Structure Sharing with Binary Trees

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Many current interfaces for natural language represent syntactic and semantic information in the form of directed graphs where attributes correspond to vectors and values to nodes. There is a simple correspondence between such graphs and the matrix notation linguists traditionally use for feature sets.

![Graph Example](image)

Figure 1

The standard operation for working with such graphs is unification. The unification operation succeeds only on a pair of compatible graphs, and its result is a graph containing the information in both contributors. When a parser applies a syntactic rule, it unifies selected features of input constituents to check constraints and to build a representation for the output constituent.

Problem: proliferation of copies

When words are combined to form phrases, unification is not applied to lexical representations directly because it would result in the lexicon being changed. When a word is encountered in a text, a copy is made of its entry, and unification is applied to the copied graph, not the original one. In fact, unification in a typical parser is always preceded by a copying operation. Because of nondeterminism in parsing, it is, in general, necessary to preserve every representation that gets built. The same graph may be needed again when the parser comes back to pursue some yet unexplored option. Our experience suggests that the amount of computational effort that goes into producing these copies is much greater than the cost of unification itself. It accounts for a significant amount of the total parsing time.

In a sense, most of the copying effort is wasted. Unifications that fail typically fail for a simple reason. If it were known in advance what aspects of structures are relevant in a particular case, some effort could be saved by first considering only the crucial features of the input.

Solution: structure sharing

This paper lays out one strategy that has turned out to be very useful in eliminating much of the wasted effort. Our version of the basic idea is due to Martin Kay. It has been implemented in slightly different ways by Kay in Interlisp-D and by Lauri Karttunen in Zeta Lisp. The basic idea is to minimize copying by allowing graphs share common parts of their structure.

This version of structure sharing is based on four related ideas:
• Binary trees as a storage device for feature graphs
• "Lazy" copying
• Relative indexing of nodes in the tree
• Strategy for keeping storage trees as balanced as possible

Binary trees

Our structure-sharing scheme depends on represented feature sets as binary trees. A tree consists of cells that have a content field and two pointers which, if not empty, point to a left and a right cell respectively. For example, the content of the feature set and the corresponding directed graph in Figure 1 can be distributed over the cells of a binary tree in the following way.

![Figure 2](image)

The index of the top node is 1; the two cells below have indices 2 and 3. In general, a node whose index is n may be the parent of cells indexed 2n and 2n + 1. Each cell contains either an atomic value or a set of pairs that associate attribute names with indices of cells where their value is stored. The assignment of values to storage cells is arbitrary; it doesn't matter which cell stores which value. Here, cell 1 contains the information that the value of the attribute cat is found in cell 2 and that of agr in cell 3. This is a slight simplification. As we shall shortly see, when the value in a cell involves a reference to another cell, that reference is encoded as a relative index.

The method of locating the cell that corresponds to a given index takes advantage of the fact that the tree branches in a binary fashion. The path to a node can be read off from the binary representation of its index by starting after the first 1 in this number and taking 0 to be a signal for a left turn and 1 as a mark for a right turn. For example, starting at node 1, node 5 is reached by first going down a left branch and then a right branch. This sequence of turns corresponds to the digits 01. Prefixed with 1, this is the same as the binary representation of 5, namely 101. The same holds for all indices. Thus the path to node 9 (binary 1001) would be LEFT-LEFT-RIGHT as signalled by the last three digits following the initial 1 in the binary numeral (see Figure 6).

Lazy copying

The most important advantage that the scheme minimizes the amount of copying that has to be done. In general, when a graph is copied, we duplicate only the operation that replaces copying in this scheme starts by duplicating the topmost node of the tree that contains it. The rest of the structure remains the same. It is other nodes are modified only if and when destructive changes are about to happen. For example, assume that we need another copy of the graph stored in the tree in Figure 2. This can be obtained by producing a tree which has a different root node, but shares the rest of the structure with its original. In order to keep track of which tree actually owns a given node, each node carries a numeral tag that indicates its parentage. The relationship between the original tree (generation 0) and its copy (generation 1) is illustrated in Figure 3 where the generation is separated from the index of a node by a colon.

![Figure 3](image)
If the node that we want to copy is not the topmost node of a tree, we need to duplicate the nodes along the branch leading to it.

When a tree headed by the copied node has to be changed, we use the generation tags to minimize the creation of new structure. In general, all and only the nodes on the branch that lead to the site of a destructive change or addition need to belong to the same generation as the top node of the tree. The rest of the structure can consist of old nodes. For example, suppose we add a new feature, say [gender: fem] to the value of agr in Figure 3 to yield the feature set in Figure 4.

| cat: np |
|---------|
| agr:    |
| person: 3rd |
| number: sg |
| gender: fem |

Figure 4

Furthermore, suppose that we want the change to affect only the copy but not the original feature set. In terms of the trees that we have constructed for the example in Figure 3, this involves adding one new cell to the copied structure to hold the value fem, and changing the content of cell 3 by adding the new feature to it.

The modified copy and its relation to the original is shown in Figure 5. Note that one half of the structure is shared. The copy contains only three new nodes.

