

1 A Succinct Four Russians Speedup for Edit 2 Distance Computation and One-against-many 3 Banded Alignment

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12 — Abstract —

13 The classical Four Russians speedup for computing edit distance (a.k.a. Levenshtein distance),
14 due to Masek and Paterson [15], involves partitioning the dynamic programming table into k -by-
15 k square blocks and generating a lookup table in $O(\psi^{2k}k^2|\Sigma|^{2k})$ time and $O(\psi^{2k}k|\Sigma|^{2k})$ space
16 for block size k , where ψ depends on the cost function (for unit costs $\psi = 3$) and $|\Sigma|$ is the size
17 of the alphabet. We show that the $O(\psi^{2k}k^2)$ and $O(\psi^{2k}k)$ factors can be improved to $O(k^2 \lg k)$
18 time and $O(k^2)$ space. Thus, we improve the time and space complexity of that aspect compared
19 to Masek and Paterson [15] and remove the dependence on ψ .

20 We further show that for certain problems the $O(|\Sigma|^{2k})$ factor can also be reduced. Using this
21 technique, we show a new algorithm for the fundamental problem of one-against-many banded
22 alignment. In particular, comparing one string of length m to n other strings of length m with
23 maximum distance d can be performed in $O(nm + md^2 \lg d + nd^3)$ time. When d is reasonably
24 small, this approaches or meets the current best theoretic result of $O(nm + nd^2)$ achieved by
25 using the best known pairwise algorithm running in $O(m + d^2)$ time [17, 22] while potentially
26 being more practical. It also improves on the standard practical approach which requires $O(nmd)$
27 time to iteratively run an $O(md)$ time pairwise banded alignment algorithm.

28 Regarding pairwise comparison, we extend the classic result of Masek and Paterson [15] which
29 computes the edit distance between two strings in $O(m^2/\log m)$ time to remove the dependence
30 on ψ even when edits have arbitrary costs from a penalty matrix. Crochemore, Landau, and
31 Ziv-Ukelson [8] achieved a similar result, also allowing for unrestricted scoring matrices, but
32 with variable-sized blocks. In practical applications of the Four Russians speedup wherein space
33 efficiency is important and smaller block sizes k are used (notably $k < |\Sigma|$), Kim, Na, Park, and
34 Sim [13] showed how to remove the dependence on the alphabet size for the unit cost version,
35 generating a lookup table in $O(3^{2k}(2k)!k^2)$ time and $O(3^{2k}(2k)!k)$ space. Combining their work
36 with our result yields an improvement to $O((2k)!k^2 \lg k)$ time and $O((2k)!k^2)$ space.

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39 method of the Four Russians

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42 **1 Introduction**

43 Edit distance (a.k.a. Levenshtein distance) is one of the most natural and ubiquitous
 44 measures of similarity between two strings. In the most common variant, *unit cost*, it counts
 45 the minimum number of edits needed to transform one string into another. Here, we use
 46 the Levenshtein definition of *edits* which include insertions, deletions, or substitutions of a
 47 single character. However, in some cases edit operations may be assigned differing costs
 48 from a penalty matrix and additional operations (e.g. inversions or transpositions) may be
 49 considered. Computing this distance is a fundamental problem with applications in many
 50 areas such as computation biology, natural language processing, and information theory.

51 The most well known algorithms use dynamic programming to solve the problem in
 52 $O(m^2)$ time where m is the length of the strings. The only improvement to this has been the
 53 Four Russians algorithm [15], running in $O(m^2/\log m)$ time. While the conditional hardness
 54 results, such as [3], suggest this is unlikely to be improved further for arbitrary strings even
 55 on small alphabets [5].

56 The problem of comparing a string against a large set of sequences is of central importance
 57 in domains such as computational biology, information retrieval, and databases. The banded
 58 alignment variant (a.k.a. the d differences approximate string matching problem), in which
 59 we only report the distance when it is at most some parameter d is also highly relevant.
 60 It's useful in numerous settings wherein we only care about finding small distances or the
 61 maximum distance between any two strings in known to be small. In gene clustering for
 62 example, solving this problem is a key subroutine in many *greedy* clustering heuristics wherein
 63 we iteratively choose a cluster center and form a cluster by recruiting all strings which are
 64 within some small maximum distance d of the center [6]. With the development of faster
 65 and cheaper DNA sequencing technologies, metagenomic sequencing datasets can contain
 66 over 1 billion sequences [7].

