

TRADING RUNNING TIME FOR MEMORY IN PHYLOGENETIC LIKELIHOOD COMPUTATIONS

Fernando Izquierdo-Carrasco¹, Julien Gagneur² and Alexandros Stamatakis¹

¹*The Exelixis Lab, Scientific Computing Group, Heidelberg Institute for Theoretical Studies
Schloss-Wolfsbrunnengasse 35, D-69118 Heidelberg, Germany*

²*European Molecular Biology Laboratory, Meyerhofstr. 1, 69117 Heidelberg, Germany*

Keywords: Memory versus runtime trade-offs, Phylogenetic likelihood function, RAxML.

Abstract: The revolution in wet-lab sequencing techniques that has given rise to a plethora of whole-genome or whole-transcriptome sequencing projects, often targeting 50 up to 1000 species, poses new challenges for efficiently computing the phylogenetic likelihood function both for phylogenetic inference and statistical post-analysis purposes. The phylogenetic likelihood function as deployed in maximum likelihood and Bayesian inference programs consumes the vast majority of computational resources, that is, memory and CPU time. Here, we introduce and implement a novel, general, and versatile concept to trade additional computations for memory consumption in the likelihood function which exhibits a surprisingly small impact on overall execution times. When trading 50% of the required RAM for additional computations, the average execution time increase because of additional computations amounts to only 15%. We demonstrate that, for a phylogeny with n species only $\log(n) + 2$ memory space is required for computing the likelihood. This is a promising result given the exponential growth of molecular datasets.

1 INTRODUCTION

The rapid accumulation of molecular sequence data that is driven by novel wet-lab techniques poses new challenges regarding the design of programs for phylogenetic inference that rely on computing the Phylogenetic Likelihood Function (PLF) for phylogenetic inference or statistical post-analysis. For instance, we are currently involved in two large-scale sequencing efforts aimed at obtaining, whole genome data for approximately 150 species and whole transcriptome data for approximately 1000 species¹.

In all popular Maximum Likelihood (ML) and Bayesian phylogenetic inference programs, the PLF dominates both, the overall execution time *as well as* the memory requirements by typically 85% - 95% (Stamatakis and Ott, 2008).

Based on our interactions with the RAxML user community, we find that memory shortages are increasingly becoming a problem and represent *the* main limiting factor for large-scale phylogenetic analyses, especially at the Genome level. At the same time, the amount of available genomic data is growing

at a faster pace than RAM (Random Access Memory) sizes and RAM access speeds. For instance, to compute the likelihood on a simulated DNA alignment with 1,481 species and 20,000,000 sites (corresponding roughly to the 20,000 genes in the human genome) on a single tree topology under a simple statistical model of nucleotide substitution within 48 hours, 1TB of memory and a total of 672 cores are required.

We have already assessed some potential solutions to this problem in previous work:

1. **Single Precision.** Deploying single precision arithmetics that can reduce memory requirements by 50% but can potentially induce numerical instability, especially on datasets with more than 1000 taxa (Berger and Stamatakis, 2010).
2. **Gappy Alignments.** Novel algorithmic solutions which rely on dataset-specific properties, that is, their efficiency/applicability depends on the specific properties of the Multiple Sequence Alignment (MSA) that is used as input (Stamatakis and Alachiotis, 2010). In particular, these algorithmic approaches do not work on multi-gene datasets that do not contain enough missing data on a per-gene basis, that is, when gene sequences for a

¹Because of non-disclosure agreements we can not provide further details about these projects at present.

certain number of taxa under study are not available. In other words, these approaches are not applicable to emerging, dense whole-genome MSAs without missing data.

3. **Out-Of-Core.** We recently introduced a generally applicable method for saving memory, that does not rely on MSA-specific characteristics. We evaluated an out-of-core (frequently also called external memory algorithm) approach that reduces RAM requirements by moving data back and forth between the RAM and the disk (Izquierdo-Carrasco and Stamatakis, 2011). However, the increase of at least one order of magnitude in overall execution times because of slow read/write operations to/from disk does not yield this approach practical, even when a high performance parallel file system is used.

Here, we introduce a novel and significantly more efficient method that is based on trading computations (running time) for memory. The method is completely independent from the MSA structure and is therefore particularly useful for reducing RAM requirements of “dense” whole-genome alignments. Moreover, it can be combined with the **single precision** and **gappy alignment** memory saving approaches (see above) to yield even further savings. When reducing the amount of available RAM to 50% of the total RAM required for computing the likelihood on a tree for a given MSA, we obtain a respective slowdown (because of additional (re-)computations) of only 15%. This represents a relatively small run-time increase that most end-users will not even notice. On the other hand, users will notice when their analyses crash because they have run out of RAM or the OS (operating system) starts paging.

The remainder of this paper is organized as follows: We briefly review related work in Section 2. In Section 3 we describe the underlying idea for the memory saving approach, demonstrate that only $\log(n) + 2$ vectors are required, and introduce two vector replacement strategies. In Section 4 we describe the experimental setup and provide corresponding results. We conclude and discuss directions of future work in Section 5.

