln(n)$ steps every time. With mutation rate $p_m = 4/n$ the (1+1) EA only managed

to find the optimal solution in about 50 % of the inputs. The remaining mutation rates did not manage to find an optimal solution in any of the runs. Another interesting fact is the average number of bits flipped in a successful step. For the other inputs the overall average number of bits flipped in any step was mostly the same as for the average value of the successful steps. Here this is not the case. All mutation rates flipped fewer bits in the successful steps than in the average step. The only exception is the standard mutation rate which is caused by the steps where the algorithm would flip no bit. Those steps decrease the number of the average case but not of the successful case as those steps were skipped. The p-value between 5/n and 10/n is 1.0 for the same reason as for the RLS_b^B variants.

4.7.3 pmut Comparison

For pmut only 1798/10000 repetitions were executed as there was a clear tendency which of the algorithms performs better.

algo type	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut
algo param	3.25	3.00	2.75	2.50	2.25	2.00	1.75	1.50	1.25
avg mut/change	1.286	1.359	1.459	1.598	1.814	2.152	2.722	3.719	5.197
avg mut/step	1.729	1.935	2.270	2.905	4.359	8.479	22.233	70.692	224.570
total avg count	142,937	153,214	167,454	181,340	206,641	243,262	302,445	422,885	698,772
avg eval count	142,937	$153,\!214$	$167,\!454$	181,340	206,641	243,262	302,445	422,885	686,374
max eval count	262,484	294,880	315,836	359,104	376,693	431,924	$615,\!506$	899,296	920,942
min eval count	97,595	107,234	$113,\!518$	126,834	$142,\!210$	162,092	205,614	$286,\!108$	472,286
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.053
avg fail dif	-	-	-	-	-	-	-	-	1
p-value	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	000

The results for the *pmut* operator are pretty similar to the results for the (1+1) EA and the RLS. The parameter $\beta=3.25$ which flips the least bits on average finds the solution the fastest. Decreasing value for β lead to more time needed for finding one of the two optimums. All variants find an optimum in every run except for $\beta=1.25$ which has a much higher value for the number of flipped bits per steps. The average number of bits flipped in a successful mutation is much lower than for the other inputs especially for the lower values for β . For the binomial and geometric input the successful average was around 100 for $\beta=1.25$ but for the OneMax equivalent it was only at 5.

4.7.4 Comparison of the best variants

For this comparison neither of the algorithms failed to find one of the two optimal solutions. The following table lists the amount of iterations the algorithms needed to find an optimal solution.

avg	20	50	100	500	1000	5000	10000	50000
RLS	56	187	445	3043	6806	42015	90965	535015
(1+1) EA (1/n)	127	440	1067	7508	16891	106262	231900	1375551
pmut (3.25)	81	279	668	4658	10482	65741	143099	851945
pmut (3.5)	78	265	633	4441	9934	62256	135511	802145

The results for this experiment are as expected. The RLS performs better than the (1+1) EA because it does only single bit flips. The $pmut_{3.25}$ performs better than the standard (1+1) EA although flipping more bits on average. This is most likely cause by the few steps where pmut flips many bits which increase the average. But pmut most likely chooses to flip only one bit more often as the (1+1) EA. In a previous chapter the $\mathcal{O}(n \log n)$ bound was proven for the (1+1) EA and the RLS (Theorem 3.3). This seems to hold in practice at least for the easiest version of this input where the small values are one (see figure 4.23). The scale of this figure is $n \ln n$ which means that a value of 2 on the y-axis means the algorithm needs $2n \ln n$ steps on average to reach the optimal solution. The standard (1+1) EA performs a bit worse than the other three algorithms and approaches $en \ln n$ instead of staying close to $n \ln n$.

Figure 4.23: Runtime for the OneMax equivalent with a $n \ln(n)$ scale

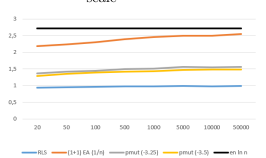
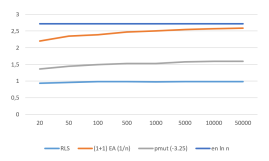


Figure 4.24: Runtime for the OneMax equivalent with uniform distribution on a $n \ln n$ scale



Another variant of this input are uniform distributed inputs for example. The small values in this case are chosen from $\sim U(1,49999)$ and the first value again is the sum of all other values. This input is harder because switching multiple small values for a big value increases the fitness but also increases the Hamming distance to the optimum. Looking at figure 4.24 the equivalent to uniform distributed linear functions looks not much harder compared to the variant where all values are one. The graphs are almost identical. It was clear for the RLS because the RLS can't switch elements which leads to the exact same behaviour. But even the other algorithms that are able to switch seem to not harm the hamming distance drastically by switching elements.