67 Another area of research surrounding the Four Russians speedup is how to apply it in
 68 practice. While the theoretical result uses a block size of $\log m$, such a large block size is
 69 impractical due the size of the lookup table exceeding hardware constraints. For the unit
 70 cost version, [13] showed how to drastically reduce the required space, especially for large
 71 alphabets, by avoiding redundant string comparisons. We show that our approach can be
 72 combined with theirs to reduce the space (and preprocessing time) requirement even further.

73 **1.1 Related Work**

74 The edit distance problem is extremely well-studied and the following related work is by no
 75 means exhaustive. We focus primarily on the aspects most related to this paper: pairwise
 76 comparison, the Four Russians speedup, and one-against-many comparison. For simplicity,
 77 we describe all results in the context wherein all strings have length exactly equal to m .

78 The most well-known approach for computing the edit distance between a pair of strings
 79 of length m uses dynamic programming and requires $O(m^2)$. This was later improved to
 80 $O(m^2/\log m)$ in 1980 using the Four Russians speedup [15, 14] and [8] achieved $O(m^2/\log m)$
 81 for unrestricted scoring matrices. The Four Russians speedup, originally proposed for matrix
 82 multiplication, has been adapted to many problems besides edit distance including: RNA
 83 folding [10], transitive closure of graphs [20], and matrix inversion [4]. On the negative
 84 side, [3] recently showed that no algorithm for edit distance can do better than $m^{2-\epsilon}$ time
 85 unless the Strong Exponential Time Hypothesis (SETH) is false and [5] extended this to

86 include strings on a binary alphabet. They accomplished this by reducing a satisfiability
87 problem to edit distance and showing that a subquadratic algorithm for edit distance implies
88 a subexponential algorithm for satisfiability. However, if we fix a maximum distance d and
89 only care about reporting the exact distance when it's less than d , we call this the *banded*
90 *alignment* problem. This problem has seen improvements to $O(md)$ time [9] and the current
91 best algorithm takes only $O(m + d^2)$ time [17, 22].

92 One-against-many edit distance comparison involves comparing a single string to a set
93 of n other strings. Here, we consider only the banded alignment version of the problem
94 wherein we seek to find the distance to all strings within maximum distance d . This problem
95 can be solved in $O(nm + nd^2)$ or $O(nmd)$ time by iteratively applying the pairwise banded
96 alignment algorithms discussed above. Heuristic approaches may run much faster in practice
97 by exploiting properties of the input strings such as prefix similarity and storing the set of
98 strings in a clever data structure such as a trie or BK-tree [9]. However, little theoretical
99 progress has been made. A popular approach to this problem in the context of spell checkers
100 employs Levenshtein automata and/or transducers [21, 16, 12]. Assuming d is a fixed constant,
101 these algorithms run in $O(nm)$ time. However, in practice they consider extremely small
102 values of d (at most 3 or 4) and their runtime appears to grow exponentially in d . In the
103 context of gene clustering in computational biology, [6] show that all pairs banded alignment
104 can be performed in $O(n^2m)$ time under the assumption that all strings are extremely
105 similar. They also use an extension of the Four Russians speedup to one-against-many
106 banded alignment, but our approach to this problem requires no assumptions on the input
107 strings.

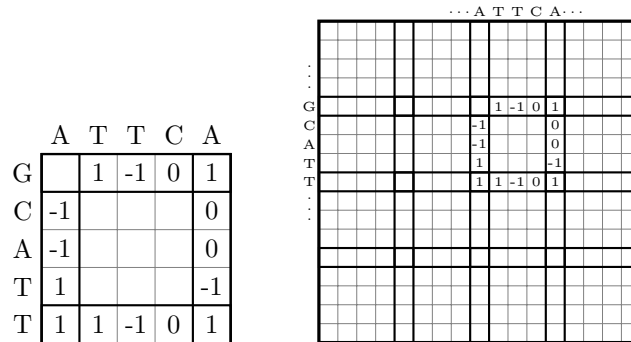
108 The Four Russian speedup is well-studied in context of the regular expression membership
109 problem where the goal is to determine if a particular string matches a given regular
110 expression. Myers[18] showed that for a regular expression of length P and a string of length
111 m , the exact regular expression membership problem (no mismatches are allowed) can be
112 solved in $O(mP/\log m)$ time using the Four Russian speedup compared to the naive $O(mP)$
113 runtime. Wu, Manber, and Myers [23] extended this result for approximate regular expression
114 membership problem where the goal is to check if a string is within an edit distance d from
115 the given regular expression. They showed that approximate regular expression matching
116 problem can be solved in $O(mP/\log_{d+2} m)$ time.

