Exact global alignment using A* with chaining seed heuristic and match pruning

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Abstract

Motivation: Sequence alignment has been at the core of computational biology for half a century. Still, it is an open problem to design a practical algorithm for exact alignment of a pair of related sequences in linear-like time (Medvedev, 2022b).

Methods: We solve exact global pairwise alignment with respect to edit distance by using the A* shortest path algorithm. In order to efficiently align long sequences with high divergence, we extend the recently proposed seed heuristic (Ivanov et al., 2022) with match chaining, gap costs, and inexact matches. We additionally integrate the novel match pruning technique and diagonal transition (Ukkonen, 1985) to improve the A* search. We prove the correctness of our algorithm, implement it in the A*PA aligner, and justify our extensions intuitively and empirically.

Results: On random sequences of divergence \(d=4\%\) and length \(n\), the empirical runtime of A*PA scales near-linearly with length (best fit \(n^{1.06}, n \leq 10^7\) bp). A similar scaling remains up to \(d=12\%\) (best fit \(n^{1.24}, n \leq 10^7\) bp). For \(n=10^7\) bp and \(d=4\%\), A*PA reaches \(500\times\) speedup compared to the leading exact aligners EdLib and BiWFA. The performance of A*PA is highly influenced by long gaps. On long \((n=500\, \text{kb})\) ONT reads of a human sample it efficiently aligns sequences with \(d<10\%\), leading to a median speedup compared to EdLib and BiWFA. When the sequences come from different human samples, A*PA performs \(1.7\times\) faster than EdLib and BiWFA.

Availability: github.com/RagnarGrootKoerkamp/astar-pairwise-aligner

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1 Introduction

The problem of aligning one biological sequence to another is known as global pairwise alignment (Navarro, 2001). Among others, it is applied to genome assembly, read mapping, variant detection, and multiple sequence alignment (Prijibelski et al., 2019). Despite the centrality and age of pairwise alignment (Needleman and Wunsch, 1970), “a major open problem is to implement an algorithm with linear-like empirical scaling on inputs where the edit distance is linear in \(n\)” (Medvedev, 2022b).

Alignment accuracy affects subsequent analyses, so a common goal is to find a shortest sequence of edit operations (single letter insertions, deletions, and substitutions) that transforms one sequence into the other. The length of such a sequence is known as Levenshtein distance (Levenshtein, 1966) and edit distance. It has recently been proven that edit distance cannot be computed in strongly subquadratic time, unless SETH is false (Backurs and Indyk, 2015). When the number of sequencing errors is proportional to the length, existing exact aligners scale quadratically both in the theoretical worst case and in practice. Given the increasing amounts of biological data and increasing read lengths, this is a computational bottleneck (Kucherov, 2019).

We solve the global alignment problem provably correct and empirically fast by using A* on the alignment graph and building on many existing techniques. Our implementation A*PA (A* Pairwise Aligner) scales near-linear with length up to \(10^7\) bp long sequences with divergence up to \(12\%\). Additionally, it shows a speedup over other highly optimized aligners when aligning long ONT reads.

Fig. 1. Computed states per algorithm. Various optimal alignment algorithms and their implementation are demonstrated on synthetic data (length \(n=500\) bp, divergence \(d=16\%\)). The colour indicates the order of computation from blue to red. (a) Band-doubling (EdLib), (b) Dijkstra, (c) Diagonal transition/DT (WFA), (d) DT with divide-and-conquer/D&C (BiWFA), (e) A*PA with gap-chaining seed heuristic (GCSH), match pruning, and DT (seed length \(k=5\) and exact matches).

1.1 Overview of method

To align two sequences \(A\) and \(B\) globally with minimal cost, we use the A* shortest path algorithm from the start to the end of the alignment graph, as first suggested by Hadlock (1988b). A core part of the A* algorithm is the heuristic function \(h(u)\) that provides a lower bound on the remaining distance from the current vertex \(u\). A good heuristic efficiently computes an accurate estimate \(h\), so suboptimal paths get penalized more and A* prioritizes vertices on a shortest path, thus reaching the target quicker. In this paper, we extend the seed heuristic by Ivanov et al. (2022) in several ways to increase its accuracy for long and erroneous sequences.
Seed heuristic (SH). To define the seed heuristic $h_s$, we split $A$ into short, non-overlapping substrings (seeds) of fixed length $k$ (Fig. 2a). Since the whole sequence $A$ has to be aligned, each of the seeds also has to be aligned somewhere in $B$. If a seed does not match anywhere in $B$ without mistakes, then at least 1 edit has to be made to align it. Thus, the seed heuristic $h_s$ is the number of remaining seeds (contained in $A_{s_j}$) that do not match anywhere in $B$. The seed heuristic is a lower bound on the distance between the remaining suffixes $A_{s_j}$ and $B_{s_j}$. In order to compute $h_s$ efficiently, we precompute all matches in $B$ for all seeds from $A$. Where Ivanov et al. (2022) uses crumbs to mark upcoming matches in the graph, we do not need them due to the simpler structure of sequence-to-sequence alignment.

Chaining (CSH). One drawback of the SH is that it may use matches that do not lie together on a path from $u$ to the end, as for example the matches for $s_3$ and $s_9$ in Fig. 2a. In the chaining seed heuristic $h_{cs}$ (Sec. 3.1), we enforce that the matches occur in the same order in $B$ as their corresponding seeds occur in $A$, i.e., the matches form a chain going down and right (Fig. 2b). Now, the number of upcoming errors is at least the minimal number of remaining seeds that cannot be aligned on a single chain to the target. When there are many spurious matches (i.e. outside the optimal alignment), chaining improves the accuracy of the heuristic, thus reducing the number of states expanded by $A^*$. To compute CSH efficiently, we subtract the maximal number of matches in a chain starting in the current state from the number of remaining seeds.

Gap costs (GCSH). The CSH penalizes the chaining of two matches by the seed cost, the number of skipped seeds in between them. This chaining may skip a different number of letters in $A$ and $B$, in which case the absolute difference between these lengths (gap cost) is a lower bound on the length of a path between the two matches. The gap-chaining seed heuristic $h_{gcs}$ (Fig. 2c) takes the maximum of the gap cost and the seed cost, which significantly improves the accuracy of the heuristic for sequences with long indels.

Inexact matches. To further improve the accuracy of the heuristic for divergent sequences, we use inexact matches (Wu and Manber, 1992; Marco-Sola et al., 2012). For each seed in $A$, our algorithm now finds all its inexact matches in $B$ with cost at most 1. The lack of a match of a seed then implies that at least $r=2$ edits are needed to align it. This doubles the potential of our heuristic to penalize errors.

Match pruning. In order to further improve the accuracy of our heuristic, we apply the multiple-path pruning observation (Poole and Mackworth, 2017): once a shortest path to a vertex $u$ has been found, no other path to $u$ can be shorter. Since we search for a single shortest path, we want to incrementally update our heuristic (similar to Real-Time Adaptive $A^*$ (Koenig and Likhachev, 2006)) to penalize further paths to $u$. We prove that once $A^*$ expands a state $u$ which is at the start or end of a match, indeed it has found a shortest path to $u$. Then we can ignore (prune) such a match, thus penalizing other paths to $u$ (Fig. 2d, Sec. 3.2). Pruning increases the heuristic in states preceding the match, thereby penalizing states preceding the “tip” of the $A^*$ search. This reduces the number of expanded states, and leads to near-linear scaling with sequence length (Fig. 1e).

Diagonal transition (DT). The diagonal transition algorithm only visits so-called furthest reaching states (Ukkonen, 1985; Myers, 1986) along each diagonal and lies at the core of WFA (Marco-Sola et al., 2021) (Fig. 1c). We introduce the diagonal transition optimization to the $A^*$ algorithm that skips states known to be not furthest reaching. This is independent of the $A^*$ heuristic and makes the exploration more “hollow”, especially speeding up the quadratic behavior of $A^*$ in complex regions.

We present an algorithm to efficiently initialize and evaluate these heuristics and optimizations (Sec. 3.3 and App. A), prove the correctness of our methods (App. B), and evaluate and compare their performance to other optimal aligners (Sec. 4 and App. C).

1.2 Related work

We first outline the algorithms behind the fastest exact aligners: DP-based band doubling (used by EDCLIB) and diagonal transition (used by BiWFA). Then, we outline methods that $A^*$PA integrates.

Dynamic programming (DP). This classic approach to aligning two sequences computes a table where each cell contains the edit distance between a prefix of the first sequence and a prefix of the second by reusing the solutions for shorter prefixes. This quadratic DP was introduced for speech signals Vintsyuk (1968) and genetic sequences (Needleman and Wunsch, 1970; Sankoff, 1972; Sellers, 1974; Wagner and Fischer, 1974). The quadratic $O(nm)$ runtime for sequences of lengths $n$ and $m$ allowed for aligning of long sequences for the time but speeding it up has been a central goal in later works. Implementations of this algorithm include SEGAN (Reinert et al., 2017) and PARASAIL (Dally, 2016).

Band doubling and bit-parallelization. When the aligned sequences are similar, the whole DP table does not need to be computed. One such output-sensitive algorithm is the band doubling algorithm of Ukkonen (1985) (Fig. 1a) which considers only states around the main diagonal of the table, in a band with exponentially increasing width, leading to $O(ns)$ runtime, where $s$ is the edit distance between the sequences. This algorithm, combined with the bit-parallel optimization by Myers (1999)
is implemented in EDLIB (Šošić and Šikić, 2017) with $O(ns/w)$ runtime, where $w$ is the machine word size (nowadays 64).

Diagonal transition (DT). The diagonal transition algorithm (Ukkonen, 1985; Myers, 1986) exploits the observation that the edit distance does not decrease along diagonals of the DP matrix. This allows for an equivalent representation of the DP table based on farthest-reaching states for a given edit distance along each diagonal. Diagonal transition has an $O(ns)$ worst-case runtime but only takes expected $O(n + s^2)$ time (Fig. 1c) for random input sequences (Myers, 1986) (which is still quadratic for a fixed divergence $d = s/n$). It has been extended to linear and affine costs in the wavefront alignment (WFA) (Marco-Sola et al., 2021) in a way similar to Gotoh (1982). Its memory usage has been improved to linear in BiWFA (Marco-Sola et al., 2023) by combining it with the divide-and-conquer approach of Hirschberg (1975), similar to Myers (1986) for unit edit costs. Wu et al. (1990) and Papamichail and Papamichail (2009) apply diagonal transition to align sequences of different lengths.

Contours. The longest common subsequence (LCS) problem is a special case of edit distance in which, with gaps are allowed but substitutions are forbidden. Contours partition the state-space into regions with the same remaining number of the LCS subtask (Fig. 3). The contours can be computed in log-linear time in the number of matching elements between the two sequences which is practical for large alphabets (Hirschberg, 1977; Hunt and Szymanski, 1977).

Shortest paths and $A^*$. An alignment that minimizes edit distance corresponds to a shortest path in the alignment graph (Vintsyuk, 1968; Ukkonen, 1985). Assuming non-negative edit costs, a shortest path can be found using Dijkstra’s algorithm (Ukkonen, 1985) (Fig. 1b) or $A^*$ (Hart et al., 1968). $A^*$ is an informed search algorithm which uses a task-specific heuristic function to direct its search, and has previously been found using Dijkstra’s algorithm (Ukkonen, 1985) (Fig. 1b) or $A^*$ (Hart et al., 1968). $A^*$ with the seed heuristic enables provably-exact alignment to the reference. When $A^*$ explores a new state, the seed heuristic sequence is split into non-overlapping substrings ($\langle S_1, S_2, \ldots, S_i, \ldots, S_n \rangle$), each of which is matched exactly to the reference. When $A^*$ explores a new state, the seed heuristic sequence is split into non-overlapping substrings ($\langle S_1, S_2, \ldots, S_i, \ldots, S_n \rangle$), each of which is matched exactly to the reference. $A^*$ with the seed heuristic function to direct its search, and has previously been found using Dijkstra’s algorithm (Ukkonen, 1985) (Fig. 1b) or $A^*$ (Hart et al., 1968). $A^*$ with the seed heuristic enables provably-exact alignment to the reference. When $A^*$ explores a new state, the seed heuristic function to direct its search, and has previously been found using Dijkstra’s algorithm (Ukkonen, 1985) (Fig. 1b) or $A^*$ (Hart et al., 1968) and Papamichail and Papamichail (2009) apply diagonal transition to align sequences of different lengths.

