Report of experiments

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1 Input properties

For various types ("mut_XMs_YMt_Z" means s and t are random identical strings of length X, and Y million respectively with mutations inserted every Z characters. "rnd_XMs_YMt" means s and t are random strings of length X, and Y million respectively) of inputs run the MS algorithm and count the number of consecutive wl() or parent() calls during the runs or ms construction phase.

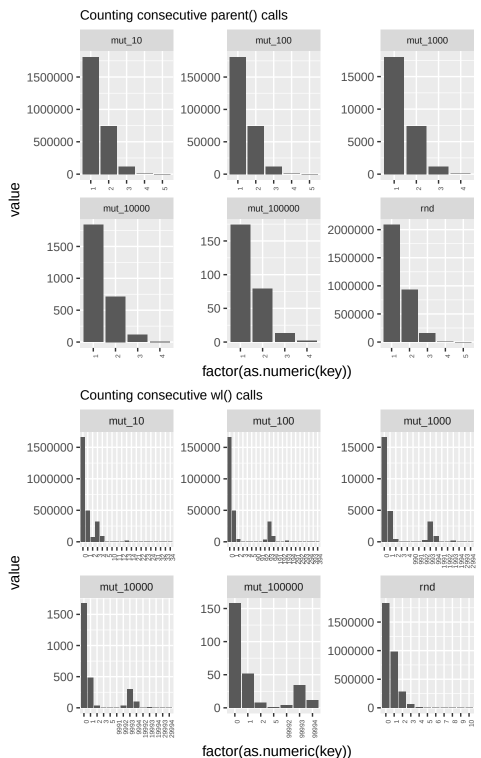


Table 1: Double rank iterations that fail for various input types.

b_path	fail	nofail	perc
rnd	690911	1818974	37.98
mut_10	1494544	2323907	64.31
mut_100	4649474	4732340	98.25
mut_1000	4965082	4973326	99.83
mut_10000	4996520	4997320	99.98
mut_100000	4999634	4999731	100.00

Table 2: Composition of the runs vector for various input types.

vector_value	mut_10	mut_100	mut_1000	mut_10000	mut_100000	rnd
0	2676093	267660	26674	2680	269	3181026
1	2323907	4732340	4973326	4997320	4999731	1818974

Table 3: Composition of the $\tt B$ vector (containing ends of maximal repeats) for various input types.

vector_value	mut_10	mut_100	mut_1000	mut_10000	mut_100000	rnd
0	36317673	36318948	36322473	36315561	36306922	26179771
1	63682328	63681053	63677528	63684440	63693079	73820230

2 Double vs. single rank

2.1 Code

The single rank and double rank implementations in sdsl: rank support v.hpp link // RANK(idx) const uint64_t* p = m_basic_block.data() + ((idx>>8)&0xFFFFFFFFFFFFFEULL); return *p + ((*(p+1)>>(63 - 9*((idx&0x1FF)>>6)))&0x1FF) +(idx&0x3F ? trait_type::word_rank(m_v->data(), idx) : 0); // DOUBLE RANK OD(i, j) $if((i>>8) == (j>>8)){$ res.first = *p + ((*(p+1))>(63 - 9*((i&0x1FF))>6)))&0x1FF) +(i&0x3F ? trait_type::word_rank(m_v->data(), i) : 0); res.second = *p + ((*(p+1))>(63 - 9*((j&0x1FF))>6)))&0x1FF) +(j&0x3F ? trait_type::word_rank(m_v->data(), j) : 0); } else { res.first = *p + ((*(p+1))>(63 - 9*((i&0x1FF)>>6)))&0x1FF) +(i&0x3F ? trait_type::word_rank(m_v->data(), i) : 0); res.second = *p + ((*(p+1))>(63 - 9*((j&0x1FF))>6)))&0x1FF) +(j&0x3F ? trait_type::word_rank(m_v->data(), j) : 0); } return res // DOUBLE RANK FC(i, j) const uint64_t* b = m_basic_block.data(); return (*pi + ((*(pi+1)>>(63 - 9*((i&0x1FF)>>6)))&0x1FF) + (i&0x3F ? trait_type::word_rank(m_v->data(), i) : 0), *pj + ((*(pj+1)>>(63 - 9*((j&0x1FF)>>6)))&0x1FF) +(j&0x3F ? trait_type::word_rank(m_v->data(), j) : 0));

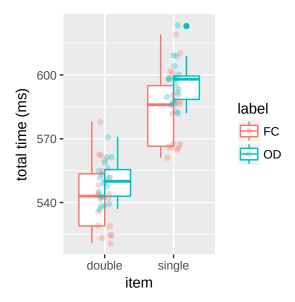
2.2 Performance

Table 4: Time (in ms) of 500K calls to wl() based on single_rank() or double_rank() methods on 100MB random DNA input; Mean/sd over 20 repetitions.