117 Space efficiency is also a major concern in practical applications of the Four Russians
118 speedup since the entire lookup table must be stored in main memory. Thus, block sizes
119 as small as $k = 4$ or 5 may be used. The classical approach for the unit cost variant uses
120 $O(3^{2k}k|\Sigma|^{2k})$ space. Kim, Na, Park, and Sim [13] showed how to remove the dependence on
121 the alphabet size, generating a lookup table in $O(3^{2k}(2k)!k^2)$ time and $O(3^{2k}(2k)!k)$ space.
122 This offers a significant improvement, for example, when $|\Sigma| = 20$ for protein sequences or
123 $|\Sigma| = 26$ for the English language.

124 1.2 Preliminaries

125 For simplicity of presentation, we assume all strings have equal length m . However, the
126 results extend easily to the case where strings have different lengths. We assume the lookup
127 table is any data structure that can perform lookups and insertions in $O(k)$ time for blocks
128 which are identified by distinct keys of length $O(k)$.

■ **Figure 1** Example of classic Four Russians. **Left:** a single block. Notice that for any input in the upper left corner, we can sum that value with one path along the edges of the block to recover the value in the lower right corner. Note that the offset value in the lower right corner may be different for the row and column vectors overlapping at that cell. In this case, the lower right cell is one more than its left neighbor and one less than its above neighbor. **Right:** the full dynamic programming table divided into sixteen 5×5 blocks. Note that the offset values in the example block may not correspond to the optimal alignment of the two substrings shown since they depend on the global alignment between the two full length strings.



129 1.2.1 The classical Four Russians speedup

130 In the classical Four Russians speedup of edit distance computation due to [15, 14], the
 131 dynamic programming table is broken up into square *blocks* of size k -by- k as shown in the
 132 right of Fig. 1. These blocks are tiled such that they overlap by one column/row on each
 133 side (for a thorough description see [11]).

134 The high level idea of the Four Russians speedup is to precompute all possible solutions
 135 to a *block function* and store them in a lookup table. The block function takes as input the
 136 two substrings to be compared in that block and the first row and column of the block itself
 137 in the dynamic programming table. It outputs the last row and column of the block. We
 138 can see in Figure 1 that given the two strings and the first row and column of the table,
 139 such a function could be applied repeatedly to compute the lower right cell of the table and
 140 therefore, the edit distance.

141 There are several tricks that reduce the number of inputs to the block function to bound
 142 the time and space requirements of the lookup table. For example, when the edits have unit
 143 cost, the input row and column for each block can be reduced to vectors in $\{-1, 0, 1\}^k$. These
 144 *offset vectors* encode only the difference between one cell and the next (see Fig. 1) which is
 145 known to be in $\{-1, 0, 1\}$. It has also been shown that the upper left corner does not need
 146 to be included in the offset vectors. This bounds the number of possible row and column
 147 inputs at 3^k each [15]. More generally, when edit costs are derived from a penalty matrix,
 148 the number of row/column inputs is bounded by ψ^k where ψ is the number of possible offset
 149 values and depends on the penalty matrix.

150 1.3 Our Contributions

151 We show a new way to store and query block functions. For a given pair of strings cor-
 152 responding to a k -by- k block in the dynamic programming table, we store an entry in the
 153 lookup table using only $O(k^2 \lg k)$ time and $O(k^2)$ space. We show how to query this entry
 154 in $O(k)$ time. By contrast, the classical approach requires $O(\psi^{2k} k^2)$ time and $O(\psi^{2k} k)$ space,
 155 where ψ is the number of possible offset values and depends on the costs of edits, to store a

lookup entry for a pair of strings since it computes the function for all possible row/column offset vectors and $O(k)$ time per query. Thus, we improve the time and space complexity of that aspect by a factor of at least ψ^{2k}/k and remove the dependence on ψ . This result is stated in Theorem 1.

