

# 1 KrakenHLL: Confident and fast metagenomics classification using 2 unique k-mer counts

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8

## 9 **Abstract**

10 *Motivation:* False positive identifications are a significant problem in metagenomics. Spurious  
11 identifications can attract many reads that often aggregate in the genomes. Genome coverage  
12 may be used to filter false positives, but fast k-mer based metagenomic classifiers only provide  
13 read counts as metrics, and re-alignment is expensive. We propose using k-mer coverage, which  
14 can be computed during classification, as proxy for genome base coverage.

15 *Results:* We present KrakenHLL, a metagenomics classifier that records the number of unique k-  
16 mers as well as coverage for each taxon. KrakenHLL is based on the ultra-fast classification  
17 engine Kraken and combines it with HyperLogLog cardinality estimators. We demonstrate that  
18 more false-positive identifications can be filtered using the unique k-mer count, especially when  
19 looking at species of low abundance. Further enhancements include mapping against multiple  
20 databases, plasmid and strain identification using an extended taxonomy, and inclusion of over  
21 100,000 additional viral strain sequences. KrakenHLL runs as fast as Kraken, and sometimes  
22 faster.

23 *Availability and Implementation:* KrakenHLL is implemented in C++ and Perl, and available  
24 under the GPL v3 license at <https://github.com/fbreitwieser/krakenhll>.

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## 26 **Introduction**

27 Metagenomic classifiers attempt to assign taxon identifiers to each read in a sample. Typically,  
28 this is done using mapping rather than alignment, which returns the read classifications but not  
29 the aligned positions in the genomes (as reviewed by Breitwieser, et al., 2017). However, read  
30 counts can be deceiving. Sequence contamination of the samples - introduced from laboratory  
31 kits or the environment during sample extraction, handling or sequencing - can yield high  
32 numbers of spurious identifications (Salter, et al., 2014; Thoendel, et al., 2017). Having only  
33 small amounts of input material can further compound the problem of contamination. In clinical  
34 diagnosis of infectious diseases, for example, often less than 0.1% of the DNA sequenced is from  
35 microbes of interest (Brown, et al., 2018; Salzberg, et al., 2016). Furthermore, spurious matches  
36 can result from low-complexity regions of genomes, and contamination in the database genomes  
37 themselves (Mukherjee, et al., 2015).

38

39 Such false positive reads typically match only small portions of the genome. Reads from  
40 microbes that are truly present should distribute relatively uniformly across the genome rather  
41 than be concentrated in one or a few locations. Genome alignment can reveal this information.  
42 However, it is resource intensive, requires the selection of specific genomes, and it is difficult to  
43 extrapolate from the alignment of one genome to higher levels in the taxonomic tree. Some  
44 metagenomics methods use coverage information for better mapping or quantification, but  
45 usually require results from much slower alignment methods as input (Dadi, et al., 2017).

