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ITHAKA: A TAXONOMIC CLASSIFIER BASED ON BLOOM FILTERS, AND SPACED SEEDS



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METAGENOMICS & NGS

Metagenomics is a powerful approach to study genetic material contained in environmental samples, which is revolutionized by high-throughput sequencing technologies. Taxonomic classification of metagenomics data sets is a common step in analysis, a step for which computational cost becomes prohibitive with the growth of metagenomic datasets.

Approaches using sequence alignment algorithms, often based on the Burrows-Wheeler transform, such as Kaiju[1], Centrifuge[2], compete successfully with k-mer based alignment-free comparison methods such as Kraken[3], Clark[4]. Improvements to k-mer based approaches include: extending contiguous k-mers with spaced seeds [5][4], using Bloom filters as an underlying data structure for storing k-mers.[6]

SPACED SEEDS

A **spaced seed** is a pattern over alphabet $A = \{\#, -\}$, where

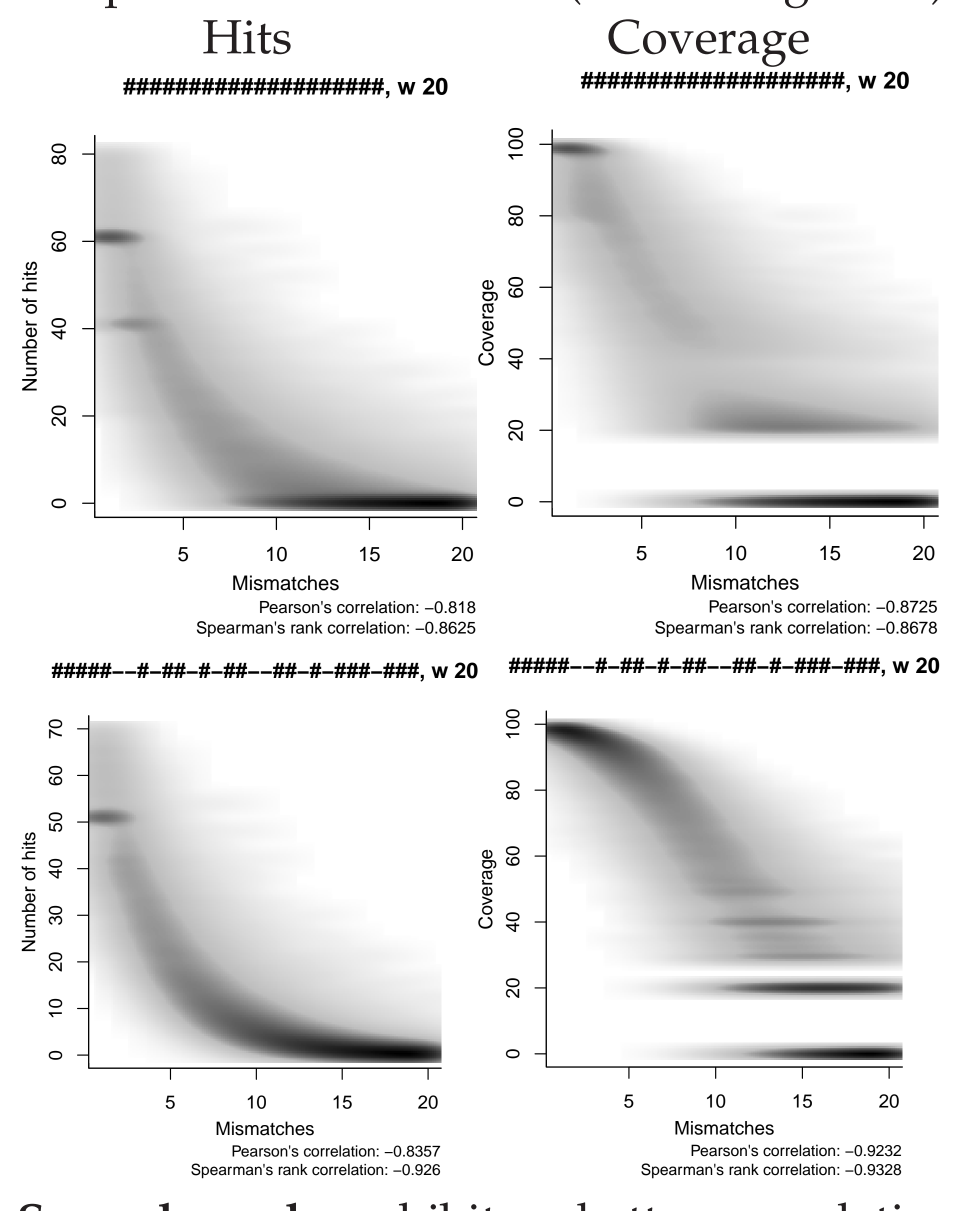
matching position, — don't care position.