► **Theorem 1.** *Given two strings corresponding to a k -by- k block, we can store a lookup entry using $O(k^2 \lg k)$ time and $O(k^2)$ space such that given any values for the first row and column of the block, we can compute the last row and column of the block in $O(k)$ time.*

We demonstrate the power of our technique for block functions by designing an algorithm for the fundamental problem of one-against-many banded alignment. In particular, comparing one string of length m to n other strings of length m where we only need to report distances within a maximum distance threshold d can be performed in $O(nm + md^2 \lg d + nd^3)$ time. When d is reasonably small, this improves on the common, naive approach which requires $O(nmd)$ time to iteratively run an $O(md)$ time pairwise banded alignment algorithm. It also approaches the best theoretic result of $O(nm + nd^2)$ achieved by using the best known pairwise algorithm running in $O(m + d^2)$ time [17, 22]. We note that the author of [17], describes the $O(m + d^2)$ time algorithm as “impractical” and “primarily of theoretical interest”. We are somewhat more optimistic, observing that our algorithm blends neatly with approaches such as [6] for comparing genetic sequences and as discussed in Section 4.3 can be implemented in a way that exploits the prefix similarity occurring in practice.

► **Theorem 2.** *Performing banded alignment with maximum distance d between a string of length m and n other strings also of length m can be done in $O(nm + md^2 \lg d + nd^3)$ time.*

We extend the classic result of [15] which computes the edit distance between two strings in $O(m^2/\log m)$ time to remove the dependence on ψ even when edits have costs derived from a penalty matrix. Here, the number of entries in the lookup table does not depend on the penalty matrix. We acknowledge that [8] also achieves the same $O(m^2/\log m)$ running time on unrestricted scoring matrices. However, there are some differences between our approach and theirs which may make one or the other more advantageous in different settings. Most notably our approach adheres more closely to the classic Four Russians speedup and uses a uniform block size which is necessary for our one-against-many algorithm. Uniform block sizes also allow our technique to be combined easily with the space-efficient approach in [13] and the gene clustering technique in [6] since both rely on splitting the dynamic programming table into uniform size blocks. In the case of [6], this is crucial to exploiting the prefix similarity among highly conserved genomic sequences. On the other hand, the blocks in [8] vary in size in a clever way to take advantage of the compressibility of the strings being compared. This yields a faster running time for pairwise comparison of strings with small entropy, $O(hn^2/\log n)$, where $h \leq 1$ is the entropy of the text.

► **Theorem 3.** *Given a penalty matrix for edit operations, the edit distance between two strings can be computed in $O(m^2/\log m)$ time.*

In practical applications of the Four Russians speedup wherein space efficiency is important and smaller block sizes k are used (notably $k < |\Sigma|$), [13] showed how to remove the dependence on the alphabet size for the unit cost version, generating a lookup table in $O(3^{2k}(2k)!k^2)$ time and $O(3^{2k}(2k)!k)$ space. Combining their work with our result yields an improvement to $O((2k)!k^2 \lg k)$ time and $O((2k)!k^2)$ space.

► **Theorem 4.** *For a block size k , a lookup table can be generated in $O((2k)!k^2 \lg k)$ time and $O((2k)!k^2)$ space such that we can find the unit cost edit distance between two strings of length m in $O(m^2/k)$ time.*

2 Storing and querying the block function

Here, we consider the crucial subroutine in our algorithms and prove Theorem 1. For a block size k , we first show how to store a lookup entry for any two strings of length k in $O(k^2 \lg k)$ time and $O(k^2)$ space. Then, we show how, given two strings of length k and the first row and column of the block, we can compute the last row and column in $O(k)$ time by querying the corresponding lookup entry. Notice that in contrast to the classical Four Russians speedup, the information we precompute and store for a block function is based only on the two strings being compared. Thus, we avoid having to store an entry for each of the 3^{2k} possible input vectors considered in [15] (For unit costs, they encode rows/columns as offset vectors in $\{-1, 0, 1\}$ since the values in adjacent cells differ by at most 1, yielding 3^k possible inputs each for the row and column vectors).

2.1 Notation

We start by defining some notation, illustrated in Figure 2. Let $U = \{u_1, u_2, \dots, u_{2k-1}\}$ be an ordered set of the cells in the first row and column of the block and let $V = \{v_1, v_2, \dots, v_{2k-1}\}$ be an ordered set of the cells in the last row and column of the block. For both sets, the ordering starts with the upper right corner and ends in the lower left corner. Thus, both u_1 and v_1 correspond to the upper right corner, u_k corresponds to the upper left corner, v_k corresponds to the lower right corner, and both u_{2k-1} and v_{2k-1} correspond to the lower left corner.

For each pair of cells (u, v) , we store the least cost $c_{u,v}$ of any path through the block from u to v . If no such path exists, we set $c_{u,v} = \infty$ and if u and v correspond to the same cell, we set $c_{u,v} = 0$. Note that $c_{u,v}$ is not necessarily based on the optimal alignment within the entire block. It corresponds to an alignment of the subset of the block with u as the upper left corner and v as the lower right. Also, recall that this block will be part of a larger dynamic programming table and the path through the block corresponding to the best global alignment may not be the same as the path corresponding to the best local alignment within the block.