2 RELATED WORK

One common approach for reducing memory requirements are out-of-core algorithms. These are specifically designed to minimize the I/O overhead via explicit, application-specific, data placement control and movement. Out-of-core algorithms mostly aim at

optimizing data movement between disk and RAM. A comprehensive review of work on out-of-core algorithms can be found in (Vitter, 2008). In phylogenetics, out-of-core algorithms have been applied to Neighbor-Joining (Wheeler, 2009; Martin Simonsen and Pedersen, 2010). Moreover, we recently introduced a generic out-of-core implementation for PLF computations (Izquierdo-Carrasco and Stamatakis, 2011). As already mentioned, the performance of our out-of-core implementation was rather disappointing and is therefore not applicable to likelihood-based inference on dense whole-genome alignments. Nonetheless, our approach was significantly more efficient than the paging strategy of the OS.

The term time-memory trade-off engineering refers to situations/approaches where memory requirements/utilization is reduced at the cost of additional (re-)computations, that is, at the expense of increased run time. This trade-off engineering approach has been applied in diverse fields such as language recognition (Dri and Galil, 1984), cryptography (Barkan et al., 2006), and packet scheduling (Xu and Lipton, 2005).

We are not aware of any applications of or experiments with time-memory trade-off engineering approaches in the field of computational phylogenetics.

3 RECOMPUTATION OF ANCESTRAL PROBABILITY VECTORS

3.1 PLF Memory Requirements

A detailed discussion on PLF Memory requirements is provided in (Izquierdo-Carrasco and Stamatakis, 2011). We include a part of this discussion, that is also relevant for the work presented here. The PLF is defined on unrooted binary trees. The n extant species/organisms of the MSA under study are located at the tips of the tree, whereas the $n - 2$ inner nodes represent extinct common ancestors. The molecular sequence data in the MSA that has a length of s sites (alignment columns) is located at the tips of the tree. The memory requirements for storing those n tip vectors of length s is not problematic, because one 32-bit integer is sufficient to store, for instance, 8 nucleotides when ambiguous DNA character encoding (requiring 4 bits to store one nucleotide) is used. Hence, for the aforementioned large dataset (1,481 species, 20,000,000 sites) only 13GB are required to store the actual sequence data compared to 1 TB for storing ancestral probability vectors.

Hence, the memory requirements of the PLF are dominated by the space for storing the ancestral probability vectors that are located at the ancestral (inner) nodes of the tree. Depending on the PLF implementation, at least one such vector (a total of $n - 2$) will need to be stored per ancestral node. For each alignment site $i, i = 1 \dots s$, an ancestral probability vector needs to hold the data for the probability of observing an A, C, G or T. Thus, under double precision arithmetics and for DNA data a total of $(n - 2) \cdot 8 \cdot 4 \cdot s$ bytes is required for the most simple statistical model of nucleotide substitution. When the less simplistic, standard Γ model of rate heterogeneity (Yang, 1994) with 4 discrete rates is deployed, this number increases by a factor of four $((n - 2) \cdot 8 \cdot 16 \cdot s)$, since we need to store 16 probabilities for each alignment site i . Furthermore, if protein data is used (that has 20 instead of 4 states) in conjunction with the Γ model of rate heterogeneity, the memory requirements for storing ancestral probability vectors increase to $(n - 2) \cdot 8 \cdot 80 \cdot s$ bytes.

To obtain the likelihood of a given, fixed tree, the PLF performs a post-order tree traversal that starts at a virtual root. Such a virtual root can be placed arbitrarily into any branch of the unrooted tree without changing the overall likelihood of the tree. By means of the post-order tree traversal, the ancestral probability vectors are computed bottom-up from the leaves toward the virtual root.

For a more detailed description of PLF computations and efficient PLF implementations please refer to (Stamatakis, 2011). For understanding the memory access pattern during the post-order traversal, let us consider equation (Felsenstein, 1981) of the Felsenstein pruning algorithm.

This equation computes the ancestral probability vector entry $\tilde{L}_A^{(p)}$ for observing the nucleotide A at site i of a parent node p , with two child nodes l and r given the respective branch lengths b_l and b_r , the corresponding transition probability matrices $P(b_l)$, $P(b_r)$, and the probability vectors of the children $\tilde{L}^{(l)}$, $\tilde{L}^{(r)}$:

$$\tilde{L}_A^{(p)}(i) = \left(\sum_{S=A}^T P_{AS}(b_l) \tilde{L}_S^{(l)}(i) \right) \left(\sum_{S=A}^T P_{AS}(b_r) \tilde{L}_S^{(r)}(i) \right) \quad (1)$$

In order to compute $L_A^{(p)}(i)$, we do not necessarily need to immediately have available in memory vectors $L_S^{(l)}(i)$ and $L_S^{(r)}(i)$, since they can be obtained by a recursive descent (a post-order subtree traversal) into the subtrees rooted at l and r using the above equation. In other words, if we do not have enough memory available, we can recursively re-compute the values of $L^{(l)}$ and $L^{(r)}$. Note that, the recursion terminates when we reach the tips (leaves) of the tree and that

memory requirements for storing tip vectors are negligible compared to ancestral vectors (see above).

If not all ancestral probability vectors fit in RAM, the required vectors for the operation at hand can still be obtained by conducting some additional computations for obtaining them by applying equation 1. We observe that, when both $L_S^{(l)}(i)$ and $L_S^{(r)}(i)$ have been used (consumed) for calculating $L_A^{(p)}(i)$, the two child vectors are not required any more. That is, those two vectors can be omitted/dropped from RAM or be overwritten in RAM to save space. Therefore, the likelihood of a tree can be computed without storing all $n - 2$ ancestral vectors. Instead, a smaller amount of space for only storing $x < (n - 2)$ vectors can be used. Those x vectors can then dynamically be assigned to a subset (varying over time) of the $n - 2$ inner nodes. This gives rise to the following question: Given a MSA with n taxa, what is the minimum number of inner vectors x_{min} that must reside in memory to be able to compute the likelihood on any unrooted binary tree topology with n taxa?