From the point of view of a process that only needs to find or print out the value of particular features, it makes no difference that the nodes containing the values belong to several trees as long as there is no confusion about the structure.

Relative addressing

Accessing an arbitrary cell in a binary tree consumes time in proportion to the logarithm of the size of the structure, assuming that cells are reached by starting at the top node and using the index of the target node as an address. Another method is to use relative addressing. Relative addresses encode the shortest path between two nodes in the tree regardless of where they are are. For example, if we are at node 9 in Figure 6.a below and need to reach node 11, it is easy to see that it is not necessary to go all the way up to node 1 and then partially retrace the same path in looking up node 11. Instead, one can stop going upward at the lowest common ancestor, node 2, of nodes 9 and 11 and go down from there.

With respect to node 2, node 11 is in the same position as 7 is with respect 1. Thus the relative address of cell 11 counted from 9 is 2,7—"two nodes
up, then down as if going to node 7°. In general, relative addresses are of the form \(<\text{up, down}>\) where \(<\text{up}>\) is the number of links to the lowest common ancestor of the origin and \(<\text{down}>\) is the relative index of the target node with respect to it. Sometimes we can just go up or down on the same branch; for example, the relative address of cell 10 seen from node 2 is simply 0,6; the path from 8 or 9 to 4 is 1,1.

As one might expect, it is easy to see these relationships if we think of node indices in their binary representation (see Figure 6.b). The lowest common ancestor 2 (binary 10) is designated by the longest common initial substring of 9 (binary 1001) and 11 (binary 1011). The relative index of 11, with respect to, 7 (binary 111), is the rest of its index with 1 prefixed to the front.

In terms of number of links traversed, relative addresses have no statistical advantage over the simpler method of always starting from the top. However, they have one important property that is essential for our purposes: relative addresses remain valid even when trees are embedded in other trees; absolute indices would have to be recalculated.

Figure 7 is a recoding of Figure 5 using relative addresses.

Keeping trees balanced
When two feature matrices are unified, the binary trees corresponding to them have to be combined to form a single tree. New attributes are added to some of the nodes; other nodes become "pointer nodes," i.e., their only content is the relative address of some other node where the real content is stored. As long as we keep adding nodes to one tree, it is a simple matter to keep the tree maximally balanced. At any given time, only the growing fringe of the tree can be incompletely filled. When two trees need to be combined, it would, of course, be possible to add all the cells from one tree in a balanced fashion to the other one but that would defeat the very purpose of using binary trees because it would mean having to copy almost all of the structure. The only alternative is to embed one of the trees in the other one. The resulting tree will not be a balanced one; some of the branches are much longer than others. Consequently, the average time needed to look up a value is bound to be worse than in a balanced tree.

For example, suppose that we want to unify a copy of the feature set in Figure 1b, represented as in Figure 2 but with relative addressing, with a copy of the feature set in Figure 8.

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Figure 8
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```
Figure 9
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Figure 7
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Figure 9
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Although the feature set in Figure 9.a is the same as the one represented by the right half of Figure 7, the structure in Figure 9.b is more complicated because it is derived by unifying copies of two separate trees, not by simply adding more features to a tree, as in Figure 7. In 9.b, a copy of 8.b has been embedded as node 6 of the host tree. The original indices of both trees remain unchanged. Because all the addresses are relative, no harm comes from the fact that indices in the embedded tree no longer correspond to the true location of the nodes. Absolute indices are not used as addresses because they change when a tree is embedded. The symbol -> in node 2 of the lower tree indicates that the original content of this node—gender 1,3—has been replaced by the address of the cell that it was unified with, namely cell 3 in the host tree.

In the case at hand, it matters very little which of the two trees becomes the host for the other. The resulting tree is about as much out of balance either way. However, when a sequence of unifications is performed, differences can be very significant. For example, if A, B, and C are unified with one another, it can make a great deal of difference, which of the two alternative shapes in Figure 10 is produced as the final result.

![Figure 10](image)

When a choice has to be made as to which of the two trees to embed in the other, it is important to minimize the length of the longest path in the resulting tree. To do this at all efficiently requires additional information to be stored with each node. According to one simple scheme, this is simply the length of the shortest path from the node down to a node with a free left or right pointer. Using this, it is a simple matter to find the shallowest place in a tree at which to embed another one. If the length of the longest path is also stored, it is also easy to determine which choice of host will give rise to the shallowest combined tree.

Another problem which needs careful attention concerns generation markers. If a pair of trees to be unified have independent histories, their generation markers will presumably be incommensurable and those of an embedded tree will therefore not be valid in the host. Various solutions are possible for this problem. The most straightforward is relate the histories of all trees at least to the extent of drawing generation markers from a global pool. In Lisp, for example, the simplest thing is to let them be CONS cells.

**Conclusion**

We will conclude by comparing our method of structure sharing with two others that we know of: R. Cohen's immutable arrays and the idea discussed in Fernando Pereira's paper at this meeting. The three alternatives involve different trade-offs along the space/time continuum. The choice between them will depend on the particular application they are intended for. No statistics on parsing are available yet but we hope to have some in the final version.

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