We can think of this set of costs as a complete, weighted bipartite graph $G = \{U, V, U \times V\}$ with weights $c_{u,v}$ on the edges. We also use c_u and c_v to denote the values stored in the corresponding cells of the block within the dynamic programming table. When we query a block function for two strings, the c_u values (input row and column) will be given as input and our goal will be to compute the c_v values (output row and column). Thus, if we consider the values stored in the cells after the full dynamic programming table has been computed, we have that $c_v = \min_{u \in U} (c_u + c_{u,v})$.

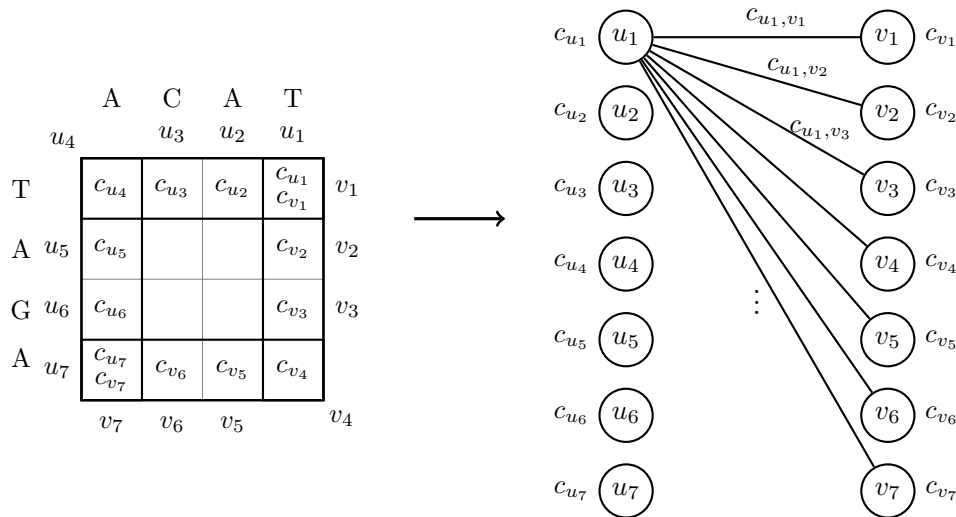
2.2 Storing lookup entries

For every pair of substrings we wish to query eventually, our lookup table will simply store the cost c_{uv} for every edge in the graph G defined by comparing those substrings. These cost values will be stored in a $|V| \times |U|$ matrix M with a row for each $v \in V$ and a column for each $u \in U$. Cell M_{ji} will contain $c_{u_i v_j}$. We now show that computing G and storing M for any pair of substrings of length k can be done in $O(k^2 \lg k)$ time.

► **Lemma 5.** *Given a pair of strings of length k , we can compute all $c_{u,v}$ in $O(k^2 \lg k)$ time.*

Proof. Note that each $c_{u,v}$ can be seen as the weight of the shortest path in a grid graph of dimension $k \times k$. Thus the algorithm of [19] can be applied. That algorithm requires

■ **Figure 2** Illustration of how the dynamic programming table is represented as a bipartite graph of least cost paths. **Left:** The dynamic programming table for a block comparing the strings “ACAT” and “TAGA” with all u, v, c_u , and c_v labeled. **Right:** The bipartite graph representation. Note that this will be a complete, weighted bipartite graph with costs $c_{u,v}$ for all pairs in $U \times V$.



245 $O(k^2 \lg k)$ preprocessing time and can then compute each of the $O(k^2)$ $c_{u,v}$ entries in $O(\lg k)$
 246 time. This leads to an overall running time of $O(k^2 \lg k)$.

248 For completeness, we also state the simple fact that the space requirement for an entry is
 249 $O(k^2)$.

250 ► **Lemma 6.** *Given a pair of strings of length k , storing the entry requires $O(k^2)$ space.*

251 **Proof.** The proof follows directly from the fact that we are simply storing the edges of a
 252 complete, weighted bipartite graph $G = \{U, V, U \times V\}$ with $|U| = |V| = 2k - 1$.

253 2.3 Querying a block function

254 Given the two substrings and the input row and column vectors, we now show how to use
 255 our lookup entry matrix M to compute the output row and column (a.k.a all c_v for $v \in V$)
 256 in $O(k)$ time.