Due to the post-order traversal of the binary tree topology, some ancestral vectors need to be stored as intermediate results. Consider an ancestral node p that has two subtrees rooted at child nodes l and r with identical subtree depth. When the computations on the first subtree for obtaining $L^{(l)}$ have been completed, this vector needs to be kept in memory until the ancestral probability vector $L^{(r)}$ has been calculated to be able to compute $L^{(p)}$.

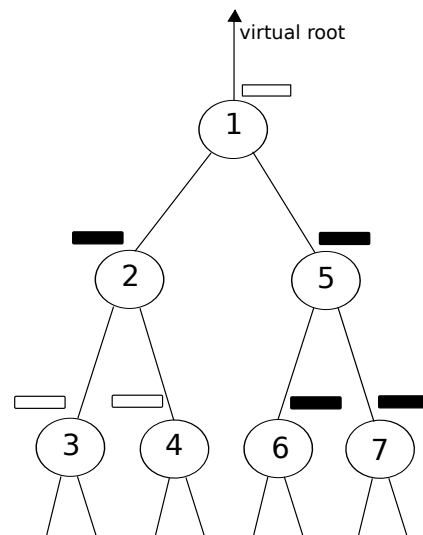


Figure 1: A balanced subtree, where the vector of inner node 1 is oriented in the direction of the virtual root. In order to compute the likelihood of vector 1, the bold vectors must be held in memory. The transparent vectors may reside in memory but are not required to.

In the following section 3.2, we demonstrate that the minimum number of ancestral vectors $x_{min}(n)$ that must reside in RAM to be able to compute the likelihood of a tree with n taxa is $\log_2(n) + 2$. For obtaining this lower memory bound, we consider the worst case tree shape, that is, a perfectly balanced unrooted binary tree with the virtual root placed into the innermost branch.

Figure 1, where we first descended into the left subtree, depicts this worst case scenario for $n = 4$. The probability vectors will be written in the following order: 3, 4, 2, 6, 7, 5, 1. Figure 1 shows the number of vectors required in memory ($\log_2(4) + 2 = 4$) at the point of time where vector 5 needs to be computed. At any other point of time during the post-order traversal, holding 3 vectors in RAM is sufficient to successfully proceed with the computations.

3.2 Minimum Memory Requirements

The underlying idea of our approach is to reduce the total number of ancestral probability vectors residing in RAM at any given point of time. Let PLF be the likelihood function (see Equation 1), which—for the sake of simplifying our notation—can be computed at a tip (essentially at zero computational cost) or an inner node given the ancestral probability vectors of its two child nodes. The PLF always returns an ancestral probability vector as result.

As a pre-processing step, we compute the number of descendants (size of the respective subtree) for each inner node. This can be implemented via a single post-order tree traversal. The memory required for storing the number of descendants as integers at each node is negligible compared to required probability vector space. Given this information, the binary tree data structure can be re-ordered such that, for every node p , the left child node l contains the larger number of descendants and the right child node r contains the smaller number of descendants. We can now compute the PLF of the tree by invoking the following recursive procedure \mathbf{f} at the virtual root p of the tree:

```

proc f( $p$ ) ≡
  if isALeaf( $p$ ) then return( $PLF(p)$ ) fi;
   $v_l := f(p.l)$ ;                               Step 1
   $v_r := f(p.r)$ ;                               Step 2
  return( $PLF(v_l, v_r)$ )                         Step 3
    
```

During this computation, the maximum number of probability vectors $x_{min}(n)$ that need to reside in memory for any tree with n leaves is $\log_2(n) + 2$. This upper bound is required if and only if the binary tree is balanced.

Proof. We demonstrate the above theorem by recursion. $x_{min}(1) = 1$, since for a tree with a single node, only the sequence data (a single probability vector) need to be stored. Now assume that $x_{min}(n-1) = \log_2(n-1) + 2$ and consider a tree with n leaves. We execute all steps of $f(p)$, where p is the root, and keep track of the number of probability vectors that are stored simultaneously in RAM:

- **Step 1.** Computing $f(p.l)$ requires storing at most $x_{min}(n-1)$ probability vectors simultaneously which is strictly less than $\log_2(n) + 2$. Once $f(p.l)$ has been computed only the result vector is stored (i.e., only one single probability vector).
- **Step 2.** Because the subtree size of the right descendant of the root $p.r$ must be $\leq n/2$, this computation needs to simultaneously store at most $\log_2(n/2) + 2 = \log_2(n) + 1$ probability vectors. Here, we need to add the number of probability vectors that required to be maintained in RAM after completion of **Step 1**. Thus, overall, **Step 2** needs to simultaneously hold at most $\log_2(n) + 2$ probability vectors in RAM. Once the results of $f(p.l)$ and $f(p.r)$ have been computed we are left with two probability vectors v_l, v_r that need to reside in memory.
- **Step 3.** This step requires 3 probability vectors to be stored at the same time, namely, v_l, v_r , and the vector to store the result of $PLF(v_l, v_r)$. Note that, to obtain the overall likelihood of the tree (a single numerical value), we need to apply a function $g()$ to the single result vector returned by $f()$, that is, $g(f(p))$. Function $g()$ simply uses the data in the result vector to calculate the likelihood score over all root vector entries.

