

Algorithms and
Data Structures

Jeffrey Vitter
Editor

Skip Lists: A Probabilistic Alternative to Balanced Trees

Skip lists are data structures that use probabilistic balancing rather than strictly enforced balancing. As a result, the algorithms for insertion and deletion in skip lists are much simpler and significantly faster than equivalent algorithms for balanced trees.

William Pugh

Binary trees can be used for representing abstract data types such as dictionaries and ordered lists. They work well when the elements are inserted in a random order. Some sequences of operations, such as inserting the elements in order, produce degenerate data structures that perform very poorly. If it were possible to randomly permute the list of items to be inserted, trees would work well with high probability for any input sequence. In most cases queries must be answered on-line, so randomly permuting the input is impractical. *Balanced* tree algorithms rearrange the tree as operations are performed to maintain certain balance conditions and assure good performance.

Skip lists are a probabilistic alternative to balanced trees. Skip lists are balanced by consulting a random number generator. Although skip lists have bad worst-case performance, no input sequence consistently produces the worst-case performance (much like quicksort when the pivot element is chosen randomly). It is very unlikely a skip list data structure will be significantly unbalanced (e.g., for a dictionary of more than 250 elements, the chance that a search will take more than three-times the expected time is less than one in a million). Skip lists have balance properties similar to that of search trees built by random insertions, yet do not require insertions to be random.

It is easier to balance a data structure probabilistically than to explicitly maintain the balance. For many applications, skip lists are a more natural representation than trees, and they lead to simpler algorithms. The simplicity of skip list algorithms makes them easier to implement and provides significant constant factor speed improvements over balanced tree and

self-adjusting tree algorithms. Skip lists are also very space efficient. They can easily be configured to require an average of $1\frac{1}{2}$ pointers per element (or even less) and do not require balance or priority information to be stored with each node.

SKIP LISTS

We might need to examine every node of the list when searching a linked list (Figure 1a). If the list is stored in sorted order and every other node of the list also has a pointer to the node two ahead of it in the list (Figure 1b), we have to examine no more than $\lceil n/2 \rceil + 1$ nodes (where n is the length of the list). Also giving every fourth node a pointer four ahead (Figure 1c) requires that no more than $\lceil n/4 \rceil + 2$ nodes be examined. If every (2^i) th node has a pointer 2^i nodes ahead (Figure 1d), the number of nodes that must be examined can be reduced to $\lceil \log_2 n \rceil$ while only doubling the number of pointers. This data structure could be used for fast searching, but insertion and deletion would be impractical.

A node that has k forward pointers is called a *level k* node. If every (2^i) th node has a pointer 2^i nodes ahead, then levels of nodes are distributed in a simple pattern: 50 percent are level 1, 25 percent are level 2, 12.5 percent are level 3 and so on. What would happen if the levels of nodes were chosen randomly, but in the same proportions (e.g., as in Figure 1e)? A node's i th forward pointer, instead of pointing 2^{i-1} nodes ahead, points to the next node of level i or higher. Insertions or deletions would require only local modifications; the level of a node, chosen randomly when the node is inserted, need never change. Some arrangements of levels would give poor execution times, but we will see that such arrangements are rare. Because these data structures are linked lists with extra pointers that skip over intermediate nodes, I named them skip lists.

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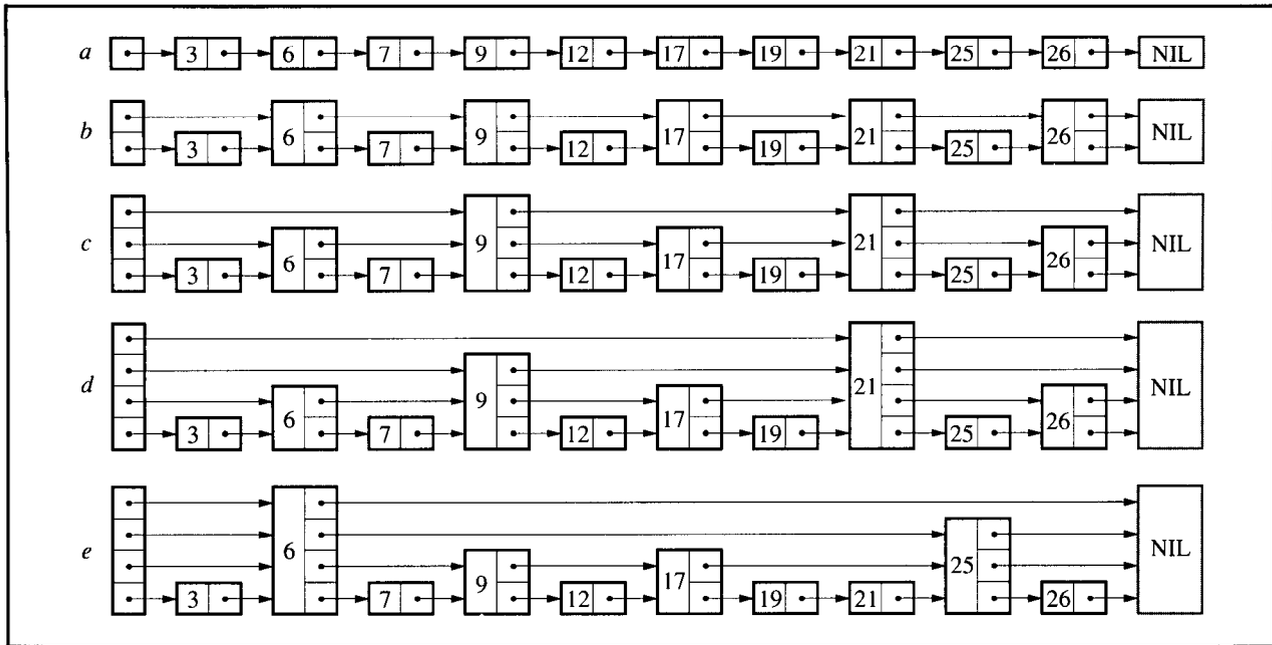