257 ► **Lemma 7.** *Given the input row and column vectors and the $O(k) \times O(k)$ lookup entry
 258 matrix M , we can compute the output row and column in $O(k)$ time using the SMAWK
 259 algorithm [2].*

260 **Proof.** Let \vec{w} be the vector of all c_u values generated from the input row and column vectors.
 261 Scaling each column of M by the corresponding cell in \vec{w} gives us a new matrix M' wherein the
 262 minimum value in each row j is our desired output value $c_{v_j} = \min_{u \in U} (c_u + c_{u,v_j})$. It is known
 263 that M' is totally monotone [1, 19] and thus we can find row minima in $O(|U|) = O(k)$ time
 264 using the classic SMAWK algorithm [2]. Note that we need not explicitly generate M' since
 265 the value of any cell we wish to query can be computed from M and \vec{w} as $M'_{ji} = M_{ji} + \vec{w}_i$.

266 The proof of Theorem 1 follows from Lemmas 5, 6, and 7.

267 **2.4 Alternatives to query a block function without SMAWK**

268 While our algorithm for banded alignment in Section 3 uses larger block sizes than the typical
 269 pairwise Four Russians approach, in many cases, the blocks will be small enough for SMAWK
 270 to be inefficient in practice. As such, we introduce a simpler query algorithm here and briefly
 271 discuss the potential for future work to speed up the query function in practice.

272 This simpler query algorithm achieves a slightly worse asymptotic running time of $O(k \lg k)$
 273 and can be described as follows. Recall that our goal is to find the minimum value of each
 274 row in the totally monotone matrix M' with $|U|$ columns and $|V|$ rows. We first find the
 275 minimum value in row $|V|/2$ and let *mincol* be the column containing that cell. We then
 276 perform the same operation recursively on two submatrices of M' . The first submatrix
 277 includes all rows up to $|V|/2$ and all columns up to (and including) *mincol*. The second
 278 includes the rows after $|V|/2$ and columns from *mincol* to $|U|$. We do not claim this simpler
 279 algorithm is a novel approach to finding row minima and include it merely to illustrate
 280 possible alternatives to SMAWK.

281 ► **Lemma 8.** *The algorithm described here runs in $O(k \lg k)$ time and outputs the correct*
 282 *result.*

283 **Proof.** For the running time, note that each recursive call nearly partitions the columns
 284 of M' (pairs of submatrices overlap at single columns), resulting in $O(|U|) = O(k)$ time
 285 spent at each level of recursion. Since we split the rows in half at each level, there will be
 286 $O(\lg |V|) = O(\lg k)$ levels total, giving a final running time of $O(k \lg k)$.

287 The correctness follows directly from the properties of totally monotone matrices also
 288 utilized in the analysis of SMAWK. ◀

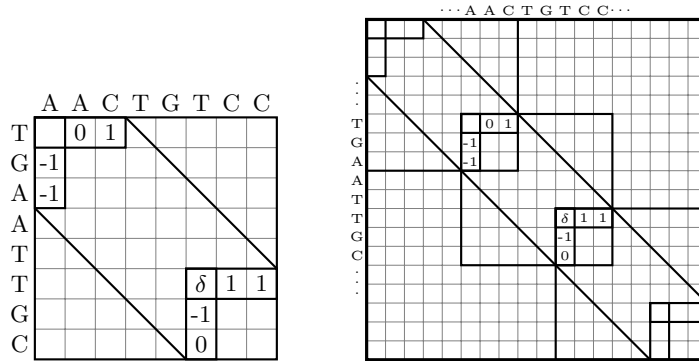
289 Looking to the future, we note that neither SMAWK nor the algorithm in this section
 290 leverage all of the specific properties of the matrix M' . For example, M' is not an arbitrary
 291 totally monotone matrix. It comes from M , a matrix which we can afford to spend k^2 time
 292 preprocessing, scaled by \vec{w} , a vector with the property that adjacent cells differ by at most 1
 293 in the unit cost setting.

294 **3 One-against-many comparison**295 **3.1 Extending the Four Russians approach to banded alignment**

296 For our algorithm for one against many banded alignment, we use the extension to banded
 297 alignment from [6] which simplifies both the analysis and practical implementation. The
 298 extension uses a slightly different block function and way of tiling blocks to cover the relevant
 299 diagonal “band” of the dynamic programming table. The blocks now overlap on a square of
 300 size $d + 1$ at the upper left and lower right corners. We will call these overlapping regions
 301 *overlap squares*. The block function still takes as input the two substrings to be compared.
 302 The set U contains only the first row and column of the the upper left overlap square and
 303 V contains only the first row and column of the lower right overlap square as well as the
 304 difference between the upper left corners of the two overlap squares.