We increase the accuracy of this heuristic in several novel ways: seeds must match in order in the chaining seed heuristic, and gaps between seeds are penalized in the gap-chaining seed heuristic. The novel match pruning technique penalizes states “lagging behind” the tip of the search and turns the otherwise quadratic algorithm into an empirically near-linear algorithm in many cases. Inexact matches (Wu and Manber, 1992; Marco-Sola et al., 2012) increase the divergence of sequences that can be efficiently aligned. We additionally apply the diagonal transition algorithm (Ukkonen, 1985; Myers, 1986), so that only the small fraction of farthest-reaching states needs to be computed. We prove the correctness of our methods, and apply contours (Hirschberg, 1977; Hunt and Szymanski, 1977) to efficiently initialize and evaluate the heuristic. We implement our method in the novel aligner A*PA.

On uniform random synthetic data with 1% divergence, the runtime of $A^*$PA scales linearly with length up to $10^7$ bp and is up to 500× faster than EDLIB and BiWFA. On $>500$ kbp long Oxford Nanopore (ONT) reads of the human genome, $A^*$PA is 3× faster in median than EDLIB and BiWFA when only read errors are present, and 1.7× faster in median when additionally genetic variation is present.

2 Preliminaries

This section provides definitions and notation that are used throughout the paper. A summary of notation is included in App. D.

Sequences. The input sequences $A = \alpha_0\alpha_1\ldots\alpha_n\alpha_n^{-1}$ and $B = \beta_0\beta_1\ldots\beta_m\beta_m^{-1}$ are over an alphabet $\Sigma$ with $4$ letters. We refer to substrings $\alpha_i\ldots\alpha_{i+k}$ as $A_{i+k}$, to prefixes $\alpha_0\ldots\alpha_i$ as $A_i$, and to suffixes $\alpha_i\ldots\alpha_n^{-1}$ as $A_{i+1}$. The edit distance $ed(A, B)$ is the minimum number of insertions, deletions, and substitutions of single letters needed to convert $A$ into $B$. The divergence is the observed number of errors per letter, $d := ed(A, B)/n$, whereas the error rate $e$ is the number of errors per letter applied to a sequence.

Alignment graph. Let state $(i, j)$ denote the subtask of aligning the prefix $A_{i+k}$ to the prefix $B_{j-l}$. The alignment graph (also called edit graph) $G(V, E)$ is a weighted directed graph with vertices $V = \{(i, j) \mid 0 \leq i \leq n, 0 \leq j \leq m\}$ corresponding to all states, and edges connecting subtasks: edge $(i, j) \to (i+1, j+1)$ has cost 0 if $a_i = b_j$ (match) and 1 otherwise (substitution), and edges $(i, j) \to (i, j+1)$ (deletion) and $(i, j) \to (i+1, j)$ (insertion) have cost 1. We denote the starting state $(0, 0)$ by $v_s$, the target state $(n, m)$ by $v_t$, and the distance between states $u$ and $v$ by $d(u, v)$. For brevity we write $f(i, j)$ instead of $f((i, j))$.

Paths and alignments. A path $\pi$ from $(i, j)$ to $(t', j')$ in the alignment graph $G$ corresponds to a (pairwise) alignment of the substrings $A_{i+k}$ and $B_{j-l}$ with cost $c_{\text{path}}(\pi)$. A shortest path $\pi^*$ from $v_s$ to $v_t$ corresponds to an optimal alignment, thus $c_{\text{path}}(\pi^*) = d(v_s, v_t)$. We write $g^*(u) := d(v_s, u)$ for the distance from the start to $u$ and $h^*(u) := d(u, v_t)$ for the distance from $u$ to the target.

Seeds and matches. We split the sequence $A$ into a set of consecutive non-overlapping substrings (seeds) $S = \{s_1, s_2, \ldots, s_k\}$, such that each seed $s_i = A_{k+i-k}$ has length $k$. After aligning the first $i$ letters of $A$, our heuristics will only depend on the remaining seeds $s_{i+1} = s_i \cup \{s_{i+1}\}$ contained in the suffix $A_{i+k}$. We denote the set of seeds between $u = (i, j)$ and $v = (t', j')$ by $S_{u,v} = S_{i+1} \cup \{s_{i+1} \mid i \leq k < j \leq k + i \}$ and an alignment of $s$ to a subsequence of $B$ by $s_v$. The alignments of seeds $s$ with sufficiently low cost (Sec. 3.1) form the set $M_s$ of matches.

Dijkstra and $A^*$. Dijkstra’s algorithm (Dijkstra, 1959) finds a shortest path from $v_s$ to $v_t$ by expanding (generating all successors) vertices in order of increasing distance $g^*(u)$ from the start. Each vertex to be expanded is chosen from a set of open vertices. The $A^*$ algorithm (Hart et al., 1968, 1972; Pearl, 1984), instead directs the search towards a target
by expanding vertices in order of increasing \( f(u) := g(u) + h(u) \), where 
\( h(u) \) is a heuristic function that estimates the distance \( h'(u) \) to the end 
and \( g(u) \) is the shortest length of a path from \( v_0 \) to \( u \) found so far. A 
heuristic is admissible if it is a lower bound on the remaining distance, 
\( h(u) \leq h'(u) \), which guarantees that \( A^* \) has found a shortest path as 
soon as it expands \( v_t \). Heuristic \( h_2 \) dominates (is more accurate than) 
another heuristic \( h_2 \) when \( h_1(u) \geq h_2(u) \) for all vertices \( u \). A dominant 
heuristic will usually, but not always (Holte, 2010), expand less vertices. 
Note that Dijkstra’s algorithm is equivalent to \( A^* \) using a heuristic that 
is always 0, and that both algorithms require non-negative edge costs. 
Our variant of the \( A^* \) algorithm is provided in App. A.1.

**Chains.** A state \( u = (i, j) \in V \) precedes a state \( v = (i', j') \in V \), 
denoted \( u \preceq v \) when \( i \leq i' \) and \( j \leq j' \). Similarly, a match \( m \) precedes 
a match \( m' \), denoted \( m \preceq m' \), when the end of \( m \) precedes the start of \( m' \). 
This makes the set of matches a partially ordered set. A state \( u \) precedes 
a match \( m \), denoted \( u \preceq m \), when it precedes the start of the match. A chain 
of matches is a (possibly empty) sequence of matches \( m_1 \preceq \ldots \preceq m_k \).

**Gap cost.** The number of indels to align substrings \( A_i \ldots A_j \) and \( B_i' \ldots B_j' \) 
is at least their difference in length: \( c_{\text{gap}}((i,j), (i',j')) := |(i'−i)−(j'−j)| \). For \( u \preceq v \), the gap cost satisfies the triangle inequality 
\( c_{\text{gap}}(u, v) \leq c_{\text{gap}}(u, w) + c_{\text{gap}}(w, v) \).

**Contours.** To efficiently calculate maximal chains of matches, contours are used. Given a set of matches \( M \), \( S(u) \) is the number of matches in the longest chain \( u \preceq m_1 \preceq \ldots \preceq m_k \) starting at \( u \). The function 
\( S(i, j) \) is non-increasing in both \( i \) and \( j \). Contours are the boundaries 
between regions of states with \( S(u) = \ell \) and \( S(u) < \ell \) (Fig. 3). Note 
that contour \( \ell \) is completely determined by the set of matches \( m \in M \) for 
which \( S(\text{start}(m)) = \ell \) (Hirschberg, 1977). Hunt and Szumianski (1977) 
give an algorithm to efficiently compute \( S \) when \( M \) is the set of single-letter 
matches between \( A \) and \( B \), and Deorowicz and Grabowski (2014) 
give an algorithm when \( M \) is the set of exact \( k \)-mer matches.

### 3 Methods

We formally define the general chaining seed heuristic (Sec. 3.1) that 
complements inexact matches, chaining, and gap costs (Fig. 2). Next, we 
introduce the match pruning (Sec. 3.2) improvement and integrate our 
\( A^* \) algorithm with the diagonal-transition optimization (App. A.2). We 
present a practical algorithm (Sec. 3.3), implementation (App. A.3) 
and proofs of correctness (App. B).

#### 3.1 General chaining seed heuristic

We introduce three heuristics for \( A^* \) that estimate the edit distance 
between a pair of suffixes. Each heuristic is an instance of a 
general chaining seed heuristic that we use to instantiate SH, CSH and 
GCSH.

**Definition 1** (General chaining seed heuristic). Given a set of matches 
\( M \), partial order \( \preceq \), and chaining cost \( \gamma \), the general chaining seed 
heuristic \( h_{\text{seed}}^{\gamma}(u) \) is the minimal sum of match costs and chaining costs 
over all \( \preceq \)-chains (indexing extends to \( m_0 := u \) and \( m_{i+1} := m_{\text{end}} \)): 

\[
h_{\text{seed}}^{\gamma}(u) := \min_{m_1 \in M} \sum_{i \in \mathbb{Z}} \left[ (m_i, m_{i+1}) + c_{\text{end}}(m_{i+1}) \right].
\]

| Heuristic          | Order | Chaining cost \( \gamma \) |
|--------------------|-------|-----------------------------|
| \( h_{\text{seed}}(u) \) | \( \preceq \) | \( c_{\text{seed}} \)          |
| \( h_{\text{CSH}}(u) \) | \( \preceq \) | \( c_{\text{seed}} \)          |
| \( h_{\text{GCSH}}(u) \) | \( \preceq \) | \( \max(c_{\text{gap}}, c_{\text{seed}}) \) |

Table 1. Definitions of our heuristic functions. SH orders the 
matches by \( i \) and uses only the seed cost. CSH orders the matches 
by both \( i \) and \( j \). GCSH additionally exploits the gap cost.

We instantiate our heuristics according to Table 1. Our admissibility 
proofs (App. B.1) are based on \( c_{\text{seed}} \) and \( \gamma \) being lower bounds on disjoint parts 
of the remaining path. Since the more complex \( h_{\text{GCSH}} \) dominates the 
other heuristics it usually expands fewer states.

**Theorem 1.** The seed heuristic \( h_{\text{seed}} \), the chaining seed heuristic \( h_{\text{CSH}}, \) 
and the gap-chaining seed heuristic \( h_{\text{GCSH}} \) are admissible. Furthermore, 
\( h_{\text{seed}}^\gamma(u) \leq h_{\text{CSH}}^\gamma(u) \leq h_{\text{GCSH}}^\gamma(u) \) for all states \( u \).
We are now ready to instantiate \( A^* \) with our admissible heuristics but we will first improve them and show how to compute them efficiently.

3.2 Match pruning

In order to reduce the number of states expanded by the \( A^* \) algorithm, we apply the multiple-path pruning observation: once a shortest path to a state has been found, no other path to this state could possibly improve the global shortest path (Poole and Mackworth, 2017). As soon as \( A^* \) expands the start or end of a match, we prune it, so that the heuristic in preceding states no longer benefits from the match, and they get deprioritized by \( A^* \). We define pruned variants of all our heuristics that ignore pruned matches:

\[ h(u) \] is a refinement of \( h(M) \) is a pruning heuristic version of \( h \).