FIGURE 1. Linked Lists with Additional Pointers

SKIP LIST ALGORITHMS

This section describes how to search for algorithms and to insert and delete elements in a dictionary or symbol table. The Search operation returns the contents of the value associated with the desired key or *failure* if the key is not present. The Insert operation associates a specified key with a new value (inserting the key if it had not already been present). The Delete operation deletes the specified key. It is easy to support additional operations such as “find the minimum key” or “find the next key.”

Each element is represented by a node, the level of which is chosen randomly when the node is inserted without regard for the number of elements in the data structure. A *level i* node has *i forward* pointers, indexed 1 through *i*. We do not need to store the level of a node in the node. Levels are capped at some appropriate constant *MaxLevel*. The level of a list is the maximum level currently in the list (or 1 if the list is empty). The header of a list has forward pointers at levels one through *MaxLevel*. The forward pointers of the header at levels higher than the current maximum level of the list point to NIL.

Initialization

An element NIL is allocated and given a key greater than any legal key. All levels of all skip lists are terminated with NIL. A new list is initialized so that the level of the list is equal to 1 and all forward pointers of the list's header point to NIL.

Search Algorithm

We search for an element by traversing forward pointers that do not overshoot the node containing the element being searched for (Figure 2). When no more progress can be made at the current level of forward

```

Search(list, searchKey)
  x := list→header
  -- loop invariant: x→key < searchKey
  for i := list→level downto 1 do
    while x→forward[i]→key < searchKey do
      x := x→forward[i]
  -- x→key < searchKey ≤ x→forward[1]→key
  x := x→forward[1]
  if x→key = searchKey then return x→value
  else return failure

```

FIGURE 2. Skip List Search Algorithm

pointers, the search moves down to the next level. When we can make no more progress at level 1, we must be immediately in front of the node that contains the desired element (if it is in the list).

Insertion and Deletion Algorithms

To insert or delete a node, we simply search and splice, as shown in Figure 3. Figure 4 gives algorithms for insertion and deletion. A vector *update* is maintained so that when the search is complete (and we are ready to perform the splice), *update*[*i*] contains a pointer to the rightmost node of level *i* or higher that is to the left of the location of the insertion/deletion.

If an insertion generates a node with a level greater than the previous maximum level of the list, we update the maximum level of the list and initialize the appropriate portions of the update vector. After each deletion, we check to see if we have deleted the maximum element of the list and if so, decrease the maximum level of the list.