Hence the peak in memory usage is reached during **Step 2** and its upper bound is $\log_2(n) + 2$. Moreover, this upper bound is reached if and only if the number of descendants in the respective left and right subtrees is identical for all nodes, that is, for a balanced tree.

3.3 Basic Implementation

While holding $\log_2(n) + 2$ vectors in RAM provides sufficient space for computing the PLF, this will induce a significant run-time increase (due to recomputations) compared to holding n vectors in memory. In practice, we need to analyze and determine a reasonable run-time versus RAM trade-off as well as an appropriate vector replacement/overwriting strategy. For instance, in the RAXML or MrBayes PLF implementations, some ancestral vectors (depending on their location in the tree) can be recomputed faster

than others. In particular, cases where the left and/or right child vectors are tip sequences can be handled more efficiently. For instance, an observed nucleotide A at a tip sequence corresponds to a simple probability vector of the form $[P(A) := 1.0, P(C) := 0.0, P(G) := 0.0, P(T) := 0.0]$. This property of tip vectors can be used for saving computations in equation 1.

Also, typical topological search operators for finding/constructing a tree topology with an improved likelihood score such as SPR (Subtree Pruning and Re-grafting), NNI (Nearest Neighbor Interchange) or TBR (Tree Bisection and Reconnection) only apply local changes to the tree topology. In other words, the majority of the ancestral probability vectors is not affected by the topological change and does therefore not need to be recomputed/updated with respect to the locally altered tree topology via a full post-order tree traversal. Therefore, if all ancestral vectors reside in RAM, only a very small part of the tree needs to be (re-)traversed after a SPR move, for instance. All standard ML-based programs such as GARLI (Zwickl, 2006), PHYML 3.0 (Guindon et al., 2010), and RAxML (Stamatakis, 2006) deploy search strategies that typically require updating only a small fraction of probability vectors in the vicinity of the topological change.

Therefore, devising an appropriate strategy (see Section 3.4) for deciding which vectors shall remain in RAM and which can be discarded (because they can be recomputed at a lower computational cost) can have a substantial impact on the induced run time overhead when holding, for instance, $x := n/2$ vectors in RAM. In the following, we will outline how to compute the PLF and conduct SPR-based tree searches with $x < n$ vectors in RAM.

Let n be the number of species, $n - 2$ the number of ancestral probability vectors, and x the number of available slots in memory, where $\log_2(n) + 2 \leq x < n$ (i.e., $n - x$ vectors are not stored, but recomputed on demand). Let w be the number of bytes required for storing an ancestral probability vector (all vectors have the same size). Our implementation will only allocate $x \cdot w$ bytes, rather than $n \cdot w$. We henceforth use the term *slot* to denote a RAM segment of w bytes that can hold an ancestral probability vector.

We define the following C data structure (details omitted) to keep track of the vector-to-slot mapping of all ancestral vectors and for implementing replacement strategies:

```
typedef struct
{
    int numVectors;
    size_t width;
    double **tmpvectors;
    int *iVector;
```

```
    int *iNode;
    int *unpinnable;
    boolean allSlotsBusy;
    unpin_strategy strategy;
}recompVectors;
```

The array `tmpvectors` is a list of pointers to a set of slots (i.e., starting addresses of allocated RAM memory) of size `numVectors` (x) and width `numVector` (w). The array `iVector` has length x and is indexed by the slot id. Each entry holds the node id of the ancestral vector that is currently stored in the indexed slot. If the slot is free, the value is set to a dedicated `SLOT_UNUSED` code. The array `iNode` has length $n - 2$ and is indexed using the unique node ids of all ancestral vectors in the tree. When the corresponding vector resides in RAM, its array entry holds the corresponding slot id. If the vector does not reside in RAM the array entry is set to the special code `NODE_UNPINNED`. Henceforth, we denote the availability/unavailability of an ancestral vector in RAM as `pinned/unpinned`. The array `unpinnable` tracks which slots are available for unpinning, that is, which slots that currently hold an ancestral vector can be overwritten, if required.

The set of ancestral vectors that are stored in the memory slots changes dynamically during the computation of the PLF (i.e., during full tree traversals and tree searches). The pattern of dynamic change in the slot vector also depends on the selected recomputation/replacement strategy. For each PLF invocation, be it for evaluating a SPR move or completely re-traversing the tree, the above data structure is updated accordingly to ensure consistency.

Whenever we need to compute a local tree traversal (following the application of an SPR move) to compute the likelihood of the altered tree topology, we initially just compute the traversal order which is part of the standard RAxML implementation. The traversal order is essentially a list that stores in which order ancestral probability vectors need to be computed. In other words, the traversal descriptor describes the partial or full post-order tree traversal required to correctly compute the likelihood of a tree. For using $x < n$ vectors, we introduce a so-called traversal order check, which extends the traversal steps (the traversal list) that assume that all n vectors reside in RAM. By this traversal order extension, we guarantee that all missing vectors (not residing in RAM) will be recomputed as needed. The effect of reducing the number of vectors residing in RAM is that, traversal lists become longer, that is, more nodes are visited and thereby run time increases. When the traversal is initiated, all vectors in the traversal list that already reside in RAM (they are pinned to a slot) are protected (marked as `unpinnable`) such that, they

will not be overwritten by intermediate vectors of the recomputation steps.