The hat over the heuristic function \( \hat{h} \) denotes the implicit dependency on the progress of the \( A^* \), where at each step a different \( h(M) \) is used. Our modified \( A^* \) algorithm (App. A.1) works for pruning heuristics by ensuring that the \( f \)-value of a state is up-to-date before expanding it, and otherwise 

\[ \text{prune} \] it in the priority queue. Even though match pruning violates the admissibility of our heuristics for some vertices, we prove that \( A^* \) is still guaranteed to find a shortest path (App. B.2). To this end, we show that our pruning heuristics are weakly-admissible heuristics (Def. 7) in the sense that they are admissible on at least one path from \( v_2 \) to \( v_1 \).

**Theorem 2.** \( A^* \) with a weakly-admissible heuristic finds a shortest path.

**Theorem 3.** The pruning heuristics \( \hat{h}_{\text{h}}, \hat{h}_{\text{cS}}, \hat{h}_{\text{GCSh}} \) are weakly admissible.

Pruning will allow us to scale near-linearly with sequence length, without sacrificing optimality of the resulting alignment.

3.3 Computing the heuristic

We present an algorithm to efficiently compute our heuristics (pseudocode in App. A.4, worst-case asymptotic analysis in App. A.5). At a high level, we rephrase the minimization of costs (over paths) to a maximization of scores (over chains of matches). We initialize the heuristic by precomputing all seeds, matches, potentials and a contours data structure used to compute the maximum number of matches on a chain. During the \( A^* \) search, the heuristic is evaluated in all explored states, and the contours are updated whenever a match gets pruned.

**Scores.** The score of a match \( m \) is \( \text{score}(m):=r-c_{au}(m) \) and is always positive. The score of a \( \text{path} \) \( m_1 \ldots m_p \) is the sum of the scores of the matches in the chain. We define the chain score of a match as

\[ S_p(m) := \max_{m_0 \in \text{path}} \left\{ \text{score}(m_0) + \ldots + \text{score}(m_p) \right\}. \]  

(1)

Since \( S_p \) is a partial order, \( S_p \) can be computed with base case \( S_p(m_u) = 0 \) and the recursion

\[ S_p(m) = \text{score}(m) + \max_{m_0 \in S_p(m) \cup S_p(m')} S_p(m'). \]  

(2)

We also define the chain score of a state \( u \) as the maximum chain score over succeeding matches \( m: S_p(u) = \max_{m_0 \in \text{path}} S_p(m), \) so that Eq. (2) can be rewritten as \( S_p(m) = \text{score}(m) + S_p(\text{end}(m)) \).

The following theorem allows us to rephrase the heuristic in terms of potentials and scores for heuristics that use \( \gamma = c_{seed} \) and respect the order of the seeds, which is the case for \( h_{\text{h}} \) and \( h_{\text{cS}} \) (proof in App. B.3):

**Theorem 4.** \( h_{\text{pruned}}(u) = P(u) - S_p(u) \) for any partial order \( S_p \) that is a refinement of \( S_1 \) (i.e. any \( S_p \) must imply \( S_1 \)).

Layers and contours. We compute \( h_{\text{h}} \) and \( h_{\text{cS}} \) efficiently using contours. Let layer \( \mathcal{L}_0 \) be the set of states \( u \) with score \( S_p(u) \geq \ell \), so that \( \mathcal{L}_0 \subseteq \mathcal{L}_{\ell-1} \). The 0th contour is the boundary of \( \mathcal{L}_0 \) (Fig. 3). Layer \( \mathcal{L}_0 \) contains exactly those states that precede a match \( m \) with score \( \ell \leq S_p(m) < \ell + r \) (Lemma 5 in App. B.2).

**Computing \( S_p(u) \).** This last observation inspires our algorithm for computing chain scores. For each layer \( \mathcal{L}_0 \), we store the set \( \text{match}(\ell) \) of matches having score \( \ell: L[\ell] = \{ m \in \mathcal{M} | S_p(m) = \ell \} \). The score \( S_p(u) \) is then the highest \( \ell \) such that layer \( L[\ell] \) contains a match \( m \) reachable from \( u \) (\( S_p(m) \)). From Lemma 5 we know that \( S_p(u) \geq \ell \) if and only if one of the layers \( L[\ell'] \) for \( \ell' \in \{ \ell, \ell + r \} \) contains a match preceded by \( u \). We use this to compute \( S_p(u) \) using a binary search over the layers \( \ell \). We initialize \( L[0] = \{ m_u \} \) (\( m_u \) is a fictive match at the target \( v_t \)), sort all matches in \( \mathcal{M} \) by \( S_p \), and process them in decreasing order (from the target to the start). After computing \( S_p(\text{end}(m)) \), we add to \( \text{match} \) \( S_p(m) = \text{score}(m) + S_p(\text{end}(m)) \). Matches that do not precede the target \( (\text{start}(m) \neq m_u) \) are ignored.

**Pruning matches from \( \mathcal{L}_0 \).** When pruning matches starting or ending in \( \text{state} \) \( u \) in layer \( \mathcal{L}_0 \), we remove all matches that start at \( u \) from layers \( L[\ell_u] \) to \( L[\ell_v] \), and all matches starting in \( v \) and ending in \( u \) from layers \( L[\ell_v] \) to \( L[\ell_u] \).

Pruning a match may change \( S_p \) in layers above \( \ell_u \), so we update them after each prune. We iterate over increasing \( \ell \) starting at \( \ell_u + 1 \) and recompute \( \ell' := S_p(m) \leq \ell \) for all matches in \( L[\ell] \). If \( \ell' \neq \ell \), we move \( m \) from \( L[\ell] \) to \( L[\ell'] \). We stop iterating when either \( \ell \) consecutive layers were left unchanged, or when all matches in \( \mathcal{L}_0 \) are pruned. In the former case, no further scores can change, and in the latter case, \( S_p \) decreases by \( \ell - \ell' \) for all matches with score \( \geq \ell \). We remove the emptied layers \( L[\ell'] \) to \( L[\ell] \) so that all higher layers shift down by \( \ell - \ell' \).

**SH.** Due to the simple structure of the seed heuristic, we also simplify its computation by only storing the start of each layer and the number of matches in each layer, as opposed to the full set of matches.

**GCSh.** Thm. 4 does not apply to gap-chaining seed heuristic since it uses chaining cost \( \gamma = c_{gap}(u, v) \) which is different from \( c_{seed}(u, v) \). It turns out that in this new setting it is never optimal to chain two matches if the gap cost between them is higher than the seed.
cost. Intuitively, it is better to miss a match than to incur additional gapcost to include it. We capture this constraint by introducing a transformation $T$ such that $u \preceq_T v$ holds if and only if $c_{\text{seed}}(u,v) \geq c_{\text{gap}}(u,v)$, as shown in App. B.4. Using an additional consistency constraint on the set of matches we can compute $h_{\text{GC}}^M$ via $S_T$ as before.

Definition 3 (Consistent matches). A set of matches $M$ is consistent when for each $m \in M$ (from $(i,j)$ to $(i',j')$) with $\text{score}(m)>1$, for each adjacent pair of existing states $((i,j)\preceq(i,j'))$ and $((i,j),(i',j'))$, there is an adjacent match with corresponding start and end, and score at least $\text{score}(m)-1$.

This condition means that for $r=2$, each exact match must be adjacent to four (or less around the edges of the graph) inexact matches starting or ending in the same state. Since we find all matches $m$ with $c_m(m) \leq r$, our initial set of matches is consistent. To preserve consistency, we do not prune matches if that would break the consistency of $M$.

Definition 4 (Gap transformation). The partial order $\preceq_T$ on states is induced by comparing both coordinates after the gap transformation

$$T: \ (i,j) \mapsto (i-j - P(i,j), j-i - P(i,j))$$

Theorem 5. Given a consistent set of matches $M$, the gap-chaining seed heuristic can be computed using scores in the transformed domain:

$$h_{\text{GC}}^M(u) = \begin{cases} P(u) - S_T(u) & \text{if } u \preceq_T v, \\
\epsilon_{\text{gap}}(u,v) & \text{if } u \npreceq_T v. \end{cases}$$

Using the transformation of the match coordinates, we can now reduce $c_{\text{gap}}$ to $c_{\text{seed}}$ and efficiently compute GCSh in any explored state.

### 4 Results

Our algorithm is implemented in the aligner A*PA\footnote{github.com/RagnarGrootKoerkamp/astar-pairwise-aligner (tag evals)} in Rust. We compare it with state of the art exact aligners on synthetic (Sec. 4.2) and human (Sec. 4.3) data\footnote{github.com/pairwise-alignment/pa-bench/releases/tag/datasets}. We justify our heuristics and optimizations by comparing their scaling and performance (Sec. 4.4).

#### 4.1 Setup

**Synthetic data.** Our synthetic datasets are parameterized by sequence length $n$, induced error rate $e$, and total number of basepairs $N$, resulting in $N/n$ sequence pairs. The first sequence in each pair is uniform-random from $\Sigma^n$. The second is generated by sequentially applying $[e-n]$ edit operations (insertions, deletions, and substitutions with equal $1/3$ probability) to the first sequence. Introduced errors can cancel each other, making the divergence $d$ between the sequences less than $e$. Induced error rates of $1\%$, $5\%$, $10\%$, and $15\%$ correspond to divergences of $0.9\%$, $4.3\%$, $8.2\%$, and $11.7\%$, which we refer to as $1\%$, $4\%$, $8\%$, and $12\%$.

**Human data.** We use two datasets of ultra-long Oxford Nanopore Technologies (ONT) reads of the human genome: one without and one with genetic variation. All reads are $500-1100\text{ kbp}$ long, with mean divergence around $7\%$. The average length of the longest gap in the alignment is $0.1\text{kbp}$ for ONT reads, and $2\text{kbp}$ for ONT reads with genetic variation (detailed statistics in App. C.5). The reference genome is CHM13 (v1.1)\footnote{github.com/RagnarGrootKoerkamp/astar-pairwise-aligner (tag evals)}. The reads used for each dataset are:

- **ONT**: 50 reads sampled from those used to assemble CHM13.
- **ONT with genetic variation**: 48 reads from another human (Bowden et al., 2019), as used in the BiWFA paper (Marco-Sola et al., 2023).

**Algorithms and aligners.** We compare SH, CSH, and GCSh (all with pruning) as implemented in A*PA to the state-of-the-art exact aligners BiWFA and EDLib. We also compare to Dijkstra’s algorithm and A* with previously introduced heuristics (gap cost and character frequencies of Hadlock (1988b)), and SH without pruning of Ivanov et al. (2022)). We exclude SEQAN and PARASAIL since they are outperformed by WFA and EDLib (Marco-Sola et al., 2021; Sošić and Šikić, 2017). We run all aligners with unit edit costs with traceback enabled.

**A*PA parameters.** Inexact matches ($r=2$) and short seeds (low $k$) increase the accuracy of GCSh for divergent sequences, thus reducing the number of expanded states. On the other hand, shorter seeds have more matches, slowing down precomputation and contour updates. A parameter grid search on synthetic data (App. C.1) shows that the runtime is generally insensitive to $k$ as long as $k$ is high enough to avoid too many spurious matches ($k \gg \log_4 n$), and the potential is sufficiently larger than edit distance ($k \ll r/d$). For $d=4\%$, exact matches lead to faster runtimes, while $d=12\%$ requires $r=2$ and $k < 2d = 16.7$. We fix $k = 15$ throughout the evaluations since this is a good choice for both synthetic and human data.

**Execution.** We use PA-BENCH on Arch Linux on an Intel Core i7-10750H processor with 64 GB of memory and 6 cores, without
hyper-threading, frequency boost, and CPU power saving features. We fix the CPU frequency to 2.6GHz, limit the memory usage to 32 Gib, and run 1 single-threaded job at a time with niceness ~20.

**Measurements.** PARChEFS first reads the dataset from disk and then measures the wall-clock time and increase in memory usage of each aligner. Plots and tables refer to the average alignment time per aligned pair, and for A*PA include the time to build the heuristic. Best-fit polynomials are calculated via a linear fit in the log-log domain using the least squares method.