Choosing a Random Level

Initially, we discussed a probability distribution where

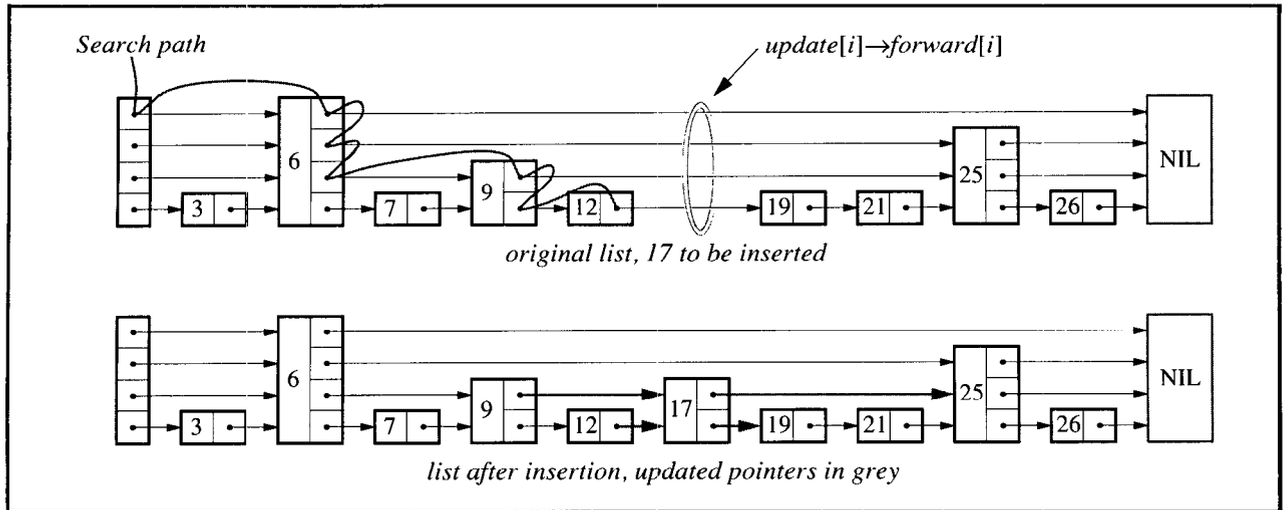


FIGURE 3. Pictorial Description of Steps Involved in Performing an Insertion

```

Insert(list, searchKey, newValue)
  local update[1..MaxLevel]
  x := list→header
  for i := list→level downto 1 do
    while x→forward[i]→key < searchKey do
      x := x→forward[i]
      -- x→key < searchKey ≤ x→forward[i]→key
      update[i] := x
    x := x→forward[1]
  if x→key = searchKey then x→value := newValue
  else
    newLevel := randomLevel()
    if newLevel > list→level then
      for i := list→level + 1 to newLevel do
        update[i] := list→header
      list→level := newLevel
    x := makeNode(newLevel, searchKey, value)
    for i := 1 to newLevel do
      x→forward[i] := update[i]→forward[i]
      update[i]→forward[i] := x

Delete(list, searchKey)
  local update[1..MaxLevel]
  x := list→header
  for i := list→level downto 1 do
    while x→forward[i]→key < searchKey do
      x := x→forward[i]
      update[i] := x
    x := x→forward[1]
  if x→key = searchKey then
    for i := 1 to list→level do
      if update[i]→forward[i] ≠ x then break
      update[i]→forward[i] := x→forward[i]
    free(x)
  while list→level > 1 and
    list→header→forward[list→level] = NIL do
    list→level := list→level - 1
    
```

FIGURE 4. Skip List Insertion and Deletion Algorithms

```

randomLevel()
  newLevel := 1
  -- random() returns a random value in [0..1)
  while random() < p do
    newLevel := newLevel + 1
  return min(newLevel, MaxLevel)
  
```

FIGURE 5. Algorithm to Calculate a Random Level

half of the nodes that have level i pointers also have level $i + 1$ pointers. To get away from magic constants, we say that a fraction p of the nodes with level i pointers also have level $i + 1$ pointers (see $p = 1/2$ for our original discussion). Levels are generated randomly by an algorithm equivalent to the one in Figure 5. Levels are generated without reference to the number of elements in the list.

At What Level do We Start a Search? Defining $L(n)$

In a skip list of 16 elements generated with $p = 1/2$, we might happen to have 9 elements of level 1; 3 elements of level 2; 3 elements of level 3; and 1 element of level 14 (this would be very unlikely, but it could happen). How should we handle this? If we use the standard algorithm and start our search at level 14, we will do a lot of useless work.

Where should we start the search? Our analysis suggests that ideally we would start a search at the level L where we expect $1/p$ nodes. This happens when $L = \log_{1/p} n$. Since we will be referring frequently to this formula, we will use $L(n)$ to denote $\log_{1/p} n$.

There are a number of solutions to the problem of deciding how to handle the case where there is an element with an unusually large level in the list.

- Don't worry, be happy. Simply start a search at the highest level present in the list. As we will see in our