Figure 4.25: Runtime of the ${\rm RLS}_k^S$ variants for the OneMax equivalent with uniform distribution on a $n \ln n$ scale

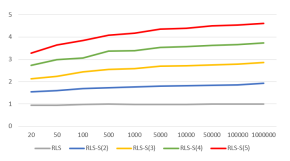


Figure 4.25 shows the average number of steps needed by the RLS_k^S for the equivalent to linear functions with uniform distribution. Lemma 3.13 proved the RLS_k^S needs time $4k + \frac{1+o(1)}{1-o(1)} \cdot kn \ln n$ but the actual expected running time seems to be lower. All RLS_k^S variants in the figure approach the value of $kn \ln n$ from below and for at least $n \leq 50,000$ do not need more time than $kn \ln n$. The RLS^B variants on the other hand have a significantly worse performance. Their average runtime is shown in Figure 4.26 with a logarithmic y-axis. A value of 2 on the y-axis here means that the algorithm needed n^2 steps on average to find an optimal solution. All RLS_k variants needs at least almost n^2 steps on average, but all variants seem to approach n^k from below (see Lemma 3.14). The actual runtime is below n^k for greater values of k and smaller input sizes which is likely a consequence of the constants. For small input sizes k! has a much higher impact but the more n grows the less noticeable the constant becomes. Figure 4.27 shows their runtime on a different scale. A value of 2 on the y-axis here means that the algorithm needed $n^2/k!$ steps on average to find an optimal solution where k is dependent on the algorithm. So all RLS_k^B variants needed about $n^k/k!$ steps on average to find an optimal solution for any input size. This is exactly the lower bound proven in Lemma 3.14.

Figure 4.26: Runtime of the RLS_k^B variants for the OneMax equivalent with uniform distribution on a n^y scale

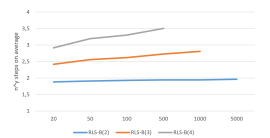
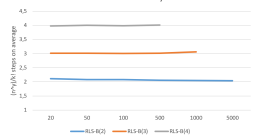


Figure 4.27: Runtime of the RLS_k^B variants for the OneMax equivalent with uniform distribution on a $n^y/k!$ scale



The bound proven for the (1+1) EA with mutation rate c/n was $4 + \frac{3+o(1)}{1-o(1)} \cdot \frac{e^c}{c} n \ln n$ (Theorem 3.3) but looking at Figure 4.28 the actual bound seems more like $\frac{e^c}{c} n \ln n$. For pmut there was no proof in this thesis but the bigger values of pmut also seem to take time $\Theta(n \ln n)$ (Figure 4.29).

Figure 4.28: Runtime of the (1+1) EA for the OneMax equivalent with uniform distribution on a $n \ln n$ scale

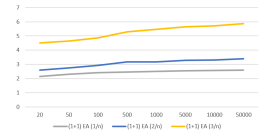
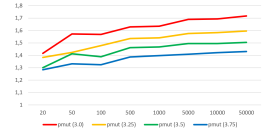


Figure 4.29: Runtime of pmut for the OneMax equivalent with uniform distribution on a $n \ln n$ scale



4.8 Carsten Witt's worst case input

This input is the worst case input from C. Witt in [Wit05] as discussed in the background section. As all experimentally researched inputs in this paper contained only integer values this input is adjusted a bit. To prevent the small values to be below zero they are instead normalised to 1. The two big values are scaled by the same factor of $((1/3 + \epsilon/2)/(n-2))^{-1}$. The higher the value for ϵ the more likely the input is to get stuck in the local optima. With increasing ϵ the local optima becomes less bad. For the small values of ϵ there were only a few cases where some algorithms did not find an optimal solution. To make this effect more visible the value of ϵ was set to $\epsilon = 0.3$.