If an ancestral vector slot needs to be written/stored by the traversal, there are three cases:

1. The vector resides in a slot (already in memory). We can just read and/or write to this slot.
2. The vector is not pinned, but there exists a free slot, which is then pinned to this vector.
3. The vector is not pinned and there is no free slot available. A residing vector must be replaced and the corresponding slot needs to be pinned to the required vector.

Since the traversal only visits each vector at most once, the corresponding children of this vector can be unpinned once it has been written to memory. Instead of unpinning them directly, they are merely marked for unpinning. The real overwrite only takes place if the slot is selected by the replacement strategy for storing another vector. Otherwise, the slot will store the values of the current vector for as long as possible for potential future re-use.

3.4 Replacement Strategies

In analogy to the replacement strategies discussed in the out-of-core implementation (Izquierdo-Carrasco and Stamatakis, 2011), there are numerous approaches for deciding which memory slot should be overwritten by a new ancestral vector that does currently not reside in RAM. We implement and analyze the following two replacement strategies.

Random. A random slot not flagged as pinned is selected. The random strategy is a naïve approach with minimal overhead and is used as baseline for performance comparisons.

MRC. Minimum Recomputation Cost. The slot with minimum subtree size (see below) and not flagged as pinned is selected.

The MRC strategy entails a slight overhead for keeping track of which vectors will be most expensive to recompute and should therefore be kept in RAM for as long as possible. Consider an unrooted binary tree T with n tips. Each inner node i in an unrooted binary tree with n taxa can be regarded as a trifurcation that defines three subtrees $T_{i,a}$, $T_{i,b}$, and $T_{i,c}$ corresponding to the three outgoing branches a , b , and c . Given a subtree $T_{i,x}$, we define its subtree size $sts(T_{i,x})$ as the number of tips beneath inner node i in the direction of the outgoing branch x . Thus, $sts(T_{i,a}) + sts(T_{i,b}) + sts(T_{i,c}) = n$ holds for any inner node i in an unrooted binary tree with n taxa/tips. When conducting likelihood computations, the tree is

always rooted by a virtual root. Hence, if the virtual root is located in the direction of branch c the relevant subtree size with respect to recomputation cost at inner node i is $sts(T_i^{rooted}) := sts(T_{i,a}) + sts(T_{i,b})$. We use this rooted subtree size $sts(T_i^{rooted})$ to determine the recomputation cost for each ancestral vector i given a virtual rooting of the tree. In particular, the case $sts(T_i^{rooted}) = 2$ (both children are tips) is particularly cheap to recompute, since tip vectors always reside in RAM *and* recomputing ancestral vector i is cheap (see above). In a perfectly balanced tree with the root placed in the innermost branch, half of the inner vectors have subtree size $sts(T_i^{rooted}) = 2$.

As already mentioned, during a partial or full tree traversal for computing the likelihood, all inner nodes (vectors) involved are oriented in a given direction toward the virtual root. Evidently, the subtree sizes will change when the topology is altered and will need to be updated accordingly.

In order to account for this, we keep an array of subtree sizes, that is, for each inner node we store a subtree size value. Whenever the topology changes, a local traversal descriptor is created. This local traversal descriptor starts at the new virtual root and recursively includes the inner nodes whose orientation has changed after the given topology change. Since this exactly corresponds to the set of nodes whose subtree size values must be updated, the subtree size array can be updated via the same recursive descent.

3.5 Implementation Details

In the following we discuss some important details of the recomputation process.

Largest Subtree First. The standard implementation of the PLF, where all ancestral vectors are available in memory, computes the PLF by conducting a post-order traversal from an arbitrary rooting of the tree. Thus, an ancestral probability vector can be computed once the respective left and right child vectors have been computed. In the standard RAxML implementation, the traversal always recursively descends into the left subtree first. The (arbitrary) choice whether to descend into the left or right subtree first, does not affect performance (nor the results) when all ancestral vectors reside in RAM.

However, when not all vectors reside in RAM, the choice whether to descend into the left or right subtree first *does* matter. This is particularly important if we use the minimum setting $x := \log_2(n) + 2$, since otherwise we may encounter situations where not enough slots are available (see Section 3.2).

Suppose that, as in the standard implementation,

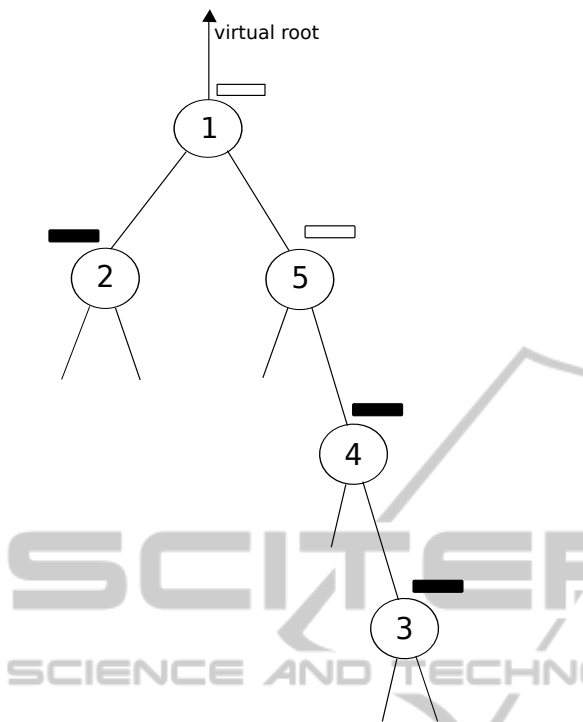


Figure 2: An unbalanced subtree, where the vector of inner node 1 is oriented in the direction of the virtual root. Bold rectangles represent vectors that must be held in memory if we first descend into the left subtree and node 4 is being written.