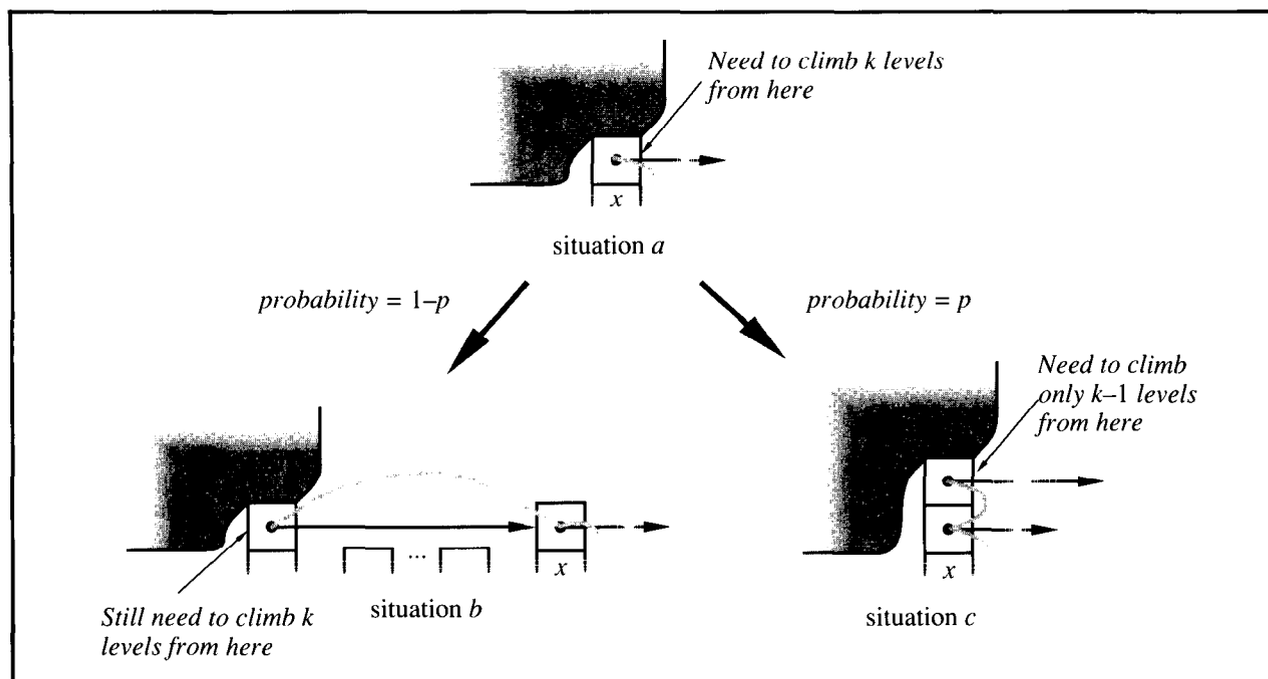


FIGURE 6. Possible Situations in Backwards Traversal of the Search Path

analysis, the probability that the maximum level in a list of n elements is significantly larger than $L(n)$ is very small. Starting a search at the maximum level in the list does not add more than a small constant to the expected search time. This is the approach used in the algorithms described in this paper.

- *Use less than you are given.* Although an element may contain room for 14 pointers, we do not need to use all 14. We can choose to utilize only $L(n)$ levels. There are a number of ways to implement this, but they all complicate the algorithms and do not noticeably improve performance, so this approach is not recommended.
- *Fix the dice.* If we generate a random level that is more than one greater than the current maximum level in the list, we simply use one plus the current maximum level in the list as the level of the new node. In practice and intuitively, this change seems to work well. However, it totally destroys our ability to analyze the resulting algorithms, since the level of a node is no longer completely random. Although programmers may implement this method, purists should avoid it.

Determining *MaxLevel*

Since we can safely cap levels at $L(n)$, we should choose $MaxLevel = L(N)$ (where N is an upper bound on the number of elements in a skip list). If $p = 1/2$, using $MaxLevel = 16$ is appropriate for data structures containing up to 2^{16} elements.

ANALYSIS OF SKIP LIST ALGORITHMS

The time required to execute the Search, Delete and

Insert operations is dominated by the time required to search for the appropriate element. For the Insert and Delete operations, there is an additional cost proportional to the level of the node being inserted or deleted. The time required to find an element is proportional to the length of the search path, which is determined by the pattern in which elements with different levels appear as we traverse the list.

Probabilistic Philosophy

The structure of a skip list is determined only by the number of elements in the skip list and the results of consulting the random number generator. The sequence of operations that produced the current skip list does not matter. We assume an adversarial user does not have access to the levels of nodes; otherwise, he could create situations with worst-case running times by deleting all nodes that were not level 1.

The probabilities of poor running times for successive operations on the same data structure are NOT independent; two successive searches for the same element will both take exactly the same time.

Analysis of Expected Search Cost

We analyze the search path backwards, travelling up and to the left. Although the levels of nodes in the list are known and fixed when the search is performed, we act as if the level of a node is being determined only when it is observed while backtracking the search path.

At any particular point in the climb, we are at a situation similar to situation *a* in Figure 6—we are at the i th forward pointer of a node x and we have no knowledge about the levels of nodes to the left of x or about the level of x , other than that the level of x must be at least i . Assume the x is not the header (this is equivalent to assuming the list extends infinitely to the left). If the level of x is equal to i , then we are in

situation *b*. If the level of *x* is greater than *i*, then we are in situation *c*. The probability that we are in situation *c* is *p*. Each time we are in situation *c*, we climb up a level. Let $C(k)$ = the expected cost (i.e., length) of a search path that climbs up *k* levels in an infinite list:

$$C(0) = 0$$

$$C(k) = (1 - p) \text{ (cost in situation } b) + p \text{ (cost in situation } c)$$

By substituting and simplifying, we get:

$$C(k) = (1 - p)(1 + C(k)) + p(1 + C(k - 1))$$

$$C(k) = 1/p + C(k - 1)$$

$$C(k) = k/p$$

Our assumption that the list is infinite is a pessimistic assumption. When we bump into the header in our backwards climb, we simply climb up it, without performing any leftward movements. This gives us an upper bound on $(L(n) - 1)/p$ on the expected length of the path that climbs from level 1 to level $L(n)$ in a list of *n* elements.

We use this analysis go up to level $L(n)$ and use a

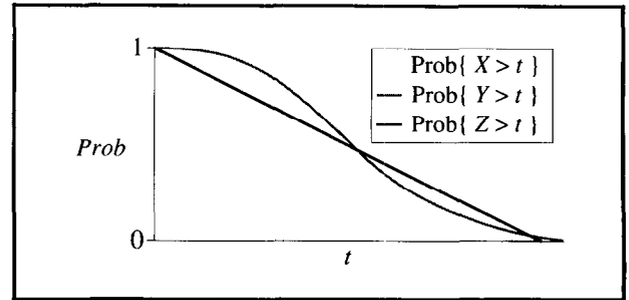


FIGURE 7. Plots of Three Probability Distributions

different analysis technique for the rest of the journey. The number of leftward movements remaining is bounded by the number of elements of level $L(n)$ or higher in the entire list, which has an expected value of $1/p$.