item	label	avg_time	sd_time
double	FC	543.11	15.88
double	OD	550.00	9.27
single	FC	584.32	17.11
single	OD	596.37	10.20

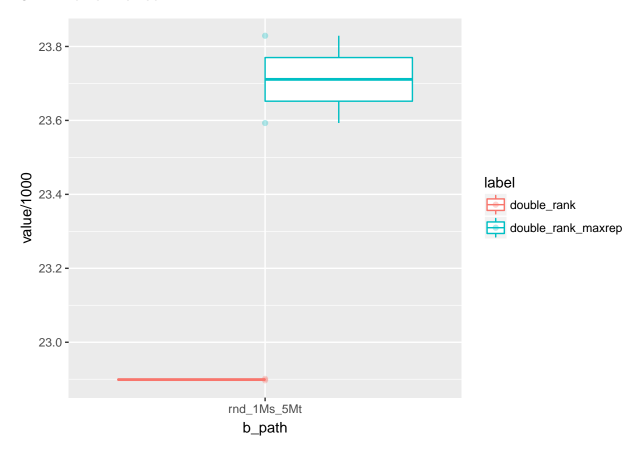
Table 5: FC vs. OD implementations. Absolute (FC / OD) and relative (100 * |FC - OD| / OD) ratios of average times

item	FC	OD	abs_ratio	rel_ratio
double	543.11	550.00	0.99	1.25
single	584.32	596.37	0.98	2.02



3 Maxrep

3.1 Performance



4 Lazy vs non-lazy

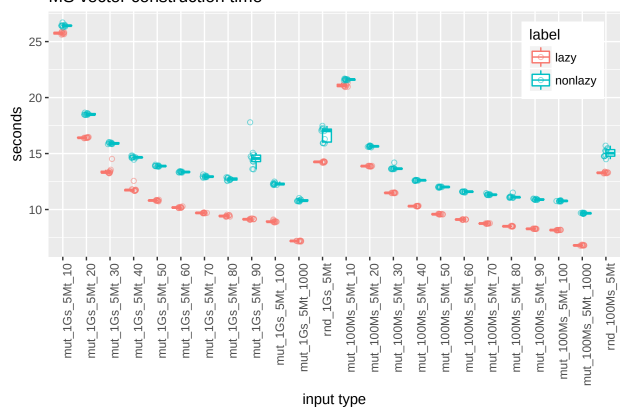
4.1 Code

The lazy and non-lazy versions differ in a couple of lines of code as follows

```
if(flags.lazy){
    for(; I.first <= I.second && h_star < ms_size; ){</pre>
        c = t[h_star];
        I = bstep_interval(st, I, c); //I.bstep(c);
        if(I.first <= I.second){</pre>
            v = st.lazy_wl(v, c);
            h_star++;
        }
    }
    if(h_star > h_star_prev) // // we must have called lazy_wl(). complete the node
        st.lazy_wl_followup(v);
} else { // non-lazy weiner links
    for(; I.first <= I.second && h_star < ms_size; ){</pre>
        c = t[h_star];
        I = bstep_interval(st, I, c); //I.bstep(c);
        if(I.first <= I.second){</pre>
            v = st.wl(v, c);
            h_star++;
        }
    }
}
```

4.2 Performance

MS vector construction time



The right panel shows the time to construct the **runs** vector. This stage is the same for both versions and is shown as a control. On the left panel it can be seen that speedup correlates positively with both the size of the indexed string and the mutation period.

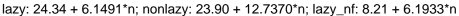
4.3 Sandbox timing

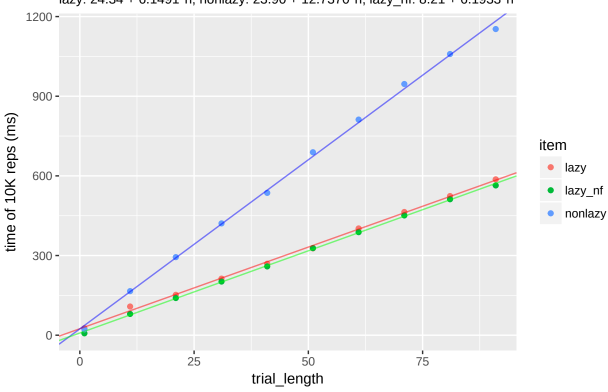
Measure the time of 10k repetitions of

- (lazy) n consecutive lazy_wl() calls followed by a lazy_wl_followup()
- (nonlazy) n consecutive wl() calls
- (lazy_nf) n consecutive lazy_wl() calls