### 4.2 Scaling on synthetic data

**Runtime scaling with length.** We compare our A* heuristics with Edlib, BiWFA, and other heuristics in terms of runtime scaling with n and d (Fig. 4, extended comparison in App. C.2). As theoretically predicted, Edlib and BiWFA scale quadratically. For small edit distance, Edlib is subquadratic due to the bit-parallel optimization. Dijkstra’s A* with the gap heuristic, character frequency heuristic (Hadlock, 1988b), or original seed heuristic (Ivanov et al., 2022) all scale quadratically. The empirical scaling of A*PA is subquadratic for d≤12 and n≤10^7, making it the fastest aligner for long sequences (n>30k bp). For low divergence (d≤4%) even the simplest SH scales near-linearly with length (best fit n<sup>1.06</sup> for n≤10^7). For high divergence (d=12%) we need inexact matches, and the runtime of SH sharply degrades for long sequences (n=10^7 bp) due to spurious matches. This is countered by chaining the matches in CSH and GCSh, which expand linearly many states (App. C.3). GCSh with DT is not exactly linear due to high memory usage and state reordering (App. C.7 shows the time spent on parts of the algorithm).

**Runtime scaling with divergence.** Fig. 4c shows that A*PA has near constant runtime in d as long as the edit distance is sufficiently less than the heuristic potential (i.e. d < r/k). In this regime, A*PA is faster than both Edlib (linear in d) and BiWFA (quadratic in d). For 1 ≤ d ≤ 6%, exact matches have less overhead than inexact matches, while BiWFA is fastest for d ≤ 1%. A*PA becomes linear in d for d ≥ r/k (App. C.4).

**Performance.** A*PA with SH with DT is >500× faster than Edlib and BiWFA for d=1% and n=10^7 (Fig. 4a). For n=10^6 and d≤12%, memory usage is less than 500 MB for all heuristics (App. C.6).

### 4.3 Speedup on human data

We compare runtime (Fig. 5, App. C.7), and memory usage (App. C.6) on human data. We configure A*PA to prune matches only when expanding their start (not their end), leaving some matches on the optimal path unpruned and speeding up contour updates. The runtime of A*PA (GCSh with DT) on ONT reads is less than Edlib and BiWFA in all quartiles, with the median being >3x faster. However, the runtime of A*PA grows rapidly when d≥10%, so we set a time limit of 100 seconds per read, causing 6 alignments to time out. In real-world applications, the user would either only get results for a subset of alignments, or could use a different tool to align divergent sequences. With genetic variation, A*PA is 1.7x faster than Edlib and BiWFA in median. Low-divergence alignments are faster than Edlib, while high-divergence alignments are slower (3 sequences with d≥10% time out) because of expanding quadratically many states in complex regions (App. C.8). Since slow alignments dominate the total runtime, Edlib has a lower mean runtime.

### 4.4 Effect of pruning, inexact matches, chaining, and DT

We visualize our techniques on a complex alignment in App. C.10.

**SH with pruning enables near-linear runtime.** Figure 4a shows that the addition of match pruning changes the quadratic runtime of SH without pruning to near-linear, giving multiple orders of magnitude speedup.

**Inexact matches cope with higher divergence.** Inexact matches double the heuristic potential, thereby almost doubling the divergence up to which A*PA is fast (Fig. 4c). This comes at the cost of a slower precomputation to find all matches.

**Chaining copes with spurious matches.** While CSH improves on SH for some very slow alignments (Fig. 5), more often the overhead of computing contours makes it slower than SH.

**Gap-chaining copes with indels.** GCSh is significantly and consistently faster than SH and CSH on human data, especially for slow alignments (Fig. 5). GCSh has less overhead over SH than CSH, due to filtering out matches m ≤ v.<

**Diagonal transition speeds up quadratic search.** DT significantly reduces the number of expanded states when the A* search is quadratic (Fig. 4a and App. C.4). In particular, this results in a big speedup when aligning genetic variation containing big indels (Fig. 5).

**CSH, GCSh, and DT only have a small impact on the uniform synthetic data, where usually either the SH is sufficiently accurate for the entire alignment and runtime is near-linear (d < r/k), or even GCSh is not strong enough and runtime is quadratic (d ≥ r/k). On human data however, containing longer indels and small regions of quadratic search, the additional accuracy of GCSh and the reduced number of states explored by DT provide a significant speedup (App. C.10).**

### 5 Discussion

**Seeds are necessary, matches are optional.** The seed heuristic uses the lack of matches to penalize alignments. Given the admissibility of our heuristics, the more seeds without matches, the higher the penalty for alignments and the easier it is to dismiss suboptimal ones. In the extreme, not having any matches can be sufficient for finding an optimal alignment in linear time (App. C.9).
**Modes:** Near-linear and quadratic. The \( A^* \) algorithm with a seed heuristic has two modes of operation that we call near-linear and quadratic. In the near-linear mode \( A^*PA \) expands few vertices because the heuristic successfully penalizes all edits between the sequences. When the divergence is larger than what the heuristic can handle, every edit that is not penalized by the heuristic increases the explored band, leading to a quadratic exploration similar to Dijkstra.

**Limitations.**

1. **Quadratic scaling.** Complex data can trigger a quadratic (Dijkstra-like) search, which nullifies the benefits of \( A^* \) (Appendices C.8 and C.10). Regions with high divergence (\( d \leq 10\% \)), such as high error rate or long indels, exceed the heuristic potential to direct the search and make the exploration quadratic. Low-complexity regions (e.g. with repeats) result in a quadratic number of matches which also take quadratic time.

2. **Computational overhead of \( A^* \).** Computing states sequentially (as in EDLIB, BiWFA) is orders of magnitude faster than computing them in random order (as in Dijkstra, \( A^* \)). \( A^*PA \) outperforms EDLIB and BiWFA (Fig. 4a) when the sequences are long enough for the linear-scaling to have an effect (\( n \geq 30 \text{kbp} \)), and there are enough errors (\( d > 1\% \)) to trigger the quadratic behaviour of BiWFA.

**Future work.**

1. **Performance.** We are working on a DP-based version of \( A^*PA \) that applies computational volumes (Spouge, 1989, 1991), block-based computations (Liu and Steinegger, 2023), and a SIMD version of EDLIB’s bit-parallelization (Meyers, 1999). This has already shown 10x additional speedup on the human data sets and is less sensitive to the input parameters. Independently, the number of matches could be reduced by using variable seed lengths and skipping seeds with many matches.

2. **Generalizations.** Our chaining seed heuristic could be generalized to non-unit and affine costs, and to semi-global alignment. Cost models that better correspond to the data can speed up the alignment.

3. **Relaxations.** At the expense of optimality guarantees, inadmissible heuristics could speed up \( A^* \) considerably. Another possible relaxation would be to validate the optimality of a given alignment instead of computing it from scratch.

4. **Analysis.** The near-linear scaling with length of \( A^* \) is not asymptotic and requires a more thorough theoretical analysis (Medvedev, 2022a).

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A Pseudocode

A.1 A* algorithm for match pruning

We present our variant of A* (Hart et al., 1968) that supports match pruning (Algorithm 1). All computed values of \( g \) are stored in a hash map, and all open states are stored in a bucket queue of tuples \((u, g(u), f(u))\) ordered by increasing \( f \). Line 14 prunes (removes) a match and thereby increases some heuristic values before that match. As a result, some \( f \)-values in the priority queue may become outdated. Line 11 solves this problem by checking if the \( f \)-value of the state about to be expanded was changed, and if so, line 12 pushes the updated state to the queue, and proceeds by choosing a next best state. This way, we guarantee that the expanded state has minimal updated \( f \). To reconstruct the best alignment we traceback from the target state using the hash map \( g \) (not shown).

Algorithm 1 A* algorithm with pruning.

| Line | Code                                                                                          |
|------|------------------------------------------------------------------------------------------------|
| 1    | **Input:** Sequences \( A \) and \( B \) and pruning heuristic \( h \)                        |
| 2    | **Output:** Edit distance between \( A \) and \( B \)                                       |
| 3    | **function** \( \text{ASTAR}(A, B, h) \)                                                  |
| 4    | \( g(v_i) \leftarrow 0 \) > Hashmap of distances; default \( +\infty \)                     |
| 5    | \( q \leftarrow \text{BucketQueue()} \) > Bucket queue of open states                      |
| 6    | \( q.push((v_s, g=0, f=0)) \)                                                             |
| 7    | repeat                                                                                       |
| 8    | \((u, g_u, f_u) \leftarrow q.pop()\) > Pop \( u \) with minimal \( f \)                     |
| 9    | if \( g_u > g(u) \) then \( u \) was already expanded                                       |
| 10   | continue                                                                                 |
| 11   | else if \( f_u < g(u) + h(u) \) then \( h(u) \) has increased                             |
| 12   | \( q.push((u, g_u, (g(u) + h(u)))) \) > Reorder \( u \)                                    |
| 13   | else \( u \) \(\text{/} Expand \( u \)                                                  |
| 14   | \( \text{Prune}(u) \)                                                                      |
| 15   | for successors \( v \) of \( u \) do                                                         |
| 16   | \( g_v \leftarrow g_u + d(u, v) \)                                                           |
| 17   | \( v \leftarrow \text{Extend}(v) \) > Greedy matching within seed                           |
| 18   | if \( g_v < g(v) \) then \( v \) is expanded                                               |
| 19   | \( g(v) \leftarrow g_v \)                                                                  |
| 20   | \( q.push((v, g_v, g_v + h(v))) \)                                                           |
| 21   | until \( v \) is expanded                                                                 |
| 22   | return \( g(v_t) \)                                                                        |

A.2 Diagonal transition for A*

For a given distance \( g \), the diagonal transition method only considers the farthest-reaching (f.r.) state \( u=(i,j) \) on each diagonal \( k=i-j \) at distance \( g \). We use \( F_{g,k} := i+j \) to indicate the antidiagonal\(^4\) of the farthest reaching state. Let \( X_{g,k} \) be the farthest state on diagonal \( k \) adjacent to a state at distance \( g-1 \), which is then extended into \( F_{g,k} \) by following as many matches as possible. The edit distance is the lowest \( g \) such that \( F_{g,n+m} \geq n + m \), and we have the recursion

\[
X_{g,k} := \max(F_{g-1,k+1}, F_{g-1,k+2}, F_{g-1,k+1+1}), \tag{3}
\]

\[
F_{g,k} = X_{g,k} + \text{LCP}(A_{g-k}(k)/2, B_{g-k}(k)/2). \tag{4}
\]

The base case is \( X_{0,0} = 0 \) with default value \( F_{g,k} = -\infty \) for \( k > |g| \), and LCP is the length of the longest common prefix of two strings. Each edge in a traceback path is either a match created by an extension (4), or a mismatch of the farthest-reaching \( A^* \) state using the hash map \( (\text{not shown}) \).