305 Thus, we can move directly from one block to the next, storing a sum of the differences
 306 between the upper left corners. In this case, reaching the final lower right cell of the table
 307 requires an additional $O(d^2)$ operation to fill in the last overlap square, but this adds only a
 308 negligible factor to the running time.

■ **Figure 3** Example of our approach to the Four Russians speedup. **Left:** a block for maximum edit distance $d = 2$. The output δ represents the offset from the upper left corner of one block to the upper left corner of the next block. Note that we only need to consider a diagonal band of the block itself. **Right:** using these blocks to cover the diagonal band of the dynamic programming table in the context of banded alignment.



309 **3.2 Our algorithm**

310 We start with some notation and definitions. For a string s , let $s_{i,i+k}$ be a length k
 311 substring starting at index i of s . We define two types of block comparisons, *identities* and
 312 *differences*, based on the strings being compared. An identity comparison is between the
 313 substring $s_{i,i+k}$ and another substring that is identical to one of the substrings $s_{j,j+k}$ for
 314 $j \in \{i - d, i - d + 1, \dots, i, \dots, i + d\}$. All other comparisons are difference comparisons.
 315 In other words, identity comparisons between two strings will come from long common
 316 subsequences between the two strings. Difference comparisons will come from the locations
 317 where an edit occurs. Note that we can stop comparing two strings once we've encountered
 318 more than d differences among their prefixes. Let S be a set of strings and let p be the single
 319 string we wish to compare to all strings in S .

320 The algorithm can be summarized as follows. We first compute and store lookup entries for
 321 all possible identity comparisons for each block in p . We then perform pairwise comparisons
 322 between p and each string in S . A pairwise comparison is computed as follows. For each
 323 block, we first query the lookup table using the corresponding substrings. If we find an
 324 entry (similarity comparison), we query it as described in Section 2. Otherwise (difference
 325 comparison), we perform standard banded alignment on the two strings with the first row
 326 and column of the table initialized to the values of the input row and column of the block.
 327 If at any time during a pairwise comparison the distance accumulated exceeds d , then we
 328 immediately halt and move on to the next pair.

329 We divide the analysis into three parts: the time to compute and store the lookup table,
 330 the time to query the lookup table during pairwise comparison, and the time to compute the
 331 block function for difference comparisons.

332 ► **Lemma 9.** *The time to compute and store the lookup table for all block identity comparisons*
 333 *in a single string p of length m and max distance d is $O(md^2 \lg d)$.*

334 **Proof.** Let the block size $k = 2d$. Then p will be divided into $m/d - 1$ blocks. For any
 335 given block, let $p_{i,i+k}$ be the substring of p corresponding to that block. Then, for every
 336 $j \in \{i - d, i - d + 1, \dots, i, \dots, i + d\}$, we need to store the comparison between $p_{i,i+k}$ and
 337 $p_{j,j+k}$. We need not compare $p_{i,i+k}$ to any substrings outside this range since that would
 338 imply an alignment of distance greater than d . Thus, for each block we need to store lookups

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339 for at most $2d + 1 = O(d)$ different identity comparisons. Computing the lookup entry for
340 each comparison takes $O(k^2 \lg k) = O(d^2 \lg d)$ time by Theorem 1. Putting it all together,
341 we have $O(m/d \cdot d \cdot d^2 \lg d) = O(md^2 \lg d)$. ◀

342 ▶ **Lemma 10.** *Excluding the time to compute block functions for difference comparisons, the*
343 *time to compare a string p of length m to n other strings using the precomputed lookup table*
344 *is $O(nm)$.*

345 **Proof.** Each pairwise comparison involves computing $m/d - 1$ block functions. If a block
346 corresponds to an identity comparison querying the block function takes $O(k) = O(d)$ time
347 by Theorem 1. Otherwise, if it's a difference comparison block, the only time will come from
348 checking the lookup table which we've assumed takes $O(d)$ time. It follows that the running
349 time for each pairwise comparison is $O(m)$ and comparing p to all n strings requires $O(nm)$
350 time. ◀

351 ▶ **Lemma 11.** *The time needed to compute block functions for difference comparisons between*
352 *p and all n other strings is $O(nd^3)$.*

353 **Proof.** Notice that each edit is present in at most two overlapping blocks. It follows that
354 for a given pair of strings, the number of block queries corresponding to differences can be
355 at most $2(d + 1) = O(d)$ since we will halt a comparison if the distance ever reaches $d + 1$
356 or more. Thus, the running time to compute the full dynamic programming for difference
357 blocks for all n pairwise comparisons is $O(n \cdot d \cdot d^2) = O(nd^3)$. ◀

358 The proof of Theorem 2 follows from combining Lemmas 9, 10, and 11.

359 4 Extensions and applications

360 In this section, we briefly show how the results of Section 2 can be applied to other settings
361 in which the Four Russians speedup is used for computing string edit distance.