We also move upwards from level $L(n)$ to the maximum level in the list. The probability that the maximum level of the list is a greater than *k* is equal to $1 - (1 - p^k)^n$, which is at most np^k . The expected max-

PROBABILISTIC ANALYSIS

In addition to analyzing the expected performance of skip lists, we can analyze their probabilistic performance. This capability will allow us to calculate the probability of an operation taking longer than a specified time. Since this analysis is based on the same concept as our analysis of the expected cost, the reader should understand that analysis first.

A random variable has a fixed but unpredictable value and a predictable probability distribution and average. If *X* is a random variable, $\text{Prob}\{X = t\}$ denotes the probability that *X* equals *t* and $\text{Prob}\{X > t\}$ denotes the probability that *X* is greater than *t*. For example, if *X* is the number obtained by throwing an unbiased die, $\text{Prob}\{X > 3\} = 1/2$.

It is often preferable to find simple upper bounds on values whose exact value is difficult to calculate. To discuss upper bounds on random variables, we need to define a partial ordering and equality on the probability distributions of non-negative random variables.

Definitions ($=_{\text{prob}}$ and \leq_{prob}). Let *X* and *Y* be non-negative independent random variables (typically, *X* and *Y* would denote the time to execute algorithms A_X and A_Y). We define $X \leq_{\text{prob}} Y$ to be true if and only if for any value *t*, the probability that *X* exceeds *t* is not more than the probability that *Y* exceeds *t*. More

formally:

$$X \leq_{\text{prob}} Y \quad \text{iff } \forall t, \text{Prob}\{X > t\} \leq \text{Prob}\{Y > t\}$$

and

$$X =_{\text{prob}} Y \quad \text{iff } \forall t, \text{Prob}\{X > t\} = \text{Prob}\{Y > t\}. \quad \square$$

For example, the graph in Figure 7 shows the probability distribution of three random variables *X*, *Y* and *Z*. Since the probability distribution curve for *X* is completely under the curves for *Y* and *Z*, $X \leq_{\text{prob}} Y$ and $X \leq_{\text{prob}} Z$. Since the probability curves for *Y* and *Z* intersect, neither $Y \leq_{\text{prob}} Z$ nor $Z \leq_{\text{prob}} Y$. Since the expected value of a random variable *X* is simply the area under the curve $\text{Prob}\{X > t\}$, if $X \leq_{\text{prob}} Y$ then the average of *X* is less than or equal to the average of *Y*.

We make use of two probability distributions:

Definition (binomial distributions— $B(t, p)$). Let *t* be a non-negative integer and *p* be a probability. The term $B(t, p)$ denotes a random variable equal to the number of successes seen in a series of *t* independent random trials where the probability of a success in a trial is *p*. The average and variance of $B(t, p)$ are tp and $tp(1 - p)$ respectively.

Definition (negative binomial distributions— $NB(s, p)$). Let *s* be a non-negative integer and *p* be a probability. The term $NB(s, p)$ denotes a random variable equal to the number of failures seen before the *s*th success in a series of random independent trials where the probability of a success in a trial is *p*. The average and variance of $NB(s, p)$ are $s(1 - p)/p$ and $s(1 - p)/p^2$ respectively.

imum level is therefore at most $L(n) + 1/(1 - p)$. Putting our results together, we find that the total expected cost to climb out of a list of n elements is at most $L(n)/p + 1/(1 - p)$, which is $O(\log n)$.

Our result is an analysis of the *length* of the search path. The number of comparisons required is one plus the length of the search path (a comparison is performed for each position in the search path, the length of the search path is the number of hops between positions in the search path).

It is also possible to analyze the probability distribution of search costs. The probabilistic analysis is somewhat more complicated (see box). From the probabilistic analysis, we can calculate an upper bound on the probability that the actual cost of a search exceeds the expected cost by more than a specified ratio. Some results of this analysis are shown in Figure 8.

Choosing p

Table I gives the relative time and space requirements for different values of p . Decreasing p also increases the variability of running times. If $1/p$ is a power of 2, it will be easy to generate a random level from a stream

Probabilistic Analysis of Search Cost

The number of leftward movements we need to make before we move up a level (in an infinite list) has a negative binomial distribution: it is the number of failures (situations b 's) we see before we see the first success (situation c) in a series of independent random trials where the probability of success is p . Using the probabilistic notation introduced here:

Cost to climb one level in an infinite list
 $=_{\text{prob}} 1 + NB(1, p)$.

We can sum the costs of climbing each level to get the total cost to climb up to level $L(n)$:

Cost to climb to level $L(n)$ in an infinite list
 $=_{\text{prob}} (L(n) - 1) + NB(L(n) - 1, p)$.