46 Notably, assembly-based methods also work, but only for highly abundant species (Quince, et  
47 al., 2017).

48

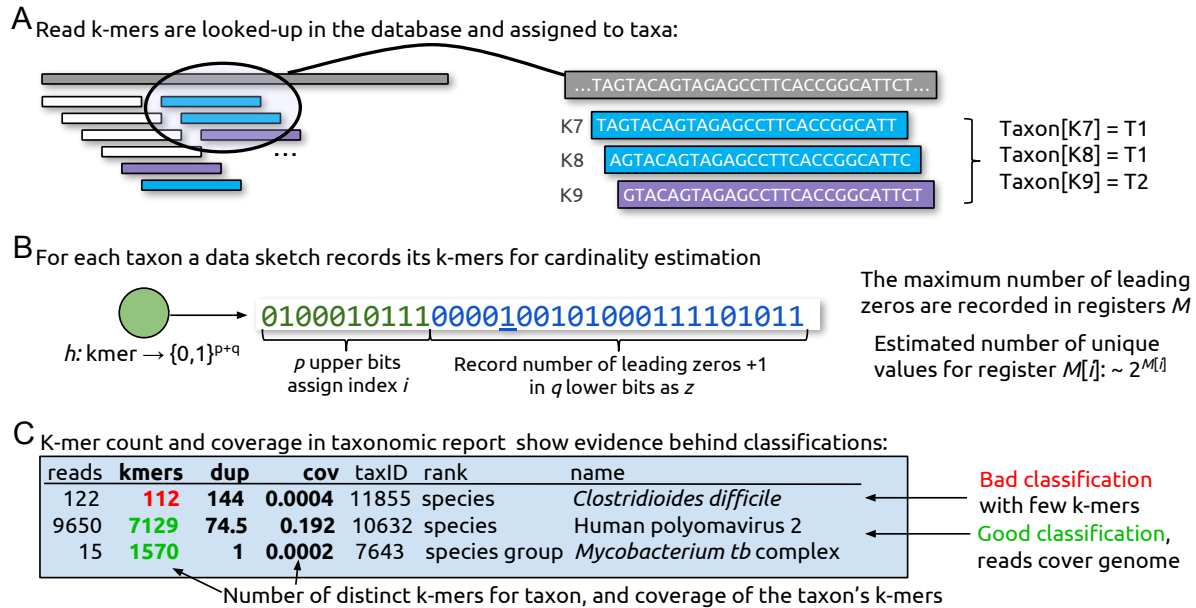
49 Here, we present KrakenHLL, a novel method that combines fast k-mer based classification with  
50 fast k-mer cardinality estimation. KrakenHLL is based on the Kraken metagenomics classifier  
51 (Wood and Salzberg, 2014) and implements fast counting of the number of unique k-mers  
52 identified for each taxon using the efficient probabilistic cardinality estimation algorithm  
53 HyperLogLog (Ertl, 2017; Flajolet, et al., 2007; Heule, et al., 2013). The count and percentage of  
54 the taxon's unique k-mers in the database that are covered by read k-mers can be used to discern  
55 false positive from true-positive sequences. Furthermore, KrakenHLL implements other new  
56 features for better metagenomics classifications: (a) searches can be done against multiple  
57 databases hierarchically, (b) the taxonomy can be extended to include nodes for strains and  
58 plasmids, thus enabling their detection, and (c) database build script enables adding over 100  
59 thousand viral strains from the NCBI Viral Genome Resource (Brister, et al., 2015). Notably,  
60 KrakenHLL, which provides a superset of the information of Kraken, is as fast or faster than  
61 Kraken while using very little additional memory during classification.

## 62 **Results**

63 KrakenHLL was developed to provide efficient k-mer coverage information for all taxa  
64 identified in a metagenomics experiment. The main workflow is as follows: As reads are  
65 processed, each k-mer is assigned a taxa from the database (Figure 1 (A)). KrakenHLL  
66 instantiates a HyperLogLog data sketch for each taxon, and adds the k-mers to it (Figure 1 (B)).  
67 After classification, KrakenHLL traverses up the taxonomic tree and merges the estimators of the  
68 child taxa to the parent. KrakenHLL reports the number of unique k-mers, and the breadth and

69 depth of k-mer coverage for each taxon in the taxonomic tree in the classification report (Figure  
 70 1 (C)).

71



72

73 Figure 1: KrakenHLL algorithm and report. (A) The taxon mappings for each k-mer of a read are  
 74 queried from the database. For each taxon, a unique k-mer counter is instantiated, and the  
 75 observed k-mers are added to it. (B) Unique k-mer counting is implemented with the  
 76 probabilistic estimation method HyperLogLog (HLL) with below 1% error in 16KB of memory  
 77 per counter. (C) The number of unique k-mers, duplicity (average time each k-mer has been  
 78 seen) and coverage are reported for each taxon in the taxonomic tree, enabling assessment of the  
 79 classification.