For n=10,000 this evaluates to $w_1=w_2=5344$ and $W=9998\cdot 1+2\cdot 5344=20686$. The input then looks like this: $[5344,5344,1,1,\ldots,1,1]$. The fitness of the local optimum is $f(x)=2\cdot 5433=10688$. To leave the local optimum the algorithm therefore has to flip at least 5433+9998-10688=4654 bits as well in the same step. The best fitness is f(x)=5344+9998/2=10343, which leads to a difference of f(localOptimum)-f(opt)=345 and a approximation ratio of f(localOptimum)/f(opt)=10688/10343=1.033. This is not really close to the worst case of 4/3 any more but with this setting at least many algorithms are stuck in the local optimum at least once for the 10000 runs.

4.8.1 RLS Comparison

algo type	RLS_b^B	RLS_s^S	RLS_b^B	RLS_b^B	RLS_s^S	RLS_s^S	RLS
algo param	b=4	s=4	b=3	b=2	s=3	s=2	-
avg mut/change	3.998	2.367	3.000	1.998	1.964	1.306	1.000
avg mut/step	4.000	2.500	3.000	2.000	2.000	1.500	1.000
total avg count	4,481	4,912	6,353	11,698	15,214	32,204	102,913
avg eval count	$4,\!297$	1,510	$1,\!295$	6,945	1,794	$2,\!157$	2,211
max eval count	$77,\!441$	$11,\!617$	10,125	80,083	$12,\!221$	13,074	12,149
min eval count	0	0	0	0	0	0	0
fail ratio	0.000	0.004	0.006	0.005	0.015	0.033	0.110
avg fail dif	595	345	380	398	345	345	345
p-value	0.0	000 0.0	0.0	0.0	0.00	0.0	000

The RLS is by far most likely to get stuck in the local optimum. The general tendency is the more bits the algorithm can flip the more unlikely the local optimum becomes. The only case where this does not hold is the RLS_2^B being better than the RLS_3^S , although the RLS_3^S can also flip 3 bits. All RLS_s^B variants had runs where they neither found the global nor one of the two local optima. The algorithms were most likely tricked into the direction of the local optimum and did not manage to leave it. But they were also not fast enough to reach the local optimum because of their low probability to flip only one bit.

4.8.2 (1+1) EA Comparison

algo type	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA
algo param	100/n	50/n	10/n	5/n	4/n	3/n	2/n	-
avg mut/change	99.924	49.989	10.038	5.084	4.106	3.143	2.218	1.441
avg mut/step	100.003	50.001	10.000	4.999	4.000	3.000	2.000	1.000
total avg count	69	104	407	801	1,166	2,766	10,495	34,147
avg eval count	69	104	407	801	982	1,294	1,855	$3,\!217$
max eval count	697	$1,\!375$	5,490	$9,\!451$	11,099	$11,\!587$	13,921	$19,\!383$
min eval count	0	0	0	0	0	0	0	0
fail ratio	0.000	0.000	0.000	0.000	0.000	0.002	0.009	0.034
avg fail dif	-	-	-	-	345	345	345	345
p-value	0.0	0.0 0.0	0.0 0.0	0.0	0.0	0.0	000 0.00	000

For the (1+1) EA the result is the inversion of the results for the OneMax equivalent. The higher the mutation rate the faster the algorithm reaches a global optimum. This holds at least up to $p_m \leq 100/n$. With mutation rate $p_m \leq 4/n$ the algorithm reaches the worst case at least once in 10000 runs. If the algorithm did not manage to find an optimal solution the fitness was always the same. So there was no run where any algorithm neither found a global nor the local optimum.

algo type	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut
algo param	1.25	1.50	1.75	2.00	2.25	2.50	2.75	3.00	3.25
avg mut/change	197.409	70.534	23.050	8.724	4.351	2.777	2.111	1.770	1.563
avg mut/step	224.442	70.480	22.299	8.470	4.368	2.906	2.271	1.934	1.729
total avg count	42	87	216	503	1,094	4,063	10,961	18,727	27,644
avg eval count	42	87	216	503	910	1,488	1,676	1,814	1,909
max eval count	303	867	2,843	6,230	10,487	916,298	8 501,346	411,742	2 12,386
min eval count	0	0	0	0	0	0	0	0	0
fail ratio	0.000	0.000	0.000	0.000	0.000	0.003	0.010	0.018	0.028
avg fail dif	-	-	-	-	345	345	345	345	345
p-value	0.00	0.0	000 0.	.0000	0.0000	0.0000	0.0000	0.0034	0.0068

For pmut the result is the exact same as for the (1+1) EA. The lower β the better the performance as more bits are flipped in each step. For the OneMax input $pmut_{1.25}$ flipped 224 bits on average per step, but the average of the successful steps was only 5. Here the average of all steps is again 224, but the average of the successful steps is at 194. The heavy tail here really increases the performance as most of the high values are accepted. The algorithm is tricked into the local optima only for $\beta \leq 2.25$. If the algorithm is on the path to the local optimum it is always fast enough to reach it within the time limit.