Our assumption that the list is infinite is a pessimistic assumption:

Cost to climb to level $L(n)$ in a list of n elements
 $\leq_{\text{prob}} (L(n) - 1) + NB(L(n) - 1, p)$.

Once we have climbed to level $L(n)$, the number of leftward movements is bounded by the number of elements of level $L(n)$ or greater in a list of n elements. The number of elements of level $L(n)$ or greater in a list of n elements is a random variable of the form $B(n, 1/np)$.

Let M be a random variable corresponding to the maximum level in a list of n elements. The probability that the level of a node is greater than k is p^k , so $\text{Prob}\{M > k\} = 1 - (1 - p^k)^n < np^k$. Since $np^k = p^{k-L(n)}$ and $\text{Prob}\{NB(1, 1 - p) + 1 > i\} = p^i$, we get a

of random bits (it requires an average of $(\log_2 1/p)/(1 - p)$ random bits to generate a random level). Since some of the constant overheads are related to $L(n)$ (rather than $L(n)/p$), choosing $p = 1/4$ (rather than $1/2$) slightly improves the constant factors of the speed of the algorithms as well. I suggest that a value of $1/4$ be used for p unless the variability of running times is a primary concern, in which case p should be $1/2$.

The expected total time for a sequence of operations is equal to the sum of the expected times of each of the operations in the sequence. Thus, the expected time for any sequence of m searches in a data structure that contains n elements is $O(m \log n)$. However, the pattern of searches affects the probability distribution of the actual time to perform the entire sequence of operations.

If we search for the same item twice in the same data structure, both searches will take exactly the same amount of time. Thus the variance of the total time will be four times the variance of a single search. If the search times for two elements are independent, the variance of the total time is equal to the sum of the variances of the individual searches. Searching

probabilistic upper bound of $M \leq_{\text{prob}} L(n) + NB(1, 1 - p) + 1$. Note that the average of $L(n) + NB(1, 1 - p) + 1$ is $L(n) + 1/(1 - p)$.

This gives a probabilistic upper bound on the cost once we have reached level $L(n)$ of $B(n, 1/np) + (L(n) + NB(1, 1 - p) + 1) - L(n)$. Combining our results to get a probabilistic upper bound on the total length of the search path (i.e., cost of the entire search):

total cost to climb out of a list of n elements
 $\leq_{\text{prob}} (L(n) - 1) + NB(L(n) - 1, p)$
 $+ B(n, 1/np) + NB(1, 1 - p) + 1$

The expected value of our upper bound is equal to

$$(L(n) - 1) + \frac{(L(n) - 1)(1 - p)}{p} + \frac{1}{p} + \frac{p}{1 - p} + 1$$

$$= \frac{L(n)}{p} + \frac{1}{1 - p}$$

which is the same as our previously calculated upper bound on the expected cost of a search. The variance of our upper bound is

$$\frac{(L(n) - 1)(1 - p)}{p^2} + \frac{(1 - 1/np)}{p} + \frac{p}{(1 - p)^2}$$

$$< \frac{(1 - p)L(n)}{p^2} + \frac{p}{(1 - p)^2} + \frac{2p - 1}{p^2}.$$

Figure 8 shows a plot of an upper bound on the probability of an actual search taking substantially longer than average, based on our probabilistic upper bound.

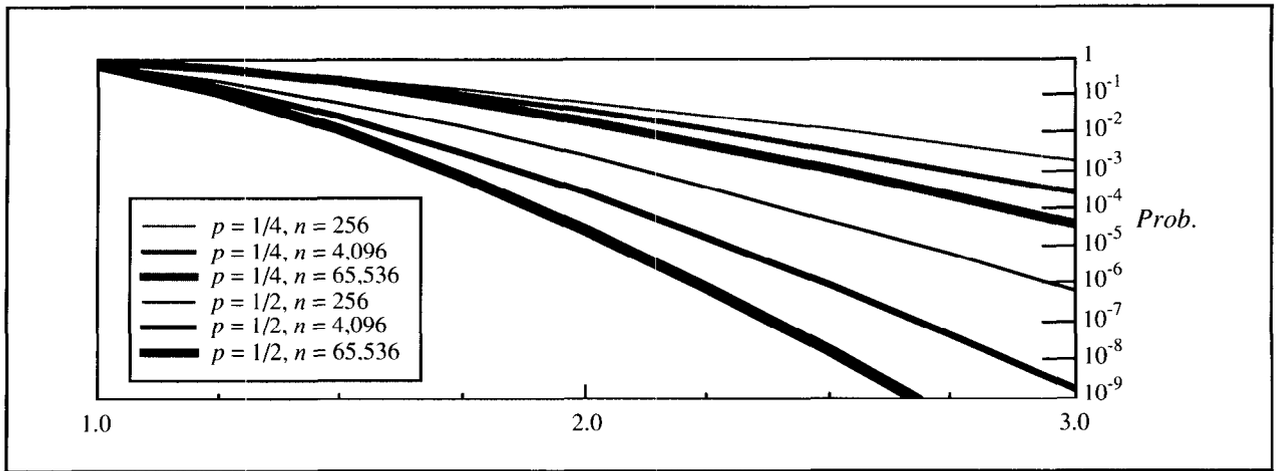


FIGURE 8. This graph shows a plot of an upper bound on the probability of a search taking substantially longer than expected. The vertical axes show the probability that the length of the search path exceeds the average length by more than the ratio on the horizontal axis. For example, for $p = 1/2$ and $n = 4096$, the probability that the search path will be more than three times the expected length is less than one in 200 million. This graph was calculated using our probabilistic upper bound.