we always descend into the left subtree first. In the example shown in Figure 2, the left subtree is significantly smaller than the right subtree. We would first descend into the left subtree, which consists of a single inner node. The child ancestral vector corresponding to node 2 must be pinned to its slot. Thereafter, we descend into the right subtree writing and pinning nodes 3, 4, 5 (always assuming that we descend to the left—smaller—subtree of *each* node first). While we keep descending into the right subtree, the ancestral vector corresponding to node 2 remains pinned, because it represents an intermediate result.

In contrast, if we initially descend into the right subtree (which is always larger in the example), there is no need to store intermediate results of the left subtree (node 2). Also, nodes 4 and 5 can be immediately unpinned as soon as their parent nodes have been computed. Thus, by inverting the descent order such as to always descend into the larger subtree first (as required by our proof), we minimize the amount of time for which intermediate vectors must remain pinned. Note that, when two subtrees have the same size, it does not matter into which subtree we descend first.

If we descend into the smaller subtrees first, there

will be more vectors that need to remain pinned for a longer time. This would also reduce the effective size of the set of inexpensive-to-recompute vectors that shall preferentially be overwritten, because more vectors that are cheap to recompute need to remain pinned since they are holding intermediate results. In this scenario more expensive-to-recompute vectors will need to be overwritten in memory and dropped from RAM.

Determining the appropriate descent order (largest subtree first) is trivial and induces a low computational overhead. When the traversal list is computed, we simply need to compare the subtree sizes of child nodes and make sure to always descend into the largest subtree first.

Priority List. For this additional optimization, we exploit a property of the SPR move technique. When a candidate subtree is pruned (removed from the currently best tree), it will be re-inserted into several branches of the tree from which it was removed to evaluate the likelihood of different placements of the candidate subtree within this tree. In the course of those insertions, the subtree itself will not be changed and only the ancestral vector at the root of the subtree will need to be accessed for computations. Hence, we maintain a dedicated list of pruned candidate subtree nodes (a unpinning priority list) that can be preferentially unpinned. Because of the design of lazy SPR moves in RAxML (similar SPR flavors are used in GARLI and PHYML) those nodes (corresponding to ancestral vectors) will not be accessed while the candidate subtree is inserted into different branches of the tree. Once this priority list is exhausted, the standard MRC recomputation strategy is applied.

Full Traversals. Full post-order traversals of the tree are required during certain phases of typical phylogenetic inference programs, for instance when optimizing global maximum likelihood model parameters (e.g., the α shape parameter of the Γ distribution or the rates in a GTR matrix) on the entire tree. Full tree traversals are also important for the post-analysis of fixed tree topologies, for instance, to estimate species divergence times. Full tree traversals represent a particular case because every inner vector of the tree needs to be visited and computed. Hence, the number of vectors that need to be computed under our memory reduction approach is exactly identical to the number of vectors that need to be computed under the standard implementation. Thus, there is no need for additional computations while a large amount of memory can be saved. While full tree traversals do not dominate run times in standard tree search algo-

rithms, they can dominate execution times in downstream analysis tools.

4 EXPERIMENTAL SETUP AND RESULTS

We have implemented the techniques described in Section 3 in RAxML-Light v1.0.4. RAxML-Light is a strapped-down dedicated version of RAxML intended for large-scale phylogenetic inferences on supercomputers. It implements a light-weight software-based checkpointing mechanism and offers fine-grained PThreads and MPI parallelizations of the PLF. It has been used to compute a tree on a dense simulated MSA with 1481 taxa and 20,000,000 sites that required 1TB of RAM and ran in parallel with MPI on 672 cores. While not yet published, RAxML-Light is available as open-source code at www.exelixis-lab.org/software.html and at <https://github.com/stamatak>. The source code of the memory saving technique presented here is available for download at www.exelixis-lab.org/software/spare.zip and is currently being integrated into the RAxML-Light development version (development snapshots available at <https://github.com/fizquierdo/vectorRecomp>).

4.1 Evaluation of Recomputation Strategies

The recomputation algorithm yields *exactly* the same log likelihood scores for the PLF as the standard algorithm. Thus, for validating the correctness of our implementation, it is sufficient to verify that the resulting trees and log likelihood scores of a ML tree search with the standard and recomputation implementations are identical. The increase of total run time depends on the number x of inner vectors that are held in memory *and* on the chosen unpinning strategy (MRC versus RANDOM). We used indelible (Fletcher and Yang, 2009) to generate simulated MSAs of 1500, 3000, and 5000 species. All experiments described in this section were conducted for these three datasets. We only show representative results for one dataset (1500 taxa, 575 base pairs).