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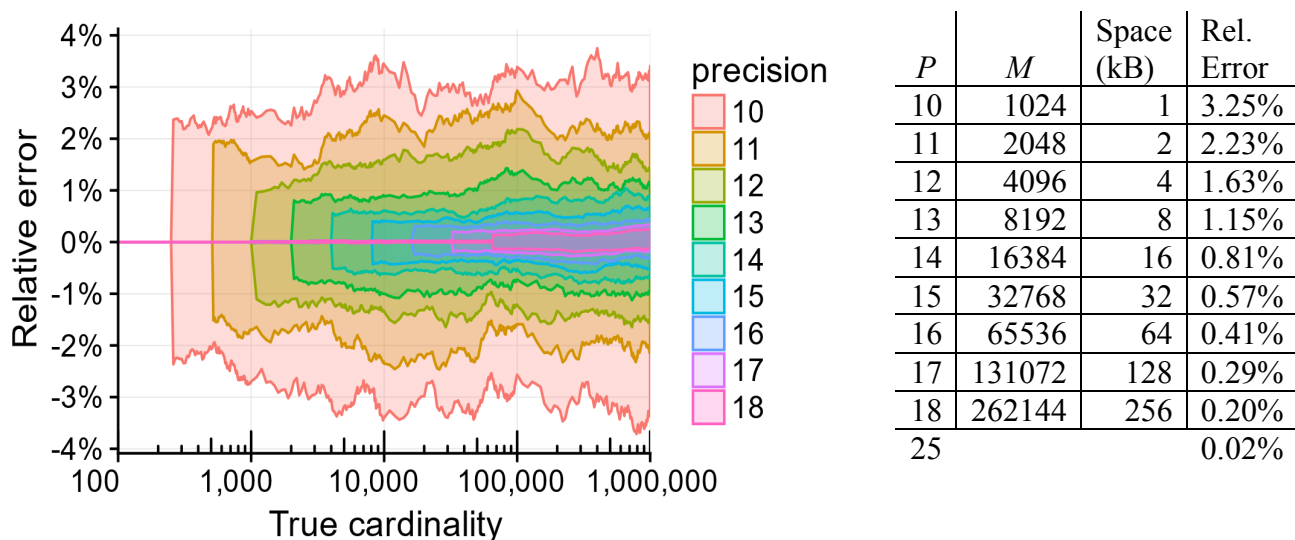
82 **Efficient k-mer cardinality estimation with HyperLogLog algorithm**

83 Exact counting of the number of unique values (cardinality) in the presence of duplicates  
84 requires memory proportional to the cardinality. Very accurate *estimation* of the cardinality,  
85 however, can be achieved using only a small amount of fixed space. The HyperLogLog  
86 algorithm (HLL), originally described by (Flajolet, et al., 2007), is currently one of the most  
87 efficient cardinality estimators, and lends itself to k-mer counting (Irber Junior and Brown,  
88 2016). The main idea behind the method is that long runs of leading zeros are unlikely in random  
89 hashes. E. g., it's expected to see every fourth hash start with one 0-bit before the first 1-bit  
90 ( $01_2$ ), and every  $32^{\text{nd}}$  hash starts with  $00001_2$ . The algorithm saves a sketch of observed data  
91 based on hashes of the k-mers in  $2^p$  one byte registers (in our implementation), where  $p$  is the  
92 precision parameter. The relative error of the estimate is  $1/\sqrt{2^p}$ . With  $p=14$ , the sketch uses  
93  $2^{14}$  one-byte registers, i.e. 16KB of space and has a relative error less than 1% (Figure 2).  
94  
95 Generating the sketch: Each k-mer is first hashed into a 64-bit string  $H$ . The sketch starts out in  
96 sparse representation which has an effective  $p$  of 25, using 4 bytes per element. See (Heule, et  
97 al., 2013) for more details on the encoding. Once  $m/4$  distinct elements have been observed, we  
98 switch to the standard representation of (Flajolet, et al., 2007): The first  $p$  bits of  $H$  are used as  
99 index  $i$  into the registers  $M$ . The later  $64-p=q$  bits are used to define the rank based on the  
100 position of the first 1-bit (or, equivalently, the count of leading zeros plus one). If all  $q$  bits are  
101 zero, the rank is  $q+1$ . The register  $M[i]$  is updated if the rank is higher than the current value of  
102  $M[i]$ .  
103 When the read classification is finished, KrakenHLL aggregates the taxon sketches up the  
104 taxonomy tree. Each taxon's sketch is merged with its children's sketches. The cardinality  
105 estimate is computed using a recently reported improved method (Ertl, 2017) that does not