TABLE I. Relative Search Speed and Space Requirements Depending on the Value of p .

p	Normalized search times (i.e., normalized $L(n)/p$)	Avg. # of pointers per mode (i.e., $1/(1-p)$)
$1/2$	1	2
$1/e$	0.94...	1.58...
$1/4$	1	1.33...
$1/8$	1.33...	1.14...
$1/16$	2	1.07...

for the same element over and over again maximizes the variance.

ALTERNATIVE DATA STRUCTURES

Balanced trees (e.g., AVL trees [5, 13]) and self-adjusting trees [11] can be used for the same problems as skip lists. All three techniques have performance bounds of the same order. A choice among these schemes involves several factors: the difficulty of implementing the algorithms, constant factors, type of bound (amortized, probabilistic or worst-case), and performance on a non-uniform distribution of queries.

For most applications, implementers generally agree skip lists are significantly easier to implement than either balanced tree algorithms or self-adjusting tree algorithms.

Constant Factors

Constant factors can make a significant difference in the practical application of an algorithm. This is particularly true for sub-linear algorithms. For example, assume that algorithms A and B both require $O(\log n)$ time to process a query, but that B is twice as fast as A : in the time algorithm A takes to process a query on a data set of size n , algorithm B can process a query on a data set of size n^2 .

There are two important but qualitatively different contributions to the constant factors of an algorithm. First, the inherent complexity of the algorithm places a

lower bound on any implementation. Self-adjusting trees are continuously rearranged as searches are performed; this imposes a significant overhead on any implementation of self-adjusting trees. Skip list algorithms seem to have very low inherent constant-factor overheads: the inner loop of the deletion algorithm for skip lists compiles to just six instructions on the 68020.

Second, if the algorithm is complex, programmers are deterred from implementing optimizations. For example, balanced tree algorithms are normally described using recursive insert and delete procedures, since that is the most simple and intuitive method of describing the algorithms. A recursive insert or delete procedure incurs a procedure call overhead. By using nonrecursive insert and delete procedures, some of this overhead can be eliminated. However, the complexity of nonrecursive algorithms for insertion and deletion in a balanced tree is intimidating and this complexity deters most programmers from eliminating recursion in these routines. Skip list algorithms are already nonrecursive and they are simple enough that programmers are not deterred from performing optimizations.

Table II compares the performance of implementations of skip lists and four other techniques. All implementations were optimized for efficiency. The AVL tree algorithms were written by James Macropol of Contel and based on those in [13]. The 2-3 tree algorithms are based on those presented in [1]. Several other existing balanced tree packages were timed and found to be

much slower than the results presented. The self-adjusting tree algorithms are based on those presented in [11]. The times in this table reflect the CPU time on a Sun-3/60 to perform an operation in a data structure containing 2^{16} elements with integer keys. The values in parenthesis show the results relative to the skip list times. The times for insertion and deletion do not include the time for memory management (e.g. in C programs, calls to *malloc* and *free*).

It should be noted that skip lists perform more comparisons than other methods (the skip list algorithms presented here require an average of $L(n)/p + 1/(1-p) + 1$ comparisons). For tests using real numbers as keys, skip lists were slightly slower than the nonrecursive AVL tree algorithms and search in a skip list was slightly slower than search in a 2-3 tree (insertion and deletion using the skip-list algorithms was still faster than using the recursive 2-3 tree algorithms). If comparisons are very expensive, it is possible to change the algorithms so that we never compare the search key against the key of a node more than once during a search. For $p = 1/2$, this produces an upper bound on the expected number of comparisons of $7/2 + 3/2 \log_2 n$. This modification is discussed in [8].

Type of Performance Bound

These three classes of algorithm have different kinds of performance bounds. Balanced trees have worst-case time bounds, self-adjusting trees have amortized time bounds and skip lists have probabilistic time bounds. With self-adjusting trees, an individual operation can take $O(n)$ time, but the time bound always holds over a long sequence of operations. For skip lists, any operation or sequence of operations can take longer than expected, although the probability of any operation taking significantly longer than expected is negligible.

In certain real-time applications, we must be sure that an operation will be completed within a certain time bound. For such applications, self-adjusting trees may be undesirable, since they can take significantly longer on an individual operation than expected (e.g., an individual search can take $O(n)$ time instead of $O(\log n)$ time). For real-time systems, skip lists may be usable if an adequate safety margin is provided: the chance that a search in a skip list containing 1000 elements takes more than five-times the expected time is about 1 in 10^{18} .