A seed acts as a mask for comparing short oligonucleotides. The number of #'s in a seed, called *weight*, defines the number k of matching nucleotides. In the above example k = 4, seed span = 6, and the matching (spaced) k-mer is **ACTC** signifying a **hit**.

A **coverage** is a number of aligned pairs covered by # from a spaced seed matches while sliding over an alignment (=4 above).

SCORES ON REAL GENOMES

We generated a set of ILLUMINA-like single-end reads: we've selected random substrings of M.tuberculosis genome of length L=100 and introduced k mismatch errors, with k random between 1 and 20. For each read, we computed: **number of hits** and **coverage** to the genome under a given seed. A typical plot error vs score (seed weight 20).

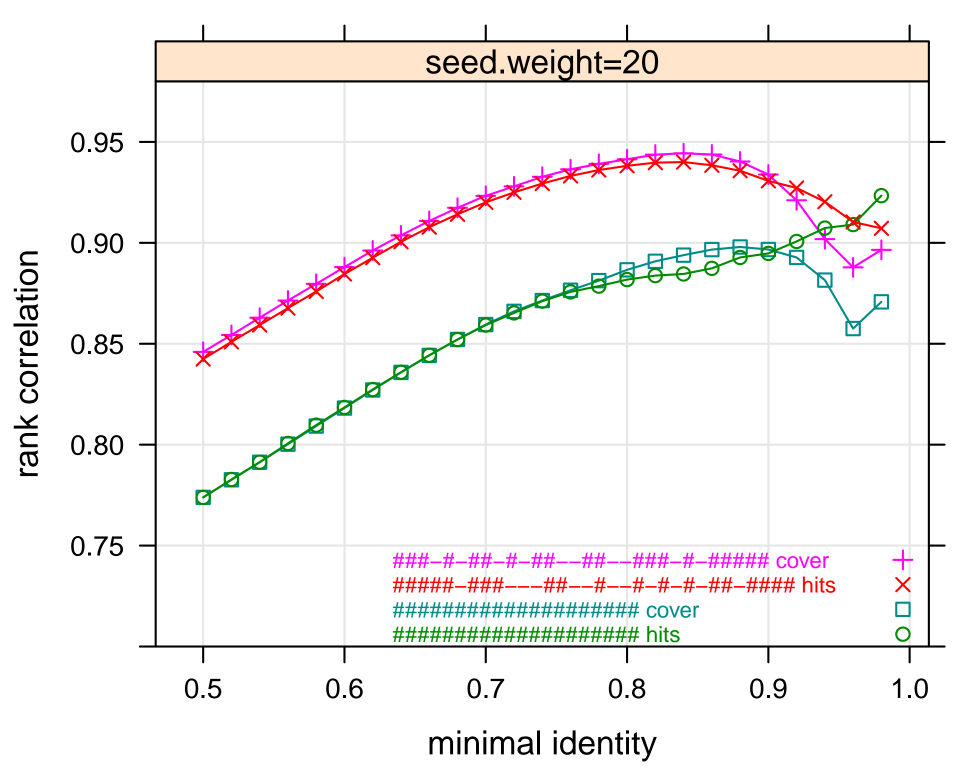


Spaced seeds exhibit a better correlation between errors and score, while *contiguous* seeds plots are more blurred.

SIMULATED ALIGNMENTS

An accurate mapping of a read to a corresponding clade requires estimating its distances to each of the genomes.

