LISA

Install

do CXX=icpc at make add -I bwa to the makefile (no need for the latest commit) add -std=c++1y to line 21 in scripts/build-rmi.linear_spline.linear.sh install rust use absolute path when building the index

exact-search-lisa.o

./exact-search-lisa.o /nfshomes/yhxu/858D/data/fasta/test.fasta /nfshomes/yhxu/858D/data/query/test.query 3 3 1

./exact-search-lisa.o

/nfshomes/yhxu/scratch/858D/858D-project/data/fasta/Chlamydia.fasta /nfshomes/yhxu/scratch/858D/858D-project/data/query/Chlamydia.32.query 4 256 1

benchmarks/bench-fixed-len-e2e-match-lisa.cpp

- [62] main
- [81] read reference sequence. Printed out and it is in ACGT form and is correct
- [86] read in queries. Printed out and it is in ACGT form and concatenated by; and is correct
- [105] add \$ to the reference seq
- [124] create rmi
- src/ipbwt rmi.h
 - [226] constructor
 - [234 332] define a function to build ipbwt from scratch
 - [339] check if ipbwt already exists. If it is, load it at [350]. If not, build it from scratch at [367]. **The code will fail at this place if it doesn't exist.**
 - [234] call build_ipbwt
 - [240] build __sa
 - [265] free __sa
 - should save sa here
 - [322] sanity check, it will fail at the last entry
 - [375] check if rmi parameters exists. If it is, load it at [384]. If not, evoke scripts to train it [405 438]
- [124] With pre-built index, rmi should return correctly
- [162] create threads
- [196] go through all queries with while loop.

- The code will end up with infinite loop if query length is not multiple of k
- Fixed by making interation condition to be num_iter > 0
- The code give wrong result if query length is not multiple of k
- If query lengths are different, the code give wrong answers for all queries except the longest ones.
- [233] program failed because of float point exception at the #pragma
 - fixed, it's because parallel_batch_size = 0 when there are few queries
- [251] call rmi.backward_extend_chunk_batched(&str_enc[i], qs_sz, &intv_all[i*2]);
- src/ipbwt_rmi.h
 - [956] backward extend chunk batched
 - [982] call process_query_one_step
 - [850] process query one step
 - [923] call last_mile_vectorized_search_final_step(meta.ipb_x[0], meta.first[0], meta.m[0]);
 - [824] last mile vectorized search final step
 - [827] call _mm512_loadu_si512, causes illegal instruction error

build-index-forward-only-lisa.o

./build-index-forward-only-lisa.o /nfshomes/yhxu/858D/858D-project/data/fasta/test.fasta 3 3

./build-index-forward-only-lisa.o

/nfshomes/yhxu/858D/858D-project/data/fasta/Ecoli.fasta 3 3

Mostly same as querying. Main code is in

- src/ipbwt rmi.h [226-438]

Fixed IPBWT entry size:

src/ipbwt_rmi.h

- [60, 63, 65] Hard-code NUM_POS_BITS = 38, NUM_CHUNK_BITS = 42, NUM IPBWT BYTES = (NUM IPBWT BITS + 7) / 8 = 10
- [289] allocate n * NUM IPBWT BYTES for IPBWT

Pufferfish

pufferfish index

./src/pufferfish index -r

/Users/henryxu/Desktop/Sp2022/858D/project/data/fasta/test.fasta -o /Users/henryxu/Desktop/Sp2022/858D/project/data/pufferfish-index/test/ -k 3

Pufferfish.cpp

- [46] main
- [313] call pufferfishIndex(indexOpt)
- PufferfishIndexer.cpp
 - [355] pufferfishIndex (pufferfish::IndexOptions&
 indexOpts)
 - [457] print ntHll estimated {} distinct k-mers, setting
 filter size to
 - [499] call buildGraphMain (args) to use TwoPaCo to build the compacted dbg
 - [523] call dumpGraphMain (args) to use TwoPaCo to serialize the compacted dbg
 - [539,540] reading GFA file created by TwoPaco and then use it them to create contig table pufferfish::BinaryGFAReader
 pf(outdir.c_str(), k 1, buildEqCls, buildEdgeVec,
 iointLog)
 - PufferfishBinaryGFAReader.cpp
 - [84] constructor
 - [96] read in the contig sequence created by TwoPaCo. Here we can print out the contig
 - [101] read in the bv created by TwoPaCo. Here we can print out the bv

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pufferfish kquery

pufferfish kquery -i <index> -q <ref>... [-p <threads>] [-v]

./src/pufferfish kquery -i

/Users/henryxu/Desktop/Sp2022/858D/project/data/pufferfish-index/test/ -q /Users/henryxu/Desktop/Sp2022/858D/project/data/query/test.query

Pufferfish.cpp

- [46] main
- [313] call pufferfishKmerQuery(kmerQueryOpt)
- PufferfishKmerQuery.cpp
 - [112] pufferfishKmerQuery
 - [150] load pufferfishIndex
 - pufferfishIndex.cpp
 - [115] load contig sequence "seq "
 - [156] call doPufferfishKmerQuery(pi, parser, iomut)
 - [25] doPufferfishKmerQuery
 - [65] call auto phits = pi.getRefPos(km, qc);
 - PufferfishIndex.cpp
 - [158] PufferfishIndex::getRefPos(CanonicalKmer& mer, pufferfish::util::QueryCache& qc)
 - [163] look up MPH table
 - [164] check this kmer exists
 - [165] look up pos vector to get the position in contig sequence 'seq_'
 - This can be replaced by querying LISA FM-Index