106 require empirically determined thresholds to account for biases and switching between linear  
 107 counting and HLL estimator (Supplementary Figures 1 and 2). Figure 2 shows the performance  
 108 and memory usage of KrakenHLL's cardinality estimator for up to one million k-mers. Suppl.  
 109 Methods Section 1 contains a more in-depth description of the algorithm and implementation.

110

111



112 Figure 2: Cardinality estimation on randomly sampled microbial k-mers using HyperLogLog.

113 (Left) Standard deviations on the relative errors of the estimate with precision  $p$  ranging from 10

114 to 18. As expected, higher values of  $p$  give lower relative error, and no systematic bias is

115 apparent. Up to cardinalities of about  $2^p/4$  the relative error is near zero, and at higher

116 cardinalities the error boundaries stay constant. (Right) The size of the registers, space

117 requirement, and expected relative error for HyperLogLog cardinality estimates with different

118 values of  $p$ . For example, with a precision  $p=14$ , the expected relative error is less than 1%. The

119 counter only requires 16 kB of space, which is three orders of magnitude less than that of an

120 exact counter (at a cardinality of a million). Up to cardinalities of  $2^p/4$ , a sparse representation of

121 the counter is used with a higher precision of 25 and an effective relative error rate of about  
122 0.02%.

123

## 124 **Results on simulated and biological data**

125 Simulated test datasets are invaluable in assessing the performance of bioinformatics algorithms.

126 Read simulators can create arbitrarily complex artificial communities and we know the source of  
127 every read. However, simulated datasets do not necessarily represent biological data.

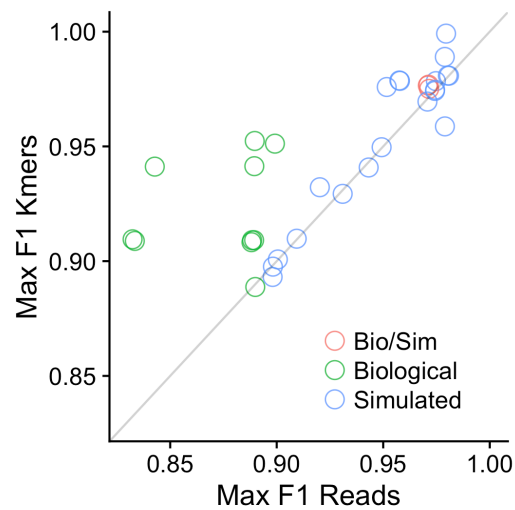
128 Specifically, laboratory and environmental contamination, a main reason behind false  
129 identifications in metagenomics samples (Salter, et al., 2014), are hard to model. Biological test  
130 datasets that are generated by mixing bacterial isolates at known quantities, on the other hand,  
131 usually have very few species and limited complexity.

132

133 (McIntyre, et al., 2017) recently reviewed eleven metagenomics classifiers and compiled a list of  
134 simulated and biological test datasets from 16 distinct sources (McIntyre-Mason, Suppl. Table  
135 2). Eleven of these datasets were from biological mock communities. The largest biological  
136 datasets consist of 23 species that were mixed at even proportions (Human Microbiome Project  
137 mock communities, sequenced with Illumina and 454 machines). We tested KrakenHLL on ten  
138 biological and 21 synthetic datasets to see if better separation of false positives and true positives  
139 can be achieved using unique k-mer counts instead of read counts (see Suppl. Table 3). Our main  
140 measure for comparison is the maximum F1 score, defined as  $2 * \text{precision} * \text{recall} / (\text{precision} +$   
141  $\text{recall})$ .