The results for the other two simulated datasets (3000 taxa with 774 base pairs and 5000 taxa with 1074 base pairs), as well as for a biological dataset with 500 taxa, are analogous and can be found in the supplementary on-line material at www.exelixis-lab.org/publications.html.

We chose to use simulated datasets to assess per-

formance for convenience. Detailed information on indelible parameters used for dataset generation are included in the supplementary on-line material. For our purely computational work it does not matter whether simulated or real data is used.

Initially, we ran the Parsimonator program (available at www.exelixis-lab.org/software.html) to generate 10 distinct randomized stepwise addition order parsimony starting trees for each MSA. For each starting tree, we then executed a standard ML tree search with RAxML-Light 1.0.4 (sequential version with SSE3 intrinsics) and ML tree searches with the recomputation version for the two replacement strategies (MRC and RANDOM) and five different RAM reduction factors ($-x$ and $-r$ options respectively). In the recomputation version used for the evaluation, both the *Largest subtree first* and the *Priority List* optimizations were activated. All experiments were executed on a 48-core AMD system with 256GB RAM. For all runs, RAM memory usage was measured every 600 seconds with `top`.

Figure 4 shows the corresponding decrease in RAM usage. Values in Figure 4 correspond to maximum observed RAM usage values.

Figure 3 depicts the run time increase as a function of available space for storing ancestral probability vectors. Clearly, the MRC strategy outperforms the RANDOM strategy and the induced run-time overhead, even for a reduction of available RAM space to only 10% is surprisingly small (approximately 40%). This slowdown is acceptable, considering that instead of analyzing a large dataset on a machine with 256GB, a significantly smaller and less expensive system with, for instance, 32GB RAM will be sufficient. Given the checkpointing capabilities of RAxML-Light, the increase in run times does not represent a problem.

4.2 Evaluation of Traversal Overhead

In order to evaluate the overhead of the extended (larger) tree traversals due to the required additional ancestral probability vector computations, we modified the source code to count the number of ancestral vector computations. We distinguish between three cases with different recomputation costs (see Section 3). For each case, there exists a dedicated PLF implementation in RAxML.

Tip/Tip Both child nodes are tips.

Tip/Vector One child node is a tip, and the other is an ancestral vector (subtree).

Vector/Vector Both child nodes are ancestral vectors.

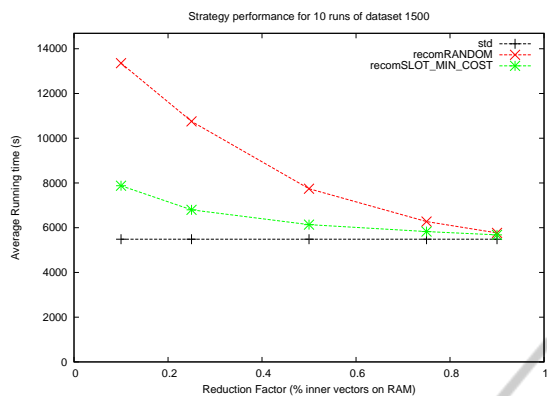


Figure 3: Different replacement strategies. The dataset was run with RAM allocations of 10%, 25%, 50%, 75%, and 90%, of the total required memory for storing all probability vectors. Run times are averaged across 10 searches with distinct starting trees.

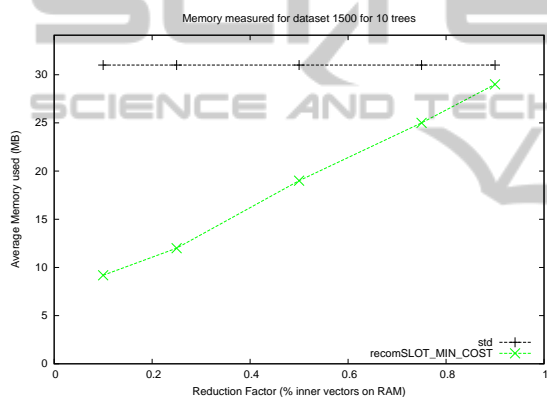


Figure 4: Overall RAM usage when allocating only 10%, 25%, 50%, 75%, and 90%, of the required ancestral probability vectors.

Table 1 shows a dramatic, yet desired increase, in the number of Tip/Tip vector computations for the MRC strategy. However, the amount of the slowest type of ancestral node computations [Vector/Vector] is only increased by 0.16% compared to the standard implementation.

4.3 Evaluation of Full Tree Traversals

We created a simple test case that parses an input tree and conducts 20 full tree traversals on the given tree. This dedicated test code is also available at <https://github.com/fizquierdo/vectorRecomp/>. We used the aforementioned starting trees and the 500, 1500, and 5000 taxon datasets. Each run was repeated 5 times and we averaged running times. All runs returned exactly the same likelihood scores.

Table 2 indicates that, even for very small R values (fraction of inner vectors allocated in memory) the run

time overhead is negligible compared to the standard implementation.

5 CONCLUSIONS AND FUTURE WORK

We have presented a generic strategy for the exact computation of log likelihood scores and ML tree searches with significantly reduced memory requirements. The additional computational cost incurred by the larger number of required ancestral vector recomputations is comparatively low when an appropriate vector replacement strategy is deployed. This will allow for computing the PLF on larger datasets than ever before, especially when the limiting factor is available RAM. In fact, the memory versus additional computations trade-off can be adapted by the users via a command line switch to fit their computational resources. We also show that, the minimum number of ancestral probability vectors for computing the PLF that need to be kept in memory for a tree with n taxa is $\log_2(n) + 2$. This result may be particularly interesting for designing equally fast, but highly memory-efficient phylogenetic post-analysis tools that rely on full tree traversals.