4.8.4 Comparison of the best variants

The $pmut_{1.25}$ and the (1+1) EA with $p_m = 100/n$ perform the best and always find an optimal solution within 700 iterations and even under 100 on the average case. The RLS₄^B performs significantly worse. In the experiment with different input sizes the mutation rate of $p_m = 100/n$ is ≥ 1 for $n \leq 100$. If the algorithm flips every bit then it won't change its solution. In these cases the mutation rate was then set to $p_m = 1/2$.

fails i	in 10000 runs	20	50	100	500	1000	5000	10000	50000
	RLS_4^B	33	12	8	2	1	2	0	1
(1+1)) EA $(100/n)$	0	0	0	0	0	0	0	0
pr	nut (1.25)	0	0	0	0	0	0	0	0

Only the RLS variant had runs where it did not reach a global optimum. This happened in less than 0.4 % of the inputs for $n \leq 100$ and even in less than 0.1 % for the remaining input sizes. For the other input sizes it also managed to reach a global optimum for all inputs.

avg	20	50	100	500	1000	5000	10000	50000
RLS_4^B	12	26	46	224	442	2171	4438	21973
(1+1) EA $(100/n)$	9	16	24	34	33	47	68	232
pmut (1.25)	7	9	12	19	23	36	43	64

For the lower input sizes the RLS is slower than the remaining algorithm even it manages to find a global optimum.

total avg	20	50	100	500	1000	5000	10000	50000
RLS_4^B	342	146	126	244	452	2255	4438	22512
(1+1) EA $(100/n)$	9	16	24	34	33	47	68	232
pmut (1.25)	7	9	12	19	23	36	43	64

The $pmut_{1.25}$ is the best variant closely followed by the (1+1) EA. The RLS version is by far slower than the other to variants for the bigger input sizes. Even for the smaller inputs it is still slower.

4.9 Multiple distributions combined

This subsection covers inputs that do not follow a specific distribution. So instead of sampling from one distribution here instead the value is chosen from a set of distributions. The used distributions were $\sim U(1,49999)$, $\sim B(10000,0.1)$, $\sim Geo(0.001)$, $\sim D_{50000}^{1.25}$. Each distribution is chosen with probability 1/8 and with remaining probability 1/2 one value is drawn from every distribution and added together. An input of this distribution is shown in figure 4.30. There are a lot of values close to 0 drawn from the powerlaw (and geometric) distribution. The amount of values then decreases to around 50. From then on the amount of generated numbers rises again until 100, the expected value of the binomial distribution. After the spike caused by the binomial distribution the values look more uniform distributed until 1200 where they fall of again. The higher the numbers get the less often they occur.

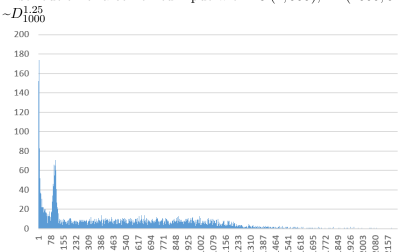


Figure 4.30: Distribution of a combined input with $\sim U(1,999)$, $\sim B(1000,0.1)$, $\sim Geo(0.01)$,

4.9.1 RLS Comparison

algo type	RLS	${\rm RLS}_s^S$	${\rm RLS}_s^S$	${\operatorname{RLS}}^S_s$	RLS_b^B	RLS^B_b	RLS^B_b
algo param	-	s=2	s=3	s=4	b=2	b=3	b=4
avg mut/change	1.000	1.431	1.831	2.214	2.000	3.000	3.999
avg mut/step	1.000	1.500	2.000	2.500	2.000	3.000	4.000
avg eval count	422	589	793	1,004	2,764	16,927	50,810
max eval count	$2,\!293$	2,934	4,879	5,722	$24,\!569$	$207,\!460$	$588,\!295$
min eval count	29	28	30	14	10	14	36
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000
p-value	0.0	0.0	0.0	0.0	0.0	0.0	000