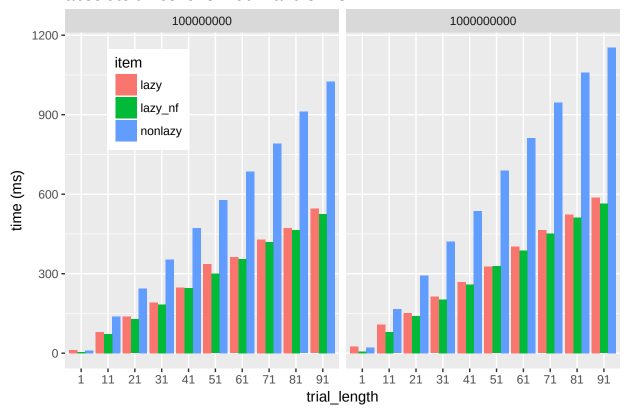
```
// lazy
for(size_type i = 0; i < trial_length; i++)
    v = st.lazy_wl(v, s_rev[k--]);
if(h_star > h_star_prev) // // we must have called lazy_wl(). complete the node
    st.lazy_wl_followup(v);
...
// non-lazy
for(size_type i = 0; i < trial_length; i++)
    v = st.wl(v, s_rev[k--]);
...
// lazy_nf
for(size_type i = 0; i < trial_length; i++)
    v = st.lazy_wl(v, s_rev[k--]);</pre>
```

indexed input size 1G





absolute times for s=100M and s=1G



4.4 Check

In the experiments above we ran the program with the "lazy" or "non-lazy" flag and measured. The total time of each experiment can be written as $t_l = l_l + a$ and $t_n = l_n + a$ for the two versions respectively; only the ts being known. Furthermore, we have \hat{l}_l and \hat{l}_n estimations – computed by combining the time / wl call with the number of with the count of wl calls in each input (Section "Input Properties"). Hence we should expect

$$\delta t = t_l - t_n = l_l + a - l_n - a = l_l - l_n \approx \delta \hat{l} = \hat{l}_l - \hat{l}_n$$

b_path	t_l	t_n	1_1	l_n	$delta_t$	$delta_l_hat$
mut_100Ms_5Mt_10	21.12	21.61	8.56	6.16	-0.49	2.39
mut_100Ms_5Mt_100	8.16	10.77	3.36	4.33	-2.60	-0.97
mut_100Ms_5Mt_1000	6.80	9.67	2.84	4.15	-2.86	-1.31
$mut_100Ms_5Mt_20$	13.87	15.64	5.66	5.14	-1.77	0.52
$mut_100Ms_5Mt_30$	11.49	13.70	4.71	4.81	-2.21	-0.10
$mut_100Ms_5Mt_40$	10.31	12.60	4.22	4.64	-2.30	-0.41
$mut_100Ms_5Mt_50$	9.58	12.01	3.93	4.53	-2.43	-0.60
$mut_100Ms_5Mt_60$	9.11	11.58	3.74	4.47	-2.48	-0.72
$mut_100Ms_5Mt_70$	8.75	11.34	3.60	4.42	-2.59	-0.81
$mut_100Ms_5Mt_80$	8.51	11.13	3.50	4.38	-2.63	-0.88
$mut_100Ms_5Mt_90$	8.28	10.90	3.42	4.35	-2.62	-0.93
$mut_1Gs_5Mt_10$	25.75	26.43	7.57	6.65	-0.68	0.92
$mut_1Gs_5Mt_100$	8.94	12.29	3.49	4.90	-3.35	-1.41

b_path	t_1	t_n	1_1	l_n	delta_t	delta_l_hat
mut_1Gs_5Mt_1000	7.19	10.82	3.08	4.72	-3.63	-1.64
$mut_1Gs_5Mt_20$	16.42	18.52	5.30	5.68	-2.10	-0.37
$mut_1Gs_5Mt_30$	13.46	15.92	4.55	5.36	-2.46	-0.81
$mut_1Gs_5Mt_40$	11.81	14.66	4.17	5.20	-2.85	-1.02
$mut_1Gs_5Mt_50$	10.81	13.89	3.95	5.10	-3.08	-1.15
$mut_1Gs_5Mt_60$	10.19	13.36	3.80	5.03	-3.17	-1.24
$mut_1Gs_5Mt_70$	9.70	12.95	3.69	4.99	-3.26	-1.30
$mut_1Gs_5Mt_80$	9.43	12.72	3.61	4.95	-3.29	-1.35
$mut_1Gs_5Mt_90$	9.14	14.74	3.55	4.93	-5.60	-1.38
rnd_100Ms_5Mt	13.29	15.07	9.65	6.55	-1.78	3.10
rnd_1Gs_5Mt	14.25	16.72	8.20	6.92	-2.48	1.28

The numbers are not identical (process dependent factors might influence the running time of function calls), but they are correlated $(corr(\delta t, \delta \hat{l}) = 0.71)$.