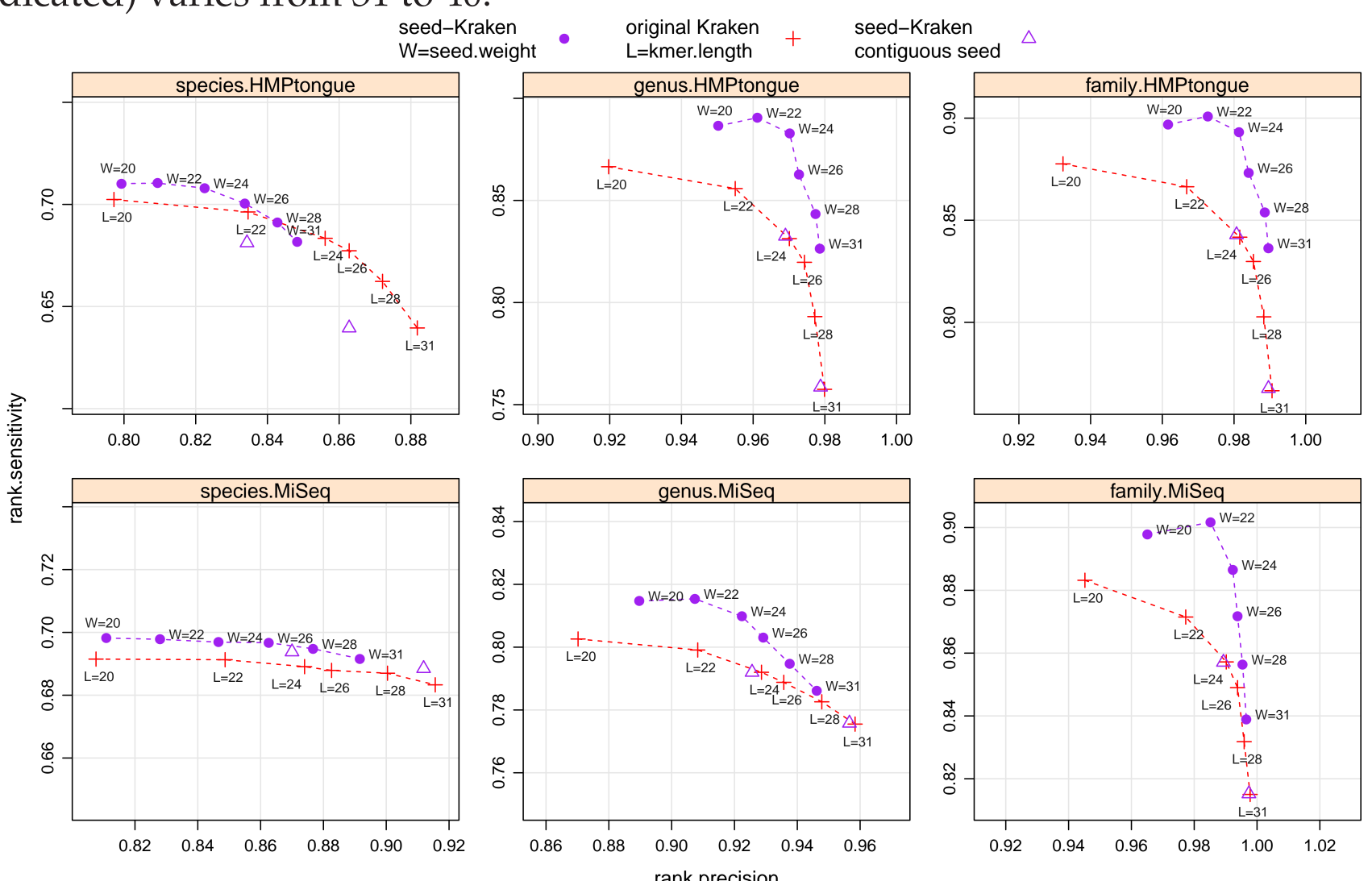
For a fixed minimal identity rate p_{id} , we randomly sampled gapless alignments of length 100 with identity rate from interval $[p_{id}..1]$, and collected pairs (number of mismatches, score), where 'score' stands for either **number of hits**, or **coverage** of a given seed. For these data, we plot Spearman's rank correlation.



In conclusion, spaced seeds provide a much better distance estimator for alignments whose score ranges over a large interval. For very high-scoring alignments (>95%) of identity, the hit number of contiguous seed becomes a better estimator.

SEED-KRAKEN

We modified KRAKEN software [3] to make it work with spaced seeds rather than with contiguous seeds only. For a set of genomes, a database of spaced k-mers matching a user-selected seed is constructed. Performances of classification of SEED-KRAKEN (spaced seed modification), and original KRAKEN, were computed on simulated metagenomes (primarily described and used in [3]): MiSeq (10 bacterial genomes, average error rate), HiSeq, and on 50K subsample of Human Microbiome Project (HMP) Tongue Dorsum wgs sample. Charted are genus precision (positive predictive value) against genus sensitivity (rate of correct assignments). Varying are k-mer length, and its spaced seed equivalent seed weight, while the seed span (not indicated) varies from 31 to 40.



SEED-KRAKEN outperforms original KRAKEN in sensitivity/precision trade-off (ROC curve characteristics) at the classification levels of genus, and family.

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https://github.com/gregorykucherov/ ithaka