There is a clear preference of algorithms with higher probability to flip only one bit per step similar to the geometric distribution. For geometric distributions all variants had a rather equal runtime but here the RLS_4^B needs more than 100 time the iterations on average than the fastest RLS variant. This type of input punishes the worse mutation operators more than geometric inputs but not as extreme as the equivalent to linear functions. Here still every variant reaches an optimal solution in every case as opposed to the linear function equivalent. For the geometric input only the RLS failed to reach an optimal value in every run but here it does not. This leads to the RLS being the best RLS variants for this type of input in contrast to the geometric inputs.

4.9.2	(1+1)	$\mathbf{E}\mathbf{A}$	Com	parison
-------	-------	------------------------	-----	---------

algo type	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM	EA-SM
algo param	_	2/n	3/n	4/n	5/n	10/n
avg mut/change	1.481	2.061	2.727	3.458	4.268	9.414
avg mut/step	1.000	2.000	3.001	3.999	5.000	10.000
total avg count	872	965	1,474	2,552	4,785	85,000
avg eval count	872	965	1,474	2,552	4,785	84,916
max eval count	4,201	5,151	8,431	18,891	37,928	856,656
min eval count	43	18	53	7	3	28
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000
avg fail dif	-	-	-	-	-	1
p-value	0.0	000 0.00	000 0.0	000 0.0	0.0 0.0	000

The results here are pretty similar to the results of the RLS. The lower the mutation rates the faster the algorithm reaches an optimal solution. For $p_m \leq 10/n$ did not find an optimal solution in $10n \ln n$ steps in of the 10000 runs. In other shorter experiments higher mutation rates reached an optimal solution less often. The highest researched rate of 100/n even failed to reach the optimal solution in about 15 % of the inputs.

4.9.3 pmut Comparison

algo type	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut	pmut
algo param	3.25	3.00	2.75	2.50	2.25	2.00	1.75	1.50	1.25
avg mut/change	1.583	1.737	2.002	2.423	3.303	5.830	12.519	30.910	73.182
avg mut/step	1.729	1.934	2.274	2.895	4.360	8.452	22.278	70.532	224.421
avg eval count	540	569	594	641	712	808	967	1,285	2,081
max eval count	3,110	2,891	3,504	3,896	5,152	4,274	5,610	6,190	14,984
min eval count	22	9	36	25	28	27	27	13	33
fail ratio	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
p-value	0.00	0.0	0000	0.0000	0.0000	0.0000	0.0000	0.0000	0.0000

Here the results are similar again, but there are a few differences. $\beta=2.5$ seems to perform better despite not being the highest value for β . Here using a worse mutation rates are has less impact on the runtime in contrast to the geometric inputs where the penalty is higher. Apart from that these inputs seem similar as every value of β also reaches an optimal value in every run.

4.9.4 Comparison of the best variants

The ranking of the algorithm is the same as for the other inputs with a similar preference of low mutation rates. The RLS variant has the best performance closely follow by $pmut_{3.25}$ and lastly be the standard (1+1) EA. For smaller values of n the results are similar too.

fails in 10000 runs	20	50	100	500	1000	5000	10000	50000
RLS	9947	9345	7432	308	7	0	0	0
RLS_2^S	9629	6237	1641	0	0	0	0	0
(1+1) EA (1/n)	8662	4391	1265	2	0	0	0	0
(1+1) EA (2/n)	7356	2549	659	2	0	0	0	0
pmut (3.25)	7812	3840	1122	2	0	0	0	0
pmut (3.0)	7598	3587	1070	2	0	0	0	0

The RLS variants perform the worst for at least $n \leq 100$. For this input it is also rather hard to find a perfect partition for $n \leq 50$. For the lower values there were probability multiple inputs generated with mostly small values and an uneven number of large values.