Non-Uniformed Query Distributions

Self-adjusting trees have the property that they adjust to nonuniform query distributions. Since skip lists are faster than self-adjusting trees by a significant constant factor when a uniform query distribution is encountered, self-adjusting trees are faster than skip lists only for highly skewed distributions. We could attempt to devise self-adjusting skip lists. However, there seems little practical motivation to tamper with the simplicity and fast performance of skip lists; in an application where highly skewed distributions are expected, either self-adjusting trees or a skip list augmented by a cache may be preferable [9].

ADDITIONAL WORK ON SKIP LISTS

I have described a set of algorithms that allow multiple processors to concurrently update a skip list in shared memory [7]. These algorithms are much simpler than concurrent balanced tree algorithms. They allow an unlimited number of readers and n busy writers in a skip list of n elements with very little lock contention.

Using skip lists, makes it easy to perform the kinds of operations you might wish to do with a balanced tree such as use search fingers, merge skip lists and allow ranking operations (e.g., determine the k th element of a skip list [8]).

Tom Papadakis, Ian Munro and Patricio Poblette [6] have done an exact analysis of the expected search time in a skip list. The upper bound described in this paper is close to their exact bound; the techniques they needed to use to derive an exact analysis are very complicated and sophisticated. Their exact analysis shows that for $p = 1/2$ and $p = 1/4$, the upper bound given in this article on the expected cost of a search does not exceed two comparisons more than the exact expected cost.

I have adapted the idea of probabilistic balancing to some other problems arising both in data structures and in incremental computation [10]. We can generate the level of a node based on the result of applying a hash function to the element (as opposed to using a random number generator). This results in a scheme where for any set S , there is a unique data structure that represents S and with high probability the data structure is approximately balanced. If we combine this idea with an *applicative* (i.e., persistent) probabilistically balanced data structure and a scheme such as hashed-consing [2] that allows constant-time structural equality tests of ap-

TABLE II. Timings of Implementations of Different Algorithms

Implementation	Search Time	Insertion Time	Deletion Time
<i>Skip lists</i>	0.051 msec(1.0)	0.065 msec(1.0)	0.059 msec(1.0)
<i>non-recursive AVL trees</i>	0.046 msec(0.91)	0.10 msec (1.55)	0.085 msec(1.46)
<i>recursive 2-3 trees</i>	0.054 msec(1.05)	0.21 msec (3.2)	0.21 msec (3.65)
<i>Self-adjusting trees:</i>			
<i>top-down splaying</i>	0.15 msec (3.0)	0.16 msec (2.5)	0.18 msec (3.1)
<i>bottom-up splaying</i>	0.49 msec (9.6)	0.51 msec (7.8)	0.53 msec (9.0)

plicative data structures, we get a number of interesting properties, such as constant-time equality tests for the representations of sequences. This scheme also has a number of applications for incremental computation. Since it is awkward to make skip lists applicative, a probabilistically balanced tree scheme is used.

Related Work

James Discroll pointed out that R. Sprugnoli suggested a method of randomly balancing search trees in 1981 [12]. With Sprugnoli's approach, the state of the data structure is *not* independent of the sequence of operations that built it. This makes it much harder or impossible to formally analyze his algorithms. Sprugnoli gives empirical evidence that his algorithm has good expected performance, but no theoretical results.

A randomized data structure for ordered sets is described in [4]. However, a search using that data structure requires $O(n^{1/2})$ expected time.

Cecilia Aragon and Raimund Seidel describe a probabilistically balanced search trees scheme [3]. They discuss how to adapt their data structure to nonuniform query distributions.

CONCLUSIONS

From a theoretical point of view, there is no need for skip lists. Balanced trees can do everything skip lists can and have good worst-case time bounds (unlike skip lists). However, the difficulty of implementing balanced trees greatly restricts their use; balanced tree algorithms are rarely implemented except as part of a programming assignment in a data structures class.

Skip lists are simple data structures that can be used in place of balanced trees for most applications. Skip-list algorithms are very easy to implement, extend and modify. Skip lists are about as fast as highly optimized balanced tree algorithms and are substantially faster than casually implemented balanced tree algorithms.

Skip-list source code libraries for both C and Pascal are available for anonymous ftp from `mimsy.umd.edu`.

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ABOUT THE AUTHOR:

WILLIAM PUGH is an assistant professor in the Computer Science Department of the University of Maryland, with a joint appointment in the University of Maryland Institute for Advanced Computer Studies. His research interests involve the design and analysis of practical data structures and algorithms, both for general problems and for problems arising in the implementation of programming languages and environments. Author's Present Address: Computer Science Department, University of Maryland, College Park, MD 20742. `pugh@cs.umd.edu`.

CORRIGENDUM

The following change should be made to the article "The Art of Navigating through Hypertext" (*Communications*, March 1990, p. 296). The instructions at the end of the article for downloading the hypertext system should be changed as follows:

Use the directory name card instead of the name hypercard.

This change in the info-mac archives and MACARCH server occurred during the period between the proofreading of the article (when the instructions worked as printed) and the publication of the March issue. The author apologizes for any inconvenience but would like to take the opportunity to point out that this change in directory names is typical for the current small practical problems with electronic publication.