Algorithm 2 A*–DT algorithm with match pruning.

| Line | Code                                                                                          |
|------|------------------------------------------------------------------------------------------------|
| 1    | **Input:** Sequences \( A \) and \( B \) and pruning heuristic \( h \)                        |
| 2    | **Output:** Edit distance between \( A \) and \( B \)                                       |
| 3    | **function** \( \text{ASTAR-DT}(A, B, h) \)                                                |
| 4    | \( F_{0,0} \leftarrow 0 \) > Hashmap of f.r. point per \( g \) and \( k \); default \( -\infty \) |
| 5    | \( q \leftarrow \text{BucketQueue()} \) > Bucket queue of open states                      |
| 6    | \( q.push((u, g=0, f=0)) \)                                                               |
| 7    | repeat                                                                                       |
| 8    | \((u, g_u, f_u) \leftarrow q.pop()\) > Pop \( u \) with minimal \( f \)                     |
| 9    | if \( g_u > g(u) \) then \( u \) was already expanded                                       |
| 10   | continue                                                                                 |
| 11   | else if \( f_u < g(u) + h(u) \) then \( h(u) \) has increased                             |
| 12   | \( q.push((u, g_u, (g(u) + h(u)))) \) > Reorder \( u \)                                    |
| 13   | else \( u \) \(\text{/} Expand \( u \)                                                  |
| 14   | \( \text{Prune}(u) \)                                                                      |
| 15   | for successors \( v \) of \( u \) do                                                         |
| 16   | \( g_v \leftarrow g_u + d(u, v) \)                                                           |
| 17   | \( v \leftarrow \text{Extend}(v) \) > Greedy matching within seed                           |
| 18   | if \( g_v < g(v) \) then \( v \) is expanded                                               |
| 19   | \( g(v) \leftarrow g_v \)                                                                  |
| 20   | \( q.push((v, g_v, g_v + h(v))) \)                                                           |
| 21   | until \( v \) is expanded                                                                 |
| 22   | return \( g(v_t) \)                                                                        |

Implementation. In Algorithm 2 we further modify the A* algorithm to only consider f.r. paths. We replace the map \( g \) that tracks the best distance by a map \( F_{g,k} \) that tracks f.r. states (lines 4, 18, and 19). Instead of \( g(u) \) decreasing over time, we now ensure that \( F_{g,k} \) increases over time. Each time a state \( u \) is opened or expanded, the check whether \( g(u) \) decreases is replaced by a check whether \( F_{g,k} \) increases (line 9). This causes the search to skip states that are known to not be farthest reaching. The proof of correctness (Thm. 2) still applies.

Alternatively, it is also possible to implement A* directly in the diagonal-transition state-space by pushing states \( F_{g,k} \) to the priority queue, but for simplicity we keep the similarity with the original A*.

A.3 Implementation notes

Here we list implementation details on performance and correctness.

**Bucket queue.** We use a hashmap to store all computed values of \( g \) in the A* algorithm. Since the edit costs are bounded integers, we implement the priority queue using a bucket queue (Hitchner, 1968; Dial, 1969; Bertsekas, 1991). Unlike heaps, this data structure has amortized constant time push and pop operations since the value difference between consecutive pop operations is bounded.

**Greedy matching of letters.** From a state \((i,j)\) where \( a_i = b_j \), it is sufficient to only consider the matching edge to \((i+1, j+1)\) (Allison, 1992; Ivanov et al., 2020), and ignore the insertion and deletion edges to \((i, j+1)\) and \((i+1, j)\). During alignment, we greedily match as many letters as possible within the current seed before inserting only the last open state in the priority queue, but we do not cross seed boundaries in order to not interfere with match pruning. This optimization is superseded by the DT-optimization. We include greedyly matched states in the reported number of expanded states.

**Priority queue offset.** Pruning the last remaining match in a layer may cause an increase of the heuristic in all states preceding the start \( u \) of the match. This invalidates \( f \) values in the priority queue and causes reordering. We skip most of the update operations by storing a global

\(^4\)Previous works indicate the column \( i \) of \( u \), but using the antidiagonal \( i+j \) keeps the symmetry between insertions and deletions.
offset to the $f$-values in the priority queue, which we update when all states in the priority queue precede $u$.

**Split vector for layers.** Pruning a match may trigger the removal of one or more layers of matches in $L$, and the shifting down of higher layers. To efficiently remove layers, we use a split vector data structure consisting of two stacks. In the first stack we store the layers before the last deleted layer, and in the second stack the remaining layers in reverse order. Before deleting a layer, we move layers from the top of one of the stacks to the top of the other, until the layer to be deleted is at the top of one of the stacks. Removing layers in decreasing order of $τ$ takes linear total time.

**Binary search speedup.** Instead of using binary search to determine the layer/score $S_p(u)$ (Algorithm 3), we first try a linear search around either the score of the parent of $u$ or a known previous score at $u$. This linear search usually finds the correct layer in a few iterations, or otherwise we fall back to binary search.

In practice, most pruning happens near the tip of the search, and the number of layers between the start $v_s$ and an open state $u$ rarely changes. Thus, to account for changing scores, we store a hint of value $S_p(v_s) - S_p(u)$ in a hashmap and start the search at $S_p(v_s) - \text{hint}$.

**Code correctness.** Our implementation $A^*PA$ is written in Rust, contains many assertions testing e.g. the correctness of our $A^*$ and layers data structure implementation, and is tested for correctness and performance on randomly-generated sequences. Correctness is checked against simpler algorithms (Needleman-Wunsch) and other aligners (EDLIB, BiWFA).

### A.4 Computation of the heuristic

Algorithm 3 shows how the heuristic is initialized, how $S_p(u)$ and $h(u)$ are computed, and how matches are pruned.

### A.5 Worst-case runtime asymptotics

Our algorithms optimize for the average case performance. Nevertheless, we discuss the worst-case time of each part of the algorithm. To analyze the worst-case asymptotics of $A^*PA$, let $M = O(n^2)$ be the number of (inexact) matches, and let $E = O(n^2)$ be the number of expanded states. Then the asymptotical runtime of our algorithm breaks down as:

1. Finding all matches takes $O(n + M)$ time.
2. Building the contours datastructure takes $O(M \log M)$ time.
3. Each expanded state is pushed onto and popped from the priority queue ($O(1)$ using a bucket queue), looked up in a hashmap ($O(1)$), and has its neighbours explored ($O(1)$), for $O(E)$ total.
4. Evaluating the heuristic requires a binary search over contours, which is bounded by $O(n \log n)$, since we test at most $\log n$ contours each containing at most $n$ matches. (This could be improved to $O(\log^2 n)$ by storing each contour as an ordered set, but it is not usually a bottleneck.)
5. Pruning a match requires a dictionary lookup and change for $O(1)$. Updating contours after pruning requires iterating over higher contours until no more changes are triggered. This could take $O(M)$ for each match that is pruned, leading to a naive upper bound of $O(M^2)$.
6. Reordering states takes $O(En)$, since the heuristic in each expanded state can decrease at most $n$ times.
7. The traceback requires $O(n)$ time.

Taking this together, we obtain $O(n + M \log M + En \log n + M^2 + En)$. When the number of matches $M$ is $\Theta(n^2)$, the updating of contours has the worst upper bound and the asymptotics becomes $O(n^4)$. Whether this upper bound can be reached in practice or whether a tighter bound exists needs further investigation. Even if the worst-case was $\Theta(n^2)$, in practice our algorithm only runs efficiently in regimes well below this bound.

**Algorithm 3 Computation of the heuristic.**

1: Input: Sequences $A$ and $B$
2: Output: Heuristic $h$
3: function INITIALIZEHEURISTIC($A, B, k$) ▷ Seeds
4: $S \leftarrow$ Non-overlapping $k$-mers of $A$
5: $H \leftarrow$ Map of all $k$-mers (and $k+1$-mers for $r=2$) of $B$
6: $M_s \leftarrow \cup_{(u,v') \in H[s']} H[s']$ ▷ Seed matches
7: $M \leftarrow \cup_{s \in S} M_s$ ▷ Matches
8: $P(i) \rightarrow r \cdot |S_{\geq i}|$ for all $i \in [0, |A|]$ ▷ Potentials
9: $L[0] = \{m_w\}$ ▷ Layers
10: for $m \in M$ in decreasing order of $s_p$ do
11: $\ell \leftarrow$ score($m$) + $S_p$(end($m$))
12: if $m \geq m_w$ then ▷ If $m$ precedes the target
13: $L[\ell].push(m)$
14: function $S_p(u)$
15: function TEST($\ell$) ▷ Binary search the highest $\ell$
16: for $\ell' \in [\ell, \ell + r]$ do
17: if $u \geq m$ for some $m \in L[\ell']$ then
18: return True
19: return False
20: return $\max(\ell \mid \text{TEST}(\ell))$
21: function $h_s(u)$ ▷ SH
22: return $P(u) - S_s(u)$
23: function $h_C(u)$ ▷ CSH
24: return $P(u) - S_s(u)$
25: function $h_{\text{GCD}}(u)$ ▷ GCSH
26: return $\max(P(u) - S_T(u), c_{\text{gap}}(u, v))$
27: function $\text{PRUNE}(u)$
28: $\ell_u \leftarrow S_p(u)$
29: for all $m \in M$: start($m$) = $u$ or end($m$) = $u$ do
30: if $h \neq h_{\text{GCD}}$ or $M_r(m)$ is consistent then
31: Remove $m$ from $M$
32: for $\ell \in [S_p(\text{start}(m)) - r + 1, S_p(\text{start}(m))]$ do
33: Remove $m$ from $L[\ell]$ if present
34: for $\ell' \in [\ell_u + 1, S_p(0, 0)]$ do
35: for all $m \in L[\ell']$ do
36: $\ell' \leftarrow S_p(m)$
37: if $\ell' \neq \ell$ then
38: Move $m$ from $L[\ell]$ to $L[\ell']$
39: if $r$ consecutive layers unchanged then
40: return
41: if $r-1 + \ell - \ell'$ consecutive layers shifted exactly $\ell - \ell'$ then
42: Remove empty layers $L[\ell' + 1], \ldots, L[\ell']$
43: return

Appendix C.1 discusses the effect of $r$ and $k$ on the number of expanded states when aligning synthetic data, and Appendix C.4 discusses the effect of the divergence $d$ on the number of expanded states.
B Proofs

B.1 Admissibility

Our heuristics are not consistent, but we show that a weaker variant holds for states at the start of a seed.

Definition 5 (Start of seed). A state \((i, j)\) is at the start of some seed when \(i\) is a multiple of the seed length \(k\), or when \(i = n\).

Lemma 1 (Weak triangle inequality). For states \(u, v, w\) with \(v\) and \(w\) at the starts of some seeds, all \(\gamma \in \{c_{\text{seed}}, c_{\text{gap}}, c_{\text{gs}}\}\) satisfy

\[
\gamma(u, v) + \gamma(v, w) \geq \gamma(u, w).
\]

Proof. Both \(v\) and \(w\) are at the start of some seeds, so for \(\gamma = c_{\text{seed}}\) we have the equality \(c_{\text{seed}}(u, w) = c_{\text{seed}}(u, v) + c_{\text{seed}}(v, w)\).

For \(\gamma = c_{\text{gap}}\),

\[
c_{\text{gap}}((i, j), (i', j')) + c_{\text{gap}}((i', j'), (i'', j'')) = (|i' - i| - |j' - j|) + (|i'' - i'| - |j'' - j'|) + c_{\text{gap}}((i, j'), (i'', j'')))\]

and lastly, for \(\gamma = c_{\text{gs}}\),

\[
c_{\text{gs}}(u, v) + c_{\text{gs}}(v, w) = \max(c_{\text{gap}}(u, v), c_{\text{seed}}(u, v)) + \max(c_{\text{gap}}(v, w), c_{\text{seed}}(v, w)) \geq \max(c_{\text{gap}}(u, v) + c_{\text{gap}}(v, w), c_{\text{seed}}(u, v) + c_{\text{seed}}(v, w)) \geq \max(c_{\text{gap}}(u, w), c_{\text{seed}}(u, w)) = c_{\text{gs}}(u, w).\]

Lemma 2 (Weak consistency). Let \(h \in \{h^M_s, h^M_c, h^M_{\text{gcd}}\}\) be a heuristic with partial order \(\succeq_p\), and let \(u \succeq_p v\) be states with \(v\) at the start of a seed. When there is a shortest path \(\pi\) from \(u\) to \(v\) such that \(M\) contains all matches of cost less than \(v\) on \(\pi\), it holds that \(h(u) \leq d(u, v) + h(v)\).