Even if the amount of large numbers is even there might still be no perfect partition, hence no algorithm is able to reach one and the run is treated as a failed run.

avg	20	50	100	500	1000	5000	10000	50000
RLS	64	149	255	407	377	389	421	555
RLS_2^S	419	1910	3534	815	589	565	595	699
(1+1) EA (1/n)	21110	16004	9721	1391	869	850	875	1046
(1+1) EA (2/n)	22856	15735	8047	1282	992	942	957	1055
pmut (3.25)	27908	16239	7953	910	542	526	546	642
pmut (3.0)	27036	16242	7978	897	576	548	573	668

The inputs get easier to solve up until n = 5000 and from then on get harder again with increasing input size. The increase for larger input sizes is much smaller than the decrease for the small values.

total avg	20	50	100	500	1000	5000	10000	50000
RLS	99470	93459	74385	3474	447	389	421	555
RLS_2^S	96305	63088	19364	815	589	565	595	699
(1+1) EA (1/n)	89444	52886	21141	1410	869	850	875	1046
(1+1) EA (2/n)	79603	37214	14107	1302	992	942	957	1055
pmut (3.25)	84226	48403	18280	930	542	526	546	642
pmut (3.0)	82474	46286	17825	917	576	548	573	668

This type of inputs are best solved by the (1+1) EA with $p_m = 2/n$ for $n \le 100$. Until n < 5000 instead using $pmut_{3.0}$ or $pmut_{3.25}$ leads to a better running time. After $n \ge 5000$ the standard RLS becomes the best option and it seems like it stays that way for the remaining input sizes.

4.10 Conclusion of empirical results

There is no clear best algorithm for every input for PARTITION and not even a best parameter for every algorithm. For inputs that are comparable to a linear function/OneMax for all base algorithms the parameter with the lowest mutation rate have the best runtime. Other instances like the worst case input of C. Witt on the other hand require much higher mutation rates for the optimal performance. Inputs generated from a powerlaw distribution showed that the optimal parameter for every algorithm is not even fixed within a specific distribution. For inputs drawn from $\sim D_{50000}^{2.75}$ the higher mutation rates reached an optimum faster than the lower mutation rate for every algorithm variant. If the input was drawn from $\sim D_{50000}^{1.25}$ then the fastest mutation rates for the (1+1) EA on $\sim D_{50000}^{2.75}$ distributed inputs then instead become the slowest.

So almost no general advice is possible, but a few points still hold for every input type. The first one is the RLS being most likely to be stuck in a local optimum especially for the

Table 4.1: Best algorithms variants for all evaluated input types

	RLS variants			(1+1)	EA var	pmut variants			
	1st	2nd	3rd	1st	2nd	3rd	1st	2nd	3rd
binomial	RLS_2^B	RLS_4^B	RLS_2^S	3/n	4/n	2/n	2.0	2.25	2.5
geometric	RLS_2^S	RLS_3^S	RLS_4^S	2/n	1/n	3/n	3.25	3.0	2.75
uniform	RLS_2^B	RLS_3^S	RLS_4^S	4/n	3/n	2/n	2.0	2.25	2.75
polwerlaw	RLS_4^S	RLS_3^B	RLS_3^S	4/n	3/n	5/n	1.5	1.75	1.25
linear function	RLS	RLS_2^S	RLS_3^S	1/n	2/n	3/n	3.5	3.25	3.0
worst case	RLS_4^B	RLS_4^S	RLS_3^B	100/n	50/n	10/n	1.25	1.5	1.75
combined	RLS	RLS_2^S	RLS_3^S	1/n	2/n	3/n	3.25	3.0	2.75

smaller input size. Even if a variant of the RLS is the fastet for the bigger input sizes it is most likely to be stuck in a local optimum for $n \le 100$ for most input types. So if the input size $n \le 100$ choosing the (1+1) EA or *pmut* mutation operator is a better choice. Another noticeable relation is that inputs that require higher mutation rates are generally easy to solve and are also solved very fast by the lower mutation rates. A lower number of iterations also does not imply a shorter runtime in every case. If the mutation rate 1/n needs only a few iterations more than 100/n it will still be much faster since one iteration is much shorter. The lower mutation rates are therefore generally a better choice as they will need less time in most cases and are still rather fast if they are not the fastest. Only if the algorithm is trapped due to its low mutation rate a higher mutation rate makes a huge difference.

Another point is that the Evolutionary Algorithms perform better for larger input sizes as there are more perfect partitions. The more perfect partition an input has, the easier it is to find one. For the lower values of n the algorithm sometimes needed 20,000 iterations on average if they managed to find a perfect partition and even longer otherwise. A runtime of $100,000 \approx 2^{14,29} \geq 2^{n-6} \geq 2^{n/2}$ is exponential in the size of the input. So for smaller values choosing other approximation algorithms or even exact algorithms will probably lead to better a better runtime. For higher values of n on easier inputs they might be efficient as well or in some cases even better.