5 Double rank and fail

5.1 Code

```
// Given subtree_double_rank(v, i, j) -> (a.first, a.second) -- to simplify code
// DOUBLE RANK: int i, int j, char c
p = bit_path(c)
result_i, result_j = i, j;
node_type v = m_tree.root();
for (1 = 0; 1 < path_len; ++1, p >>= 1) {
 a = subtree_double_rank(v, m_tree.bv_pos(v) + result_i, m_tree.bv_pos(v) + result_j);
  if(p&1){ // left child
      if(result_i > 0) result_i = a.first;
      if(result_j > 0) result_j = a.second;
  } else { // right child
      if(result_i > 0) result_i -= a.first;
      if(result_j > 0) result_j -= a.second;
 v = m_tree.child(v, p&1); // goto child
return(result_i, result_j)
// DOUBLE RANK AND FAIL
p = bit_path(c)
result_i, result_j = i, j;
node_type v = m_tree.root();
for (1 = 0; 1 < path_len; ++1, p >>= 1) {
  a = subtree_double_rank(v, m_tree.bv_pos(v) + result_i, m_tree.bv_pos(v) + result_j);
  if(p&1){ // left child
      if(result_i > 0) result_i = a.first;
      if(result_j > 0) result_j = a.second;
  } else { // right child
      if(result_i > 0) result_i -= a.first;
      if(result_j > 0) result_j -= a.second;
  if(result_i == result_j) // Weiner Link call will fail
   return(0, 0)
  v = m_tree.child(v, p&1); // goto child
return(result_i, result_j)
```

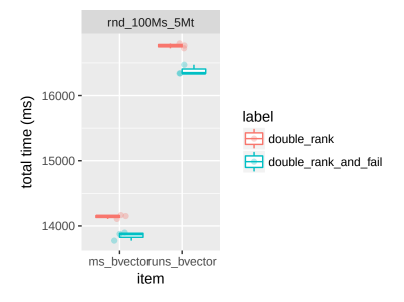
5.2 Performance

Table 7: Time (in ms) of 500K calls to wl() based on single_rank() or double_rank() methods on 100MB random DNA input; Mean/sd over 20 repetitions.

item	label	b_path	avg_time	sd_time
ms_bvector	double_rank	rnd_100Ms_5Mt	14142.00	30.27
$ms_bvector$	double_rank_and_fail	rnd_100Ms_5Mt	13850.33	66.16
$runs_bvector$	$double_rank$	rnd_100Ms_5Mt	16763.67	37.69
$runs_bvector$	$double_rank_and_fail$	rnd_100Ms_5Mt	16384.00	76.22

Table 8: Single vs. double rank. Absolute (double / single) and relative (100 * |double - single| / single) ratios of average times.

item	b_path	double_rank	double_rank_and_fail	abs_ratio	rel_ratio
ms_bvector	rnd_100Ms_5Mt	14142.00	13850.33	0.98	2.06
$runs_bvector$	rnd_100Ms_5Mt	16763.67	16384.00	0.98	2.26



6 Parallelization

6.1 Code

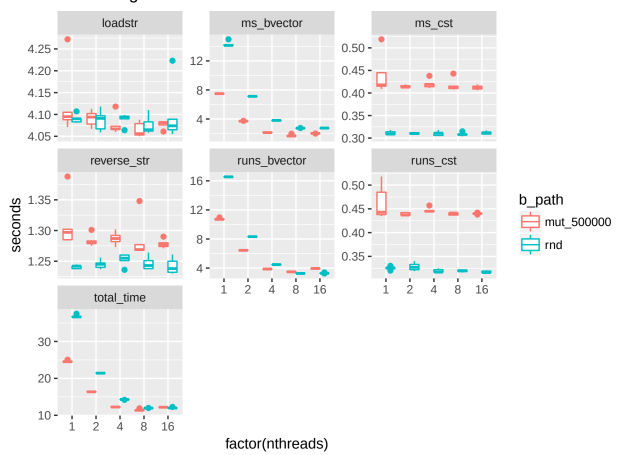
See the pseudo-code in the repo (link)

6.2 Performance

Run the MS construction program on the same input (random strings s of length 100M and t of length 5M) with varying parallelization degree (nthreads = number of threads).

The time is reported over 5 runs for each fixed number of threads.

Time usage



Space in MB for the same settings as above.

Each thread allocates its own ms vector with initial size |t|/nthreads then it resizes by a factor of 1.5 each time it needs to. Resizing will always result in a vector smaller than 2|t| elements.