Proof. The path \(\pi\) covers each seed in \(S_u, \ldots, v\) that must to be fully aligned between \(u\) and \(v\). Since the seeds do not overlap, their shortest alignments \(\pi_u^*\) do not have overlapping edges. Let \(u \leq m_1 \succeq_p \cdots \succeq_p m_t \succeq_p v\) be the chain of matches \(m_i \in M\) corresponding to those \(\pi_u^*\) of cost less than \(v\) (Fig. 6). Since the matches and the paths between them are disjoint, \(c_{\text{path}}(\pi_u^*)\) is at least the cost of the matches \(c_{\text{m}}(m_{i+1}, i) = d(\text{start}(m_{i+1}), \text{end}(m_{i+1}))\) plus the cost to chain these matches \(\gamma(\text{end}(m_i), \text{start}(m_{i+1})) \leq d(\text{end}(m_i), \text{start}(m_{i+1}))\).

Putting this together:

\[
\gamma(u, m_1) + \cdots + c_{\text{m}}(m_i) + \gamma(m_{i+1}) \leq d(u, \text{start}(m_1)) + d(\text{start}(m_1), \text{end}(m_1)) + \cdots + d(\text{end}(m_i), \text{start}(m_{i+1})).
\]

Now let \(v \succeq_p m_{t+1} \succeq_p \cdots \succeq_p m_t \succeq_p v\) be a chain of matches minimizing \(h(v)\) (Def. 1) with \(u := \text{start}(m_{i+1})\). This chain also minimizes \(h(w)\) and thus \(h(v) = \gamma(v, w) + h(w)\). We can now bound the cost of the joined chain from \(u\) to \(v\) and from \(v\) to \(e\) and get our result via \(\gamma(m_{i, v}) \leq \gamma(m_{i, v}) + \gamma(v, w)\) (Lemma 1)

\[
h(u) \leq \gamma(u, m_1) + \cdots + \gamma(m_{i+1}) + c_{\text{m}}(m_{i+1}) + \cdots + \gamma(m_t, v) + h(w) \leq \gamma(u, m_1) + \cdots + \gamma(m_t, v) + \gamma(v, w) + (h(w) - \gamma(v, w)) \leq d(u, v) + h(v).\]

\[\square\]

Theorem 1. The seed heuristic \(h_s\), the chaining seed heuristic \(h_c\), and the gap-chaining seed heuristic \(h_{\text{gcd}}\) are admissible. Furthermore, \(h_s^M(u) \leq h_c^M(u) \leq h_{\text{gcd}}^M(u)\) for all states \(u\).

Proof. We will prove \(h_s^M(u) \leq h_c^M(u) \leq h_{\text{gcd}}^M(u) \leq h^*(u)\), which implies the admissibility of all three heuristics.

(1) Note that \(u \succeq v\) implies \(u \succeq v\) and hence any \(s\)-chain is also a \(s\)-chain. A minimum over the superset of \(s\)-chains is at most the minimum of the subset of \(s\)-chains, and hence \(h_s^M = h_{c,\text{seed}}^M \leq h_{\text{gcd}}^M = h_{c,\text{gcd}}^M\).

(2) The only difference between \(h_{\text{gcd}}^M\) and \(h_{c,\text{seed}}^M\) is that the former uses \(c_{\text{seed}}\) and the latter uses the gap-seed cost \(c_{\text{gap}} := \max(c_{\text{gap}}, c_{\text{seed}})\). Since \(c_{\text{seed}} \leq c_{\text{gs}}\), we have \(h_{\text{gcd}}^M = h_{c,\text{seed}}^M \leq h_{c,\text{gcd}}^M = h_{\text{gcd}}^M\).

(3) When \(M\) is the set of all matches with costs strictly less than \(r\), admissibility follows directly from Lemma 2 with \(v = v_t\) via

\[
h_{\text{gcd}}^M(u) \leq d(u, v_t) + h_{\text{gcd}}^M(v_t) = d(u, v_t) = h^*(u).\]

\[\square\]

B.2 Match pruning

During the \(A^*\) search, we continuously improve our heuristic using match pruning. The pruning increases the value of our heuristics and breaks their admissibility. Nevertheless, we prove in two steps that \(A^*\) with match pruning still finds a shortest path. First, we introduce the concept of a weakly-admissible heuristic and show that \(A^*\) using a weakly-admissible heuristic finds a shortest path (Thm. 2). Then, we show that our pruning heuristics are indeed weakly admissible (Thm. 3).

\(A^*\) with a weakly-admissible heuristic finds a shortest path.

Definition 6 (Fixed vertex). A vertex \(u\) is fixed if it is expanded and \(A^*\) has found a shortest path to it, that is, \(g(u) = g^*(u)\).

A fixed vertex cannot be opened again (Algorithm 1, line 18), and hence remains fixed.

Definition 7 (Weakly admissible). A heuristic \(\hat{h}\) is weakly admissible if at any moment during the \(A^*\) search there exists a shortest path \(\pi^*\) from \(v_s\) to \(v_t\) in which all vertices \(u \in \pi^*\) after its last fixed vertex \(v^*\) satisfies \(\hat{h}(u) \leq h^*(u)\).

To prove that \(A^*\) finds a shortest path when used with a weakly-admissible heuristic, we follow the structure of Hart et al. (1968). First we restate their Lemma 1 in our notation with a slightly stronger conclusion that follows directly from their proof.

Lemma 3 (Lemma 1 of Hart et al. (1968)). For any unfixed vertex \(v\) and for any shortest path \(\pi^*\) from \(v_s\) to \(v_t\) there exists an open vertex \(v^*\) on \(\pi^*\) with \(g(v^*) = g^*(v^*)\) such that \(\pi^*\) does not contain fixed vertices after \(v^*\).

Next, we prove that in each step the \(A^*\) algorithm can proceed along a shortest path to the target:

Corollary 1 (Generalization of Corollary to Lemma 1 of Hart et al. (1968)). Suppose that \(h\) is weakly admissible, and that \(A^*\) has not
terminated. Then, there exists an open vertex $n'$ on a shortest path from $v_s$ to $v_t$ with $f(n') \leq g^*(v_t)$.

**Proof.** Let $\pi'$ be the shortest path from $v_s$ to $v_t$ given by the weak admissibility of $h$ (Def. 7). Since $A^*$ has not terminated, $v_t$ is not fixed. Substitute $n = v_t$ in Lemma 3 to derive that there exists an open vertex $n'$ on $\pi'$ with $g(n') = g^*(n')$. By definition of $f$ we have $f(n') = g(n') + h(n')$. Since $\pi'$ does not contain any fixed vertices after $n'$, the weak admissibility of $h$ implies $h(n') \leq h^*(n')$. Thus, $f(n') = g(n') + h(n') \leq g^*(n') + h^*(n') = g^*(v_t)$.

**Theorem 2.** $A^*$ with a weakly-admissible heuristic finds a shortest path.

**Proof.** The proof of Theorem 1 in Hart et al. (1968) applies, with the instead of an arbitrary shortest path, we use the specific path $\pi$ given by the weak admissibility and the specific vertex $n'$ given by Corollary 1.

Our heuristics are weakly admissible. A consistent heuristic finds the correct distance to each vertex as soon as it is expanded. While our heuristics are not consistent, this property is true for states at the starts of seeds (when $i$ is a multiple of the seed length $k$, or when $i = n$).

**Lemma 4.** For $h \in \{h_1, h_{cs}, h_{gcs}\}$, every state at the start of a seed becomes fixed immediately when $A^*$ expands it.

**B.4 Computation of the (chaining) seed heuristic**

In this section we prove (Thm. 5) that we can change the dependency of GCSH on $c_{gs}$ to $c_{seed}$ by introducing a new partial order $\succeq_T$ on the matches. This way, Thm. 4 applies and we can efficiently compute GCSH. Recall that the gap-seed cost is $c_{gs} = \max(c_{gap}, c_{seed})$, and that the gap transformation is:

**Definition 4** (Gap transformation). The partial order $\succeq_T$ on states is induced by comparing both coordinates after the gap transformation

$$T: (i, j) \rightarrow (i-j-P(i, j), j-i-P(i, j))$$

The following lemma allows us to determine whether $c_{gap}$ or $c_{seed}$ dominates the cost $c_{gs}$ between two matches, based on the relation $\succeq_T$.

**Lemma 6.** Let $u$ and $v$ be two states with $v$ at the start of some seed. Then $u \succeq_T v$ if and only if $c_{gap}(u, v) \leq c_{seed}(u, v)$. Furthermore, $u \succeq_T v$ implies $u \preceq v$.

**Proof.** Let $u = (i, j)$ and $v = (i', j')$. By definition, $u \succeq_T v$ is equivalent to

$$\begin{align*}
&[(i-j-P(u) \leq i'-j-P(u) \Rightarrow P(u) - P(v)) \land \neg (i'-j) \leq P(u) - P(v)) \\
&[j-i-P(u) \leq j'-i-P(v) \Rightarrow (i'-i) - (j'-j) \leq P(u) - P(v)).
\end{align*}$$

This simplifies to

$$c_{gap}(u, v) = |(i'-i) - (j'-j)| \leq P(u) - P(v) = c_{seed}(u, v),$$

where the last equality holds because $v$ is at the start of a seed.

For the second part, $u \succeq_T v$ implies $0 \leq c_{gap}(u, v) \leq c_{seed}(u, v) = P(u) - P(v)$ and hence $P(u) \geq P(v)$. Since $v$ is at the start of a seed, this directly implies $i \leq i'$. Since seeds have length $k \geq r$ we have

$$|(i'-i) - (j'-j)| \leq P(u) - P(v) = r \cdot |S_i..i'| \leq r \cdot (i'-i) / k \leq i'-i.$$

This implies $j'-j \geq 0$ and hence $j \leq j'$, as required.
A direct corollary is that for \( u \preceq v \) with \( v \) at the start of some seed, we have
\[
    c_{gs}(u,v) = \begin{cases}  
c_{seed}(u,v) & \text{if } u \preceq_T v, \\
c_{gap}(u,v) & \text{if } u \not\preceq_T v.
\end{cases}
\]  
(5)

A second corollary is that start(m) \( \preceq_T \) end(m) for all matches \( m \in \mathcal{M} \), since a match from \( u \) to \( v \) satisfies \( c_{gap}(u,v) < r = c_{seed}(u,v) \) by definition.

**Lemma 7.** When the set of matches \( \mathcal{M} \) is consistent, \( h_{\mathcal{M}}^{g}(u) \) can be computed using \( \preceq_T \)-chains only:
\[
h_{\mathcal{M}}^{g}(u) := h^{g}_{\mathcal{M},c_{gs}}(u) = \begin{cases}  
h_{\mathcal{M}}^{g}(u) & \text{if } u \preceq_T v_t, \\
c_{gap}(u,v_t) & \text{if } u \not\preceq_T v_t.
\end{cases}
\]

**Proof.** We write \( h := h^{g}_{\mathcal{M}} = h^{g}_{\mathcal{M},c_{gs}} \) (Def. 1) and \( h' := h^{M,c_{gs}} \).

Case 1: \( u \not\preceq_T v_t \). Let \( u \preceq m_1 \cdots \preceq m_l \preceq v_t \) be a chain minimizing \( h(u) \) in Def. 1, so
\[
h(u) = c_{gs}(u,m_1) + c_{gs}(m_1,m_2) + \cdots + c_{gs}(m_l,v_t).
\]

By definition, \( c_{gs} \geq c_{gap} \), and \( c_{gs}(m_l) \geq c_{gap}(\text{start}(m_l),\text{end}(m_l)) \), so the weak triangle inequality (Lemma 1) for \( c_{gap} \) gives
\[
h(u) \geq c_{gap}(u,m_1) + c_{gap}(m_1,m_2) + \cdots + c_{gap}(m_l,v_t) \geq c_{gap}(u,v_t).
\]

Since \( u \not\preceq_T v_t \), the empty chain \( u \not\preceq_T v_t \) has cost \( h(u) \leq c_{gs}(u,v_t) = c_{gap}(u,v_t) \). Combining the two inequalities, \( h(u) = c_{gap}(u,v_t) \).