Now to round this paper up there are two tables that summarise the previous results. For each input type and each algorithm the best three variants are listed in Table 4.1 ordered by their average runtime. This implies a general tendency of better algorithms but is not necessarily a complete insight as the best parameter and algorithm changes depending on n. Table 4.2 list my personal preference based on the previous results depending on the distribution and size of the input.

Table 4.2: Suggestions for the fastest algorithm on each input depending on the input size (the (1+1) EA is listed as EA to make the table shorter)

input size n	100	500	100	0 500	50,	000			
geometric	F	$\mathrm{EA}_{\ 2/n}$		pm	$ut_{3.25}$	RLS			
binomial	EA _{3/r}	i		RLS	zB				
uniform	$EA_{4/r}$	i		ПД	$^{\circ}2$				
powerlaw		$pmut_{1.5} \text{ or } pmut_{1.75}$							
linear function		RLS							
worst case	$pmut_{1.25}$								
combined	EA _{2/n}	$pmut_{3.25}$							

Bibliography

- [BCP01] Christian Borgs, Jennifer Chayes, and Boris Pittel. Phase transition and finite-size scaling for the integer partitioning problem. Random Structures & Algorithms, 19(3-4):247-288, 2001.
- [Dev06] Luc Devroye. Nonuniform random variate generation. *Handbooks in operations* research and management science, 13:83–121, 2006.
- [DJL23] Carola Doerr, Duri Andrea Janett, and Johannes Lengler. Tight runtime bounds for static unary unbiased evolutionary algorithms on linear functions. arXiv preprint arXiv:2302.12338, 2023.
- [DLMN17] Benjamin Doerr, Huu Phuoc Le, Régis Makhmara, and Ta Duy Nguyen. Fast genetic algorithms. In Proceedings of the genetic and evolutionary computation conference, pages 777–784, 2017.
- [Fer05] Henning Fernau. Parameterized algorithmics: A graph-theoretic approach. PhD thesis, Citeseer, 2005.
- [FGQW18] Tobias Friedrich, Andreas Göbel, Francesco Quinzan, and Markus Wagner. Evolutionary algorithms and submodular functions: Benefits of heavy-tailed mutations. arXiv preprint arXiv:1805.10902, 2018.
- [Gra66] Ronald L Graham. Bounds for certain multiprocessing anomalies. *Bell system technical journal*, 45(9):1563–1581, 1966.
- [Gun05] Christian Gunia. On the analysis of the approximation capability of simple evolutionary algorithms for scheduling problems. In *Proceedings of the 7th annual conference on Genetic and evolutionary computation*, pages 571–578, 2005.
- [Hay02] Brian Hayes. Computing science: The easiest hard problem. American Scientist, 90(2):113-117, 2002.
- [KMPS03] Hans Kellerer, Renata Mansini, Ulrich Pferschy, and Maria Grazia Speranza. An efficient fully polynomial approximation scheme for the subset-sum problem. Journal of Computer and System Sciences, 66(2):349–370, 2003.
- [Kor98] Richard E Korf. A complete anytime algorithm for number partitioning. Artificial Intelligence, 106(2):181–203, 1998.
- [Kor09] Richard Earl Korf. Multi-way number partitioning. In Twenty-First International Joint Conference on Artificial Intelligence, 2009.
- [SN12] Andrew M Sutton and Frank Neumann. A parameterized runtime analysis of simple evolutionary algorithms for makespan scheduling. In *International Conference on Parallel Problem Solving from Nature*, pages 52–61. Springer, 2012.

- [Wit05] Carsten Witt. Worst-case and average-case approximations by simple randomized search heuristics. In *Annual Symposium on Theoretical Aspects of Computer Science*, pages 44–56. Springer, 2005.
- [Wit13] Carsten Witt. Tight bounds on the optimization time of a randomized search heuristic on linear functions. *Combinatorics, Probability and Computing*, 22(2):294–318, 2013.
- [Wit14] Carsten Witt. Fitness levels with tail bounds for the analysis of randomized search heuristics. *Information Processing Letters*, 114(1-2):38–41, 2014.