Furthermore, the concepts presented here can be applied to all PLF-based programs such as GARLI, PHYML, MrBayes, etc. On the software engineering side, the implementation of the recomputation strategy in RAxML is encapsulated in such a way, that it can be combined with any other memory saving techniques such as out-of-core computations, Subtree Equality Vectors, or the use of lossless compression algorithms for ancestral probability vectors.

We plan to further refine/tune the MRC strategy and fully integrate it into the up-to-date release of RAxML-Light v1.0.5 and the already existing memory saving techniques. This will allow to infer trees and compute likelihood scores on huge datasets, that would previously have required a supercomputer, on a single multi-core system.

ACKNOWLEDGEMENTS

Fernando Izquierdo-Carrasco is funded by the German Science Foundation (DFG). The authors would like to thank Simon Berger for pointing out and discussing favorable trade-offs regarding recomputation of vectors and the potential use of lossless compression algorithms to decrease memory requirements.

Table 1: Frequency of ancestral vector cases for the standard implementation and the recomputation strategies (50% of ancestral vectors allocated).

Strategy	Tip/Tip	Tip/Vector	Vector/Vector	Total	Runtime (s)
Standard	11,443,484	57,884,490	76,325,233	145,653,207	5678
MRC (0.5)	20,368,957	61,224,562	76,444,874	158,038,393	6453
Random (0.5)	37,778,575	85,303,730	104,398,910	227,481,215	7999

Table 2: Average run times in seconds for 20 full traversals averaged across 5 runs.

Dataset	Standard	R:=0.1	R:=0.9
500	0.122	0.121	0.130
1500	0.430	0.430	0.434
5000	2.402	2.412	2.438

REFERENCES

- Barkan, E., Biham, E., and Shamir, A. (2006). Rigorous bounds on cryptanalytic time/memory tradeoffs. In *In Advances in Cryptology CRYPTO 2006, volume 4117 of LNCS*, pages 1–21. Springer-Verlag.
- Berger, S. and Stamatakis, A. (2010). Accuracy and performance of single versus double precision arithmetics for Maximum Likelihood Phylogeny Reconstruction. *Springer Lecture Notes in Computer Science*, 6068:270–279.
- Dri, P. and Galil, Z. (1984). A time-space tradeoff for language recognition. *Theory of Computing Systems*, 17:3–12. 10.1007/BF01744430.
- Felsenstein, J. (1981). Evolutionary trees from DNA sequences: a maximum likelihood approach. *J. Mol. Evol.*, 17:368–376.
- Fletcher, W. and Yang, Z. (2009). INDELible: a flexible simulator of biological sequence evolution. *Molecular biology and evolution*, 26(8):1879–1888.
- Guindon, S., Dufayard, J., Lefort, V., Anisimova, M., Hordijk, W., and Gascuel, O. (2010). New algorithms and methods to estimate maximum-likelihood phylogenies: assessing the performance of PhyML 3.0. *Systematic biology*, 59(3):307.
- Izquierdo-Carrasco, F. and Stamatakis, A. (2011). Computing the phylogenetic likelihood function out-of-core. In *In Proceedings of the IPDPS-HiCOMB 2011*.
- Martin Simonsen, T. M. and Pedersen, C. N. S. (2010). Building very large neighbour-joining trees. Proceedings of Bioinformatics 2010, to appear in Springer bioinformatics lecture notes.
- Stamatakis, A. (2006). RAxML-VI-HPC: maximum likelihood-based phylogenetic analyses with thousands of taxa and mixed models. *Bioinformatics*, 22(21):2688–2690.
- Stamatakis, A. (2011). *Phylogenetic Search Algorithms for Maximum Likelihood*, pages 547–577. John Wiley & Sons, Inc.
- Stamatakis, A. and Alachiotis, N. (2010). Time and memory efficient likelihood-based tree searches on phylogenomic alignments with missing data. *Bioinformatics*, 26(12):i132.
- Stamatakis, A. and Ott, M. (2008). Exploiting fine-grained parallelism in the phylogenetic likelihood function with mpi, pthreads, and openmp: a performance study. *Pattern Recognition in Bioinformatics*, pages 424–435.
- Vitter, J. S. (2008). Algorithms and data structures for external memory. *Found. Trends Theor. Comput. Sci.*, 2(4):305–474.
- Wheeler, T. (2009). Large-scale neighbor-joining with ninja. In Salzberg, S. and Warnow, T., editors, *Algorithms in Bioinformatics*, volume 5724 of *Lecture Notes in Computer Science*, pages 375–389. Springer Berlin / Heidelberg.
- Xu, J. and Lipton, R. J. (2005). On fundamental tradeoffs between delay bounds and computational complexity in packet scheduling algorithms. *IEEE/ACM Trans. Netw.*, 13:15–28.
- Yang, Z. (1994). Maximum likelihood phylogenetic estimation from DNA sequences with variable rates over sites. *J. Mol. Evol.*, 39:306–314.
- Zwickl, D. (2006). *Genetic Algorithm Approaches for the Phylogenetic Analysis of Large Biological Sequence Datasets under the Maximum Likelihood Criterion*. PhD thesis, University of Texas at Austin.