Case 2: \( u \preceq_T v_t \). First rewrite \( h \) and \( h' \) recursively as
\[
h(u) = \min_{u \preceq_T m \preceq_T v_t} (c_{gs}(u,m) + c_{gs}(m) + h(\text{end}(m))) \quad (6)
\]
\[
h'(u) = \min_{u \preceq_T m \preceq_T v_t} (c_{gs}(u,m) + c_{gs}(m) + h'(\text{end}(m))), \quad (7)
\]
both with base case \( h(v_t) = h'(v_t) = 0 \) after eventually taking \( m_\omega \). We will show that
\[
h(u) = \min_{u \preceq_T m \preceq_T v_t} (c_{gs}(u,m) + c_{gs}(m) + h(\text{end}(m))),
\]
(8)

which is exactly the recursion for \( h' \), so that by induction \( h(u) = h'(u) \).

By Lemma 6, every \( \preceq_T \)-chain is a \( \preceq_S \)-chain, so \( h(u) \preceq_T h'(u) \). To prove \( h(u) = h'(u) \), it remains to show the reverse inequality, \( h'(u) \preceq_T h(u) \). To this end, choose a match \( m \) that

- (priority 0) minimizes \( h(u) \) in Eq. (6), and among those, has
- (priority 1) maximal \( c_{seed}(u,m) \), and among those, has
- (priority 2) minimal \( c_{gap}(u,m) \), and among those, has
- (priority 3) minimal \( c_{gap}(m,v_t) \).

We show that \( u \preceq_T m \) (in 2.A) and \( m \preceq_T v_t \) (in 2.B), which proves Eq. (8).

Part 2.A: \( u \preceq_T m \). Let \( s \) and \( t \) be the begin and end of \( m \) (Fig. 7), and let \( m' \) be a match minimizing \( h(t) \) in Eq. (6) so
\[
h(t) = c_{gs}(t,m') + c_{gs}(m') + h(\text{end}(m')).
\]
Since \( m' \) comes after \( t \) we have \( c_{seed}(u,m') > c_{seed}(u,m) \) (p.1) and hence \( m' \) does not minimize \( h(u) \) (p.0):
\[
h(u) < c_{gs}(u,m') + c_{gs}(m') + h(\text{end}(m')).
\]
Using the minimality of \( m \), the non-minimality of \( m' \), and the triangle inequality we get
\[
c_{gs}(u,s) + c_{gs}(m) + h(t) = h(u)
\]
\[
< c_{gs}(u,m') + c_{gs}(m') + h(\text{end}(m'))
\]
\[
\leq c_{gs}(u,t) + c_{gs}(t,m') + c_{gs}(m') + h(\text{end}(m'))
\]
\[
= c_{gs}(u,t) + h(t)
\]
so we have
\[
c_{gs}(u,m) + c_{gs}(m) < c_{gs}(u,t). \quad (9)
\]
From the triangle inequality for \( c_{gap} \), from \( c_{gap}(u,s) \leq c_{gs}(u,s) \), and from Eq. (9) we obtain
\[
c_{gap}(u,t) \leq c_{gap}(u,s) + c_{gap}(s,t) \leq c_{gs}(u,s) + c_{gs}(m) < c_{gs}(u,t).
\]
This implies \( c_{gs}(u,t) = c_{seed}(u,t) \) and hence reusing Eq. (9)
\[
c_{gap}(u,s) + c_{gs}(m) \leq c_{gs}(u,s) + c_{gs}(m)
\]
\[
< c_{seed}(u,t) = c_{seed}(u,s) + c_{seed}(s,t).
\]
We have \( c_{gs}(m) = c_{seed}(s,t) - \text{score}(m) \), so the above simplifies to \( c_{gap}(u,s) < c_{seed}(u,s) + \text{score}(m) \) and since these are integers \( c_{gap}(u,s) \leq c_{seed}(u,s) + \text{score}(m) - 1 \).

When \( \text{score}(m) = 1 \), this implies \( c_{gap}(u,s) < c_{seed}(u,s) \) and \( u \preceq_T s = \text{start}(m) \) by Lemma 6.

When \( \text{score}(m) > 1 \), suppose that \( c_{gap}(u,s) > c_{seed}(u,s) \geq 0 \).
That means that \( u \) is either above or below the diagonal of \( s \). Let \( s_1 = (s_1,s_j \neq 1) \) be the state adjacent to \( s \) on the same side of this diagonal as \( u \). This state exists since \( u \preceq s_1 \preceq s \). Then \( c_{gap}(u,s_1) = c_{gap}(u,s) - 1 \), and by consistency of \( \mathcal{M} \) there is a match \( m_1 \) from \( s_1 \) to \( t \) with \( c_{gs}(m_1) \leq c_{gs}(m) + 1 \). Then
\[
c_{gs}(u,s_1) + c_{gs}(m_1) + h(t)
\]
\[
\leq c_{gs}(u,s) - 1 + c_{gs}(m) + 1 + h(t) = h(u),
\]
showing that \( m_1 \) minimizes \( h(u) \) (p.0). Also \( c_{seed}(u,m_1) = c_{seed}(u,m_1) \) (p.1) and \( c_{gap}(u,m_1) < c_{gap}(u,m) \) (p.2), so that \( m_1 \) contradicts the minimality of \( m \). Thus, \( c_{gap}(u,s) > c_{seed}(u,s) \) is impossible and \( u \preceq_T s \).

Part 2.B: \( m \preceq_T v_t \). When there is some match \( m' \) succeeding \( m \) in the chain, we have \( m \preceq m' \preceq v_t \) and hence \( m \preceq v_t \). Thus, suppose that \( m \) is the only match in the chain \( u \preceq m \preceq v_t \) minimizing \( h(u) \). We repeat the proofs of Part 2.A in the reverse direction to show that \( m \preceq_T v_t \).
Since $c_{seed}(u, m)$ is maximal, $h(u) < c_{gap}(u, m)$ and thus

$$h(u) = c_{gap}(u, s) + c_m(m) + c_{gap}(m, v_t) < c_{gap}(u, v_t).$$

By assumption we have $u \preceq_T v_t$ and thus $c_{gap}(u, v_t) = c_{seed}(u, v_t)$. This gives

$$c_{seed}(u, s) + c_{seed}(s, t) - \text{score}(m) + c_{gap}(t, v_t) < c_{seed}(u, s) + c_{seed}(s, t) + c_{gap}(t, v_t).$$

Cancelling terms we obtain $c_{gap}(t, v_t) < c_{seed}(t, v_t) + 1$ and since they are integers $c_{gap}(t, v_t) \leq c_{seed}(t, v_t) + \text{score}(m) - 1$. When $\text{score}(m) = 1$, this implies $t \preceq_T v_t$, as required.

When $\text{score}(m) > 1$, suppose that $c_{gap}(t, v_t) > c_{seed}(t, v_t)$. Let $t_2 = (t_i, t_j \neq 1)$ be the state adjacent to $t$ on the same side of the diagonal as $v_t$. By consistency of $M$ there is a match $m_2$ from $s$ to $t_2$ with $\text{score}(m_2) \geq \text{score}(m) - 1$ and $c_{gap}(t_2, v_t) = c_{gap}(t, v_t) - 1$. Then $m_2$ minimizes $h(u)$ (p.0)

$$c_{gap}(u, s) + c_m(m_2) + c_{gap}(t_2, v_t) \leq c_{gap}(u, s) + c_m(m) + 1 + c_{gap}(t, v_t) - 1 = h(u),$$

and furthermore $c_{seed}(u, m_2) = c_{seed}(u, m)$ (p.1), $c_{gap}(u, m_2) = c_{gap}(u, m)$ (p.2), and $c_{gap}(m_2, v_t) < c_{gap}(m, v_t)$ (p.3), contradicting the choice of $m$, so $c_{gap}(t, v_t) > c_{seed}(t, v_t)$ is impossible and $t \preceq_T v_t$. \qed

Theorem 5. Given a consistent set of matches $M$, the gap-chaining seed heuristic can be computed using scores in the transformed domain:

$$h_{\text{gap}}^M(u) = \begin{cases} P(u) - S_T(u) & \text{if } u \preceq_T v_t, \\ c_{gap}(u, v_t) & \text{if } u \not\preceq_T v_t. \end{cases}$$

Proof. Write $h := h^M_{\text{gap}}$ and $h' := h^M_{\text{gap}, c_{gap}}$. When $u \not\preceq_T v_t$, $h(u) = c_{gap}(u, v_t)$ by Lemma 7. Otherwise, when $u \preceq_T v_t$, we have $h(u) = h'(u)$. Let $u \preceq_T m_1 \preceq_T \ldots \preceq_T v_t$ be a $T$-chain for $h'$ as in Def. 1. All terms in $h'$ satisfy $\text{end}(m_i) \preceq_T \text{start}(m_{i+1})$, so $c_{gap} \leq c_{seed}$ and by Lemma 6 $c_{gap}(m_i, m_{i+1}) = c_{seed}(m_i, m_{i+1})$. Thus, $h' = h_{\preceq_T, c_{seed}}$ and $h_{\text{gap}}^M(u) = P(u) - S_T(u)$ by Thm. 4. \qed
C Further results

C.1 A*PA parameter grid search

We ran a parameter grid search over various $r$ and $k$ with 5 parallel jobs for both synthetic data and human data.

**Synthetic data.** Figs. 8a and 8b compare the runtime of A*PA with GCSH with DT for varying $k$ and $r$ at $n \in \{10^3, 10^4, 10^5\}$ bp and (a) $d=4\%$ and (b) $d=12\%$. Black lines indicate $r=1$ and green lines indicate $r=2$. Missing datapoints are either due to timing out at $n/N \cdot 1000\, s$, or in 2 cases due to exceeding the 32 GiB memory limit. Red dots indicate alignments where $k$ is too small or large (see App. C.1).

**Human data.** Fig. 9 shows that $r=2$ and $k=15$ minimize the runtime of the third quartile on both datasets. On the ONT reads without genetic variation, choosing a larger $k$ slightly reduces the median runtime, but increases the number of timeouts, making $k=15$ a reasonable choice.

**Bounds on $k$.** The runtime is generally not very sensitive to the exact choice of $k$ as long as it falls within two bounds. First, too many spurious matches are prevented by setting $k \geq \log_4 n$ when $r=1$ and $k \geq 2 + \log_4 n$ when $r=2$. For example, $n=10^6$ would require $k \geq 10$ when $r=1$ or $k \geq 12$ when $r=2$. Secondly, the potential $P \approx r \cdot n/k$ of the heuristic should be sufficiently larger than the divergence, which implies $k$ should be a bit less than $r/d$. So, for $d=12\%$ we would choose $k \leq 1/0.12 = 8.3$ when $r=1$ and $k \leq 2/0.12 = 16.7$ when $r=2$.

Figs. 8 and 9 show alignments where $k$ is not within these bounds in red, and indeed these alignments are relatively slow.

C.2 Runtime scaling with length

In Fig. 4a we compare our A* heuristics with Edlib and BWA in terms of runtime scaling with sequence length $n$.