142

143 Unique k-mer count thresholds worked very well in biological datasets, performing better than  
144 the read count threshold in nine out of ten datasets, with a tie in one (Figure 3 and Suppl. Table  
145 3). On average, the maximum F1 was 0.05 higher when using k-mer instead of read thresholds,  
146 improving from 0.87 to 0.92. As expected, the difference was not as clear in simulated datasets,  
147 even though the k-mer count still performed better than the read count. In eight out of the 21  
148 datasets, both metrics performed equally well, as the datasets were easily separated into true and  
149 false identifications. In eight datasets k-mer count achieved better F1 scores, and in five read  
150 count achieved better F1 scores. The average F1 with k-mer count was slightly higher with 0.945  
151 against 0.940. This difference in difference in performance is likely due to simulated datasets  
152 lacking some features of biological data.

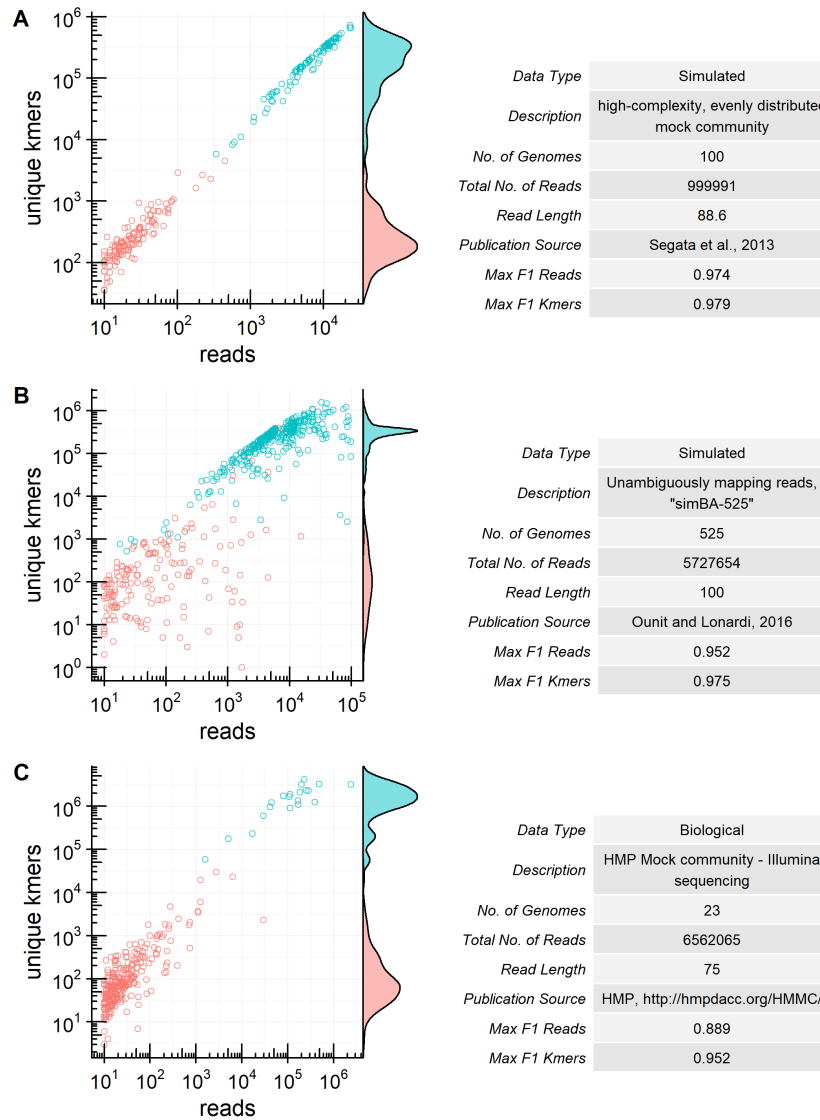


153  
154 Figure 3: Using unique k-mers as thresholds instead of reads can give higher F1 scores. Each dot  
155 is a dataset described in McIntyre, et al. Excludes (for visual purposes) dataset LC5 of Segata et  
156 al. with a F1 read score of 0.73 and F1 Kmer score of 0.75.