362 4.1 Comparing two arbitrary strings with a penalty matrix

363 As with the classical Four Russians, when the alphabet size is constant, we can choose the
364 block length k to be an appropriate logarithmic function of the string length m such that
365 the lookup table can be computed efficiently. For an alphabet Σ , there are $|\Sigma|^{2k}$ pairs of
366 string of length k . By Theorem 1, each pair requires $O(k^2 \lg k)$ time to compute the lookup
367 entry regardless of the costs of the edits. Thus, the preprocessing for $k = (\log_{|\Sigma|} m)/2$ takes
368 $O(m \log^2 m \log \log m)$ time. Since the total number of blocks in the dynamic programming
369 table is $O(m^2/k^2)$ and computing each block function from the lookup table takes $O(k)$
370 time by Theorem 1, the running time to compute the distance using the lookup table is
371 $O(m^2/\log m)$. This completes the proof of Theorem 3.

372 4.2 Improved space-efficiency

373 The approach in Section 2 can be combined with the work of [13] to achieve the improved
374 time and space bound in Theorem 4 for computing the lookup table. Notice that Theorem 1
375 gives a time and space bound for each pair of substrings for which we need to compute
376 a block function. Specifically, each pair of strings contributes $O(k^2 \lg k)$ time and $O(k^2)$
377 space. As a complement, [13] showed how to encode strings in such a way that we reduce the
378 number of redundant string comparisons. There, the number of strings compared is reduced
379 to $O((2k)!)$. Theorem 4 follows from these simple observations.

380 4.3 Exploiting prefix similarity in one-against-many comparison

381 Since the one-against-many banded alignment algorithm in Section 3 uses the same extension
 382 to banded alignment as [6], it can be combined with other techniques from that paper. In
 383 particular, they divide all of the strings in the database S into blocks and store the blocks in
 384 a trie-like data structure. This allows them to exploit prefix similarity of the strings of S and
 385 further improve the running time in practice. Additionally, that uses *lazy computation*, the
 386 technique of computing and storing the lookup table on-the-fly rather than precomputing it
 387 to heuristically avoid comparing substrings which don't actually appear in the dataset. In
 388 the context of Theorem 2, that could potentially reduce the md^3 factor.

389 5 Conclusion and future directions

390 In this paper, we provided an approach to storing and querying block functions in the
 391 Four Russians speedup for edit distance computation using less time and space than the
 392 original method. We demonstrated how this approach can lead to an algorithm for the
 393 one-against-many banded alignment problem. Finally, we showed how our approach can
 394 easily be combined with prior work to gain additional improvements such as space-efficiency.

395 The problems of comparing two similar strings and one-against-many comparison of
 396 highly similar strings have applications in variety of domains. For example, searching a query
 397 sequence against the database of multiple sequence within a certain similarity threshold is
 398 one of the basic tasks in designing database management systems. In the case of document
 399 plagiarism detection, the task is to compare two documents which are assumed to be highly
 400 similar to each other. In the case of computational biology, sequence similarity detection is a
 401 ubiquitous task in most analysis. Although there have been efficient algorithms proposed in
 402 literature, they are not very easy or practical to implement on a routine basis. Our algorithm
 403 may bridge this gap and be easier to implement while yielding similar theoretical bounds.

404 There are many questions and potential future directions following this work. One natural
 405 question is whether the techniques in this paper can be applied to other problems yielding a
 406 Four Russians speedup. In many cases, such as boolean matrix multiplication, the answer is
 407 no. However, problems more closely related to edit distance may yield some improvement.
 408 Regarding the specific problems in this paper, the $O(nd^3)$ term in the one-against-many
 409 result can likely be improved to $O(nd^2)$ to match [17] and doing so using practical techniques
 410 would be a nice addition to this work. Similarly, improving the constant factors in the
 411 query by using a more specialized algorithm than SMAWK (even an asymptotically worse
 412 algorithm) could enhance the practical applications of our approach. On the hardness side,
 413 which of these results are tight?

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