C.3 Expanded states and equivalent band

The main benefit of an A* heuristic is a lower number of expanded states, which translates to faster runtime. Instead of evaluating the runtime scaling with length (Fig. 10), we can judge how well a heuristic approximates the edit distance by directly measuring the *equivalent band* (Fig. 11) of each alignment: the number of expanded states divided by sequence length $n$, or equivalently, the number of expanded states per base pair. The theoretical lower bound is an equivalent band of $1$, indicating that the number of expanded states is linear on the given domain. The equivalent band of SH with inexact matches starts to grow around $n \geq 3 \cdot 10^6$ at divergence $d=4\%$, and around $n \geq 5 \cdot 10^6$ at $d=12\%$. Because of the chaining, CSH and GCSH cope with spurious matches and remain constant in equivalent band (i.e. linear expanded states with $n$). The equivalent band for GCSH is lower than CSH due to better accounting for indels. The DT variants expand fewer states by skipping non-farthest reaching states, also lowering the equivalent band.
C.4 Runtime scaling with divergence: two modes

Figure 12 shows the runtime scaling with divergence for various heuristics. We notice two regimes of operation, depending on whether the heuristic potential \( P \) is sufficient to compensate for the edit distance: near-linear in \( n \) (constant in \( d \)) and quadratic in \( n \) (linear in \( d \)). The edit distance becomes larger than the potential \( P \) around \( d = r/k \). For \( k=15 \) as in Fig. 12, the threshold is near \( d=1/k=0.7\% \) for exact matches and near \( d=2/k=13.3\% \) for inexact matches. Every error not accounted for by the heuristic triggers a search “to the side”, causing A* to explore \( O(n) \) additional states. When using DT, only \( O(s-P) \) additional farthest reaching states are explored instead, where \( s \) is the edit distance. This leads to observed runtimes of \( O(n+n\cdot\max(s-P,0)) \) without DT, and \( O(n+\max(s-P,0)^2) \) with DT. These are similar to Edlib’s \( O(ns) \) and BiWFA’s \( O(n+s^2) \), but skipping over the first \( P \) errors.

---

**Table 2. Human datasets statistics.** ONT reads only include short gaps, while genetic variation also includes long gaps. Cnt: number of sequence pairs. Max gap: longest gap in the reconstructed alignment.

| Dataset    | Length [kbp] | Divergence [%] | Max gap [kbp] |
|------------|-------------|----------------|--------------|
|             | min | mean | max   | min | mean | max   | min | mean | max   |
| ONT        | 50  | 500 | 594  | 849 | 2.7  | 6.3   | 18.0 | 0.02 | 0.1  | 1    |
| ONT+gen.var. | 48  | 502 | 632  | 1053| 4.4  | 7.4   | 19.8 | 0.05 | 1.9  | 42   |

**Table 3** compares memory usage of aligners on synthetic sequences of length \( n=10^8 \). Alignments with inexact matches (\( d\geq8\% \)) use significantly...
Fig. 12. Runtime scaling with divergence. The figure shows the same results as Fig. 4c, but zoomed out to show scaling for high $d$, where runtime of $\text{A}^\ast\text{PA}$ degrades from constant in $d$ to linear in $d$, $r=1$ and $r=2$ indicate exact and inexact matches, respectively. Red dots show where the heuristic potential is less than the edit distance. Missing datapoints timed out after 100 s.

More memory than those with exact matches because more $k$-mer hashes need to be stored to find inexact matches. Table 4 compares memory usage on the human data sets.

| Aligner | $d=1\%$ | $d=4\%$ | $d=8\%$ | $d=12\%$ |
|---------|----------|----------|----------|-----------|
| $\text{EDLIB}$ | 2 | 2 | 2 | 2 |
| $\text{BiWFA}$ | 15 | 13 | 14 | 18 |
| SH | 50 | 59 | 151 | 480 |
| CSH | 52 | 59 | 151 | 261 |
| GCSH | 49 | 50 | 151 | 255 |
| SH + DT | 49 | 49 | 151 | 150 |
| CSH + DT | 49 | 49 | 151 | 150 |
| GCSH + DT | 48 | 46 | 152 | 150 |

Table 3. Memory usage per algorithm (synthetic data, $n=10^6$). Exact matches are used when $d \leq 4\%$, and inexact matches when $d \geq 8\%$.

| Aligner | ONT reads | + genetic var. |
|---------|-----------|----------------|
|         | Median | Max | Median | Max |
| $\text{EDLIB}$ | 2 | 5 | 2 | 6 |
| $\text{BiWFA}$ | 11 | 19 | 15 | 24 |
| $\text{A}^\ast\text{PA (GCSH + DT)}$ | 160 | 3478 | 270 | 6926 |

Table 4. Memory usage [MB] of aligners on human data. Medians are over all alignments; maximums are over alignments not timing out.

C.7 Runtime profile of $\text{A}^\ast\text{PA}$

Figures 13a and 13b compare the time used by stages of $\text{A}^\ast\text{PA}$. On synthetic data with exact matches ($r=1$), the runtime is spread over all parts of the algorithm. When using inexact matches, precomputation takes a significant fraction of the total time and updating contours becomes slower due to the increased number of matches.

On human data, faster alignments spend a large fraction of their time on the precomputation, followed by the updating of contours after pruning matches. Slower alignments on the other hand are limited by the performance of the $\text{A}^\ast$ algorithm, and spend a large fraction of time on opening and expanding states, and evaluating the heuristic.

C.8 Quadratic scaling in complex regions

Figure 14 shows the effect of complex regions in the sequences on GCSH (and thus on all our heuristics).

C.9 Linear mode without matches

States are penalized by the number of remaining seeds that cannot be matched. So, curiously, matches are not always needed to direct the $\text{A}^\ast$ search to an optimal path. In fact, when each seed contains exactly one mutation seed heuristic scales linearly even though there are no matches, as shown by the artificial example in Fig. 15.

C.10 Comparison of heuristics and techniques

Figure 16 shows the effect of our heuristics and optimizations for aligning complex short sequences. The effect of pruning is most noticeable for CSH and GCSH without DT. GCSH is our most accurate heuristic, so, as expected, it leads to the lowest number of expanded states.
Fig. 13. Runtime distributions per stage of A’PA (GCSH with DT) (stages do not overlap). Stage A’ includes expanding and opening states. Pruning matches includes consistency checks. Updating contours includes updating of contours after pruning. (a) On synthetic data \((n=10^6 \text{ bp}, N=10^7 \text{ bp total}, k=15)\). The circle area is proportional to the total runtime. Figures for \(r=1\) and \(d\geq 8\%\) are skipped due to timeouts (100 s). (b) On human data \((r=2)\). Alignments are sorted by total runtime (timeouts not shown).

Fig. 14. Quadratic exploration behavior for complex alignments (GCSH with DT, \(r=2, k=10\), synthetic sequences, \(n=1000\)). (a) A highly divergent region, (b) a deletion, and (c) a short repeated pattern inducing a quadratic number of matches. The colour corresponds to the order of expansion, from blue to red.

Fig. 15. Artificial example of A’ with seed heuristic with no matches \((n=m=50, r=1, k=5, 80\% \text{ similarity}, 1 \text{ mutation per seed alternating substitution, insertion, and deletion})\). The background colour indicates \(h_s(u)\) with higher values darker. Expanded states are green (a), open states blue (a).
Fig. 16. Expanded states for various heuristics and techniques, on a sequence containing a noisy region, a repeat, and an indel \((n=1000, d=17.5\%\)). The colour shows the order of expanding, from blue to red. The sequences include a highly divergent region, a repeat, and a gap. Matches are shown as black diagonals, with inexact matches in grey and pruned matches in red. The final path is black. Dijkstra does not have pruning variants, and Dijkstra with DT is equivalent to WFA. More accurate heuristics reduce the number of expanded states by more effectively punishing repeats (CSH) and gaps (GCSH). Pruning reduces the number of expanded states before the pruned matches, and diagonal transition reduces the density of expanded states in quadratic regions.
D Notation

| Object            | Notation |
|-------------------|----------|
| **Sequences**     |          |
| Alphabet          | $\Sigma = \{A, C, G, T\}$ |
| Sequences         | $A = a_1 \ldots a_n \in \Sigma^*$ |
| Substring         | $A_{i..j} = a_i \ldots a_{j-1}$ |
| Prefix            | $A_{i:j} = a_i \ldots a_{j-1}$ |
| Edit distance     | $d = ed(A, B)$ |
| Divergence        | $d = ed(A, B)/n$ |
| Error rate        | $e$ |
| **Alignment graph** |         |
| Vertices (states) | $(u, v) \in V = \{(i, j) \mid 0 \leq i \leq n, 0 \leq j \leq m\}$ |
| Edges             | $(i, j) \rightarrow (i+1, j)$ |
| Path, shortest path | $\pi, \pi'$ |
| Cost              | $c_{\text{Path}}(\pi)$ |
| **Diagonal transition** |       |
| Farthest-reaching state | $F_{ak} = i+j$ on diagonal $k=i-j$ |
| **A***            |          |
| Start and target state | $v_s = (0,0), v_t = (n,m)$ |
| Distance from $v_s$ | $g = d(v_s, \cdot)$ |
| Distance to $v_t$  | $h = d(\cdot, v_t)$ |
| Heuristic          | $h$ |
| Best distance from start | $g$ |
| Estimated distance | $f = g + h$ |
| Admissible heuristic | $h(u) \leq d(u, v) + h(v)$ |
| Consistent heuristic | $h(u) \leq d(u, v) + h(v)$ |
| Expanded states   | $E$ |

| Object            | Notation |
|-------------------|----------|
| **Seeds and matches** |          |
| Seed length       | $k$ |
| Seed potential    | $r$ |
| Seeds             | $s \in S, s_1 = A_{lk..tk+k}$ |
| Seeds in suffix   | $S_{ni} = \{s_j \in S \mid k \geq i\}$ |
| Alignment of seed | $\pi_s$ |
| Matches (per seed) | $m \in M, M_s, M = |M|$ |
| Terminal match    | $m_u$, from $v_{t1}$ to $v_{t2}$ |
| Cost of match     | $0 \leq c_{\text{gap}}(m) < r$ |
| Score of match    | $0 < \text{score}(m) = r - c_{\text{gap}}(m) \leq r$ |
| Score of seed     | $\text{score}(s) = \max_{m \in M, s \in S} \text{score}(m)$ |
| **Chains**        |          |
| Preceding states  | $(i, j) \leq (i', j')$ when $i \leq i'$ and $j \leq j'$ |
| Preceding matches | $m \leq m'$ when $\text{end}(m) = \text{start}(m')$ |
| Partial order     | $u \leq m$ when $u \leq \text{start}(m)$ |
| i-order           | $(i, j) \preceq (i', j')$ when $i \leq i'$ |
| $s_p$-chain       | $m_1 \preceq s_p \cdots s_p m_t \preceq s_p v_{t2}$ |

| Object            | Notation |
|-------------------|----------|
| **Chaining costs** |          |
| Chaining cost     | $\gamma(m, m')$ |
| Gap cost          | $c_{\text{gap}}((i, j), (i', j')) = |(i'-i)-(j'-j)|$ |
| Seed cost         | $c_{\text{seed}}(u, v) = r - |S_u - v|$ |
| Gap-seed cost     | $c_{\text{gap}} = \max(c_{\text{gap}}, c_{\text{seed}})$ |

| Object            | Notation |
|-------------------|----------|
| **Scores**        |          |
| Potential         | $P(i, j) = r \cdot |S_{\text{end}}|$ |
| Chain score       | $S_{\mu}(m) = \max_{ps_p \cdots s_p v_{t2}} \text{score}(m) + \ldots + S_{\mu}(u) = \max_{ps_p \cdots s_p v_{t2}} S_{\mu}(m)$ |
| Computation       | $h_{p, \text{seed}}(u) = P(u) - S_{\mu}(u)$ |

| Object            | Notation |
|-------------------|----------|
| **Heuristics**    |          |
| SH                | $h_d(u) = P(u) - S_i(u)$ |
| CSH               | $h_d(u) = P(u) - S_i(u)$ |
| GCSH              | $h_{\text{CSH}}(u) = \max(c_{\text{gap}}(u, v_t), P(u) - S_T(u))$ |
| Pruning heuristic | $h_{\text{pruning}}(i, j) = (i-j-P(i, j), j-i-P(i, j))$ |

| Object            | Notation |
|-------------------|----------|
| **Layers**        |          |
| Layer             | $L_{\ell} = \{u \mid S_u(u) \geq \ell\}$ |
| Dominant state    | $u \in L_{\ell}$ s.t. $v \in L_{\ell} \mid u \leq v = \{u\}$ |