157  
158 Figure 4 shows the results on two simulated and one biological datasets. In simple simulated and  
159 biological datasets, the true species often separate nearly perfectly using either a read count or a



160 unique k-mer count threshold (Figure 3 (A)). In more complex datasets, however, read count  
 161 thresholds often contain more false species than k-mer thresholds (Figure 3 (B) and (C)).  
 162



163  
 164 Figure 3: Unique k-mer counts separate true identifications better from false ones. The plot  
 165 shows the number of reads vs the number of unique k-mers in two simulated and one biological  
 166 test datasets. Each point is a species identification. Blue dots are the ‘true’ species, red dots are  
 167 false positive or background identifications. (A) Relatively easy case of a simulated test dataset  
 168 with 100 genomes. The true and false positives separate nearly perfectly with either read or k-

169 mer count. (B) Largest simulated test dataset shows better separation by unique k-mer counts.  
170 (C) Largest biological dataset also separates well between true and false positives.

171

172

### 173 **Results on biological samples for infectious disease diagnosis**

174 Metagenomics is increasingly used to find species of low abundance. A special case is the  
175 emerging use of metagenomics for the diagnosis of infectious diseases (Simner, et al., 2017;  
176 Zhang, et al., 2015). Host tissue or body fluids are used to find the likely culprit of a disease.  
177 Usually, most (often 95% and more) of the reads match to the host, and maybe 10 to 100 out of  
178 the millions of reads are matched to the target species. Skin bacteria from the patient, physician  
179 or lab personal and other contamination from sample collection or preparation can easily  
180 accumulate a similar number of reads, and thus cloud the detection of the pathogen.

181

182 To assess if the unique k-mer count metric can be used to rank and identify pathogen  
183 identification, we reanalyzed ten patient samples (Salzberg, et al., 2016). (See Supplementary  
184 Methods for details on the database, which also contains over 100 thousand viral strain  
185 sequences.). (Salzberg, et al., 2016) sequenced spinal cord mass and brain biopsies from ten  
186 patients in the intensive care unit, for whom routine tests for pathogens returned inconclusive. In  
187 three out of the ten cases, a likely diagnosis could be made with the help of metagenomics, and  
188 in a fourth case, a diagnosis could be made with an updated database. For confirmation of  
189 metagenomics class, the authors re-aligned pathogen reads to individual genomes.

190

191 Table 1 shows the results of our reanalysis for the confirmed identifications in the four patients,  
192 including the number of reads and unique k-mers of the pathogen, as well as the number of  
193 covered bases of a re-alignment. Even though the read numbers are low in some cases, the  
194 number of unique k-mers suggests that they are distributed across the genome. For example, in  
195 PT8, 15 reads are matching 1570 k-mers, and re-alignment shows 2201 covered base pairs. In  
196 contrast, Table 2 shows examples of identifications in the same dataset that are not well  
197 supported by a high unique k-mer count.

198

199 Table 1: Pathogen identifications in patients with suspected neurological infections. The  
200 pathogens were identified with as little as 15 reads, but those mapped to a high number of unique  
201 k-mers, indicating random distribution of the reads on the genome. “Bases” are the number of  
202 covered bases in the re-alignment of a selected genome. Interestingly, the k-mer count in PT5  
203 reveals that there seems to be more than one viral strain present, as the k-mers cover more than  
204 one genome.

Sample	Name	Reads	K-mers	Bases
PT5	Human polyomavirus 2	9650	7129*	5130
PT7	<i>Elizabethkingia genomosp. 3</i>	403	20724	52921
PT8	<i>Mycobacterium tuberculosis</i>	15	1570	2201
PT10	Human gammaherpesvirus 4	20	2084	2780

205

206 Table 2: Dubious identifications have few k-mers. Note that the viral identifications in PT4 and  
207 PT10 stem from non-RefSeq viral genomes from the NCBI Viral Genome Resource. Since the  
208 KrakenHLL reports sequence-level matches, the source genomes are easy to find.

Sample	Name	Reads	K-mers
PT3	<i>Clostridioides difficile</i>	122	126
PT4	Hepatitis C virus JF343788.1 Recombinant Hepatitis C virus	101	3
PT5	<i>Akkermansia muciniphila</i>	936	136
PT10	Human betaherpesvirus 5 JN379815.1 UNVERIFIED: Human herpesvirus 5 strain U04, partial genome	63	5

209

## 210 **Storing strain genomes with assembly project and sequence accessions**

211 Kraken stores a NCBI taxonomic identifier for each k-mer in its database. This strategy worked  
212 well when new taxonomy IDs were assigned to each new microbial strain in GenBank. However,  
213 in 2014 the NCBI Taxonomy project stopped giving new IDs to microbial strains – only novel  
214 species get new taxonomy IDs (Federhen, et al., 2014). New strains, therefore, have the  
215 taxonomy ID of the species, or the taxonomy ID of a strain that was added before 2014.  
216 Microbes that have been intensively surveyed, such as *Escherichia coli* or *Salmonella spp.*, have  
217 up to hundreds of genomes indexed with the same taxonomy ID, and are thus indistinguishable  
218 by Kraken. The new way of identifying microbial strains is to use the Bioproject, Biosample and  
219 Assembly accession codes (Breitwieser, et al., 2017). KrakenHLL thus adds new nodes to the  
220 taxonomy tree as children of the assigned taxon. A taxonomic node may also be added for each  
221 sequence – e.g. specific bacterial chromosomes or plasmids. Those new nodes in the taxonomy  
222 tree are given taxonomy IDs starting at 1,000,000,000. Having these extended nodes can help  
223 identify specific strains as well as bad database sequences (see Table 2 and Suppl. Table 3).

224

## 225 **Hierarchical read classification with multiple databases**

226 KrakenHLL allows using multiple databases hierarchically in order of confidence. In the  
227 following example each k-mer is matched first against the HOST, then the PROK, then the  
228 EUK\_DRAFT database.

229  
230 `krakenhll --db HOST --db PROK --db EUK_DRAFT`

231  
232 Note that all database need to share the same taxonomy database. If taxIDs are added for  
233 genomes or sequences, then it is necessary that the databases are consecutively constructed with  
234 the same taxonomy database.

235

### 236 **Timing and memory requirements**

237 The additional features of KrakenHLL come without a runtime penalty. In fact, due to code  
238 improvements, KrakenHLL can run faster than Kraken especially when most of the reads are  
239 from one species (See Suppl. Table 2 for timings on patient data, Suppl. Table 3 for timings on  
240 the test datasets). On the patient data, the processing speed (base-pairs per minute) was on  
241 average 57% higher with KrakenHLL compared to Kraken, while it was 8% higher. Overall wall  
242 clock time was slower, too, when comparing the runtime of both kraken and kraken-report with  
243 krakenhll (which generates the report with the classification binary). The average additional  
244 memory requirements were less than 1GB. On the patient datasets, the average maximum  
245 memory usage went from 118 to 118.35GB, and for the test datasets, the usage went up from  
246 46.28 to 46.99GB.

## 247 Conclusions

248 We present a novel method that combines fast k-mer based classification with efficient  
249 cardinality estimation. We demonstrated that unique k-mer counts can help discard false  
250 identifications in real samples. When the reads from a species yield many unique k-mers, we are  
251 more confident that the taxon is truly present, while a low number of unique k-mers suggests a  
252 possible false positive identification. It is important to note that choice of the appropriate  
253 threshold will depend on the application. For example, in infectious disease diagnosis, unique k-  
254 mers can be used for ranking of the identifications. Conversely, in microbial ecology, a global  
255 threshold on the number of unique k-mers can be applied at any desired taxonomic rank. We  
256 believe that the ability to summarize to higher levels of the tree is a great advantage of the k-mer  
257 count over using covered bases in a genome alignment. In summary, KrakenHLL gives more  
258 confident identifications by reporting the unique k-mer count and coverage, without any runtime  
259 penalty.

260

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265

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269

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