

# A Role for Chordless Cycles in the Representation and Retrieval of Information

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## ABSTRACT

This paper explains how very large network structures can be reduced, or consolidated, to an assemblage of chordless cycles (cyclic structures without cross connecting links), that is called a trace, for storage and later retrieval. After developing a basic mathematical framework, it illustrates the reduction process using a most general (with directed and undirected links) network.

A major theme of the paper is that this approach appears to model actual biological memory, as well as offering an attractive digital solution.

## Keywords

Closure, biological memory, reduction, consolidation, recall, trace, subgraph matching, directed graphs

## 1. INTRODUCTION

A central tenet of database theory is that it is the relationships between various entities that constitute real information. It is the foundation of the relational model [7]. But, in those situations where there are millions of entities and the relationships are relatively sparse, the familiar array style representation of data simply won't work. We turn to graph, or network, type representations. In other situations, such as social network analysis, the data is naturally graph structured. In any case, we are concerned with the retrieval of specific "chunks" of the stored information.

We will contend that retrieval of information from a storage medium is not a single, unified operation; that there are at least two distinct phases. First the desired information must be identified and located within the storage medium. We call this "information access" and address it in Section 3. Then it must be read out or, in the case of biological memory, reconstituted. This latter step we call "recall" and discuss it in Section 4.

However, this paper is less concerned with actual retrieval than discovering graph structures which facilitate this process. In Section 2.3 we introduce the concept of "chordless

cycles" which we believe play a prominent role in the representation of biological information, and which we believe can be exploited in computer applications as well. In Section 2.1 we sketch the mathematical foundations for this notion, which is based on concepts of "closure". In Section 2.2 we present computer code that will reduce any network to its constituent chordless cycles. Then in Section 2.3 we can describe the desirable properties of representing information by these cycles and indicate their role in biological memory.

Our goal in this paper is to use some relatively abstract graph-theoretic concepts to effectively model, and bring together, both biological and computer applications.

## 2. REPRESENTATION AND STORAGE OF INFORMATION

Our basic understanding of the world, whether visual, oral, or tactile, is neural in nature. The representation of data in regular arrays is, to a large extent, an artifact of computer architecture and the ease of algebraic manipulation. So we begin with the assumption that all "information" is a graph structure of some form.

To our knowledge, there has been no agreement as to what neural configurations correspond to any specific empirical sensations or mental concepts; but there have been many studies documenting that all perception and cognition correlate with neural activity, even detailing its occurrence with specific locations within the brain [11, 16, 15, 18, 21, 39, 40]. However, we are fairly confident that whatever neural networks do correspond to specific sensations, concepts or knowledge, they are very large — probably in the millions of elements, or possibly even billions, as discussed in [43]. We are talking about a very large data space!

The most general mathematical model of neural activity appears to be a directed graph. So the fundamental assumption of this paper is that, at its most basic level, information retrieval involves locating and obtaining rich directed graph structures based on simpler representations which serve as query keys. We are not retrieving array-type information to display on a spread sheet.

For this paper we assume that the network structure is the "information", independent of any edge or node labels.

### 2.1 Information as Relationships

Information and knowledge are essentially relational. Elements, or neurons, are related to other elements. To our knowledge, neurons are not "labeled". In contrast, most subgraph matching algorithms assume their nodes, or elements, are labeled [8, 24, 43]. These labels can be used to prune



which represents a considerable storage compaction.

The reduction process,  $\omega$ , has a number of desirable computational properties. Because the closure operator,  $\varphi$ , is local (it need only interrogate adjacent elements), it can be executed as a parallel process. (The code of Figure 2 that we have actually used is sequential<sup>1</sup>, but it is evident how to convert the outer loop.) This is particularly important for biological information storage and retrieval. The parallel application of a similar closure based process within the visual pathway is described in [36].

We have devoted considerable space to a description of  $\omega$  because, in addition to preparing information for storage, we believe it plays a prominent role in retrieval as described in the next sections.

In the literature devoted to human “memory”, there has been considerable research suggesting that there is a process that converts short-term memory into long-term memory. It is usually called “consolidation” [1, 4, 9, 25, 28]. It appears to be quite similar to the reduction process described above, whence the subsection heading is “consolidation”.

### 2.3 Chordless Cycles

A chordless cycle is most easily visualized as a string of pearls with no cross connections. More precisely, a *chordless cycle* is a sequence  $\langle y_1, y_2, \dots, y_n, y_1 \rangle$  of elements  $y_i, 1 \leq i \leq n$  of length  $n \geq 4$  where there exist no links (chords) of the form  $(y_i, y_k)$   $k \neq i \pm 1, i, k \neq 1, n$ .<sup>2</sup> It is better to just think of them as paths with no single edge cross connections. In Figure 3, the sequence, or path,  $\langle c, g, m, i, d, c \rangle$  is a chordless cycle of length 5. The sequence  $\langle n, q, p, u, z, A, B, x, t, n \rangle$  is a chordless cycle of length 9.

If the relation  $\eta$  between elements is symmetric, it can be proven [34, 35] that if  $\mathcal{N}$  is irreducible (*i.e.*, every node is a closed set) then every node  $y$  is either (1) isolated, (2) an element of a chordless cycle of length  $\geq 4$ , or (3) an element on a path between two such chordless cycles. Thus the trace (consolidation or reduction) of a symmetric relational network is a collection of interlinked chordless cycles. In Granovetter’s analysis of social networks [19], these are the “weak ties”.

When  $\eta$  is symmetric,  $\omega$  readily preserves path connectivity within  $\mathcal{T}$ . It also preserves the distance between elements as measured by shortest paths [35], together with the center of  $\mathcal{N}$  as determined by distance which will be found in  $\mathcal{T}$ , as will those centers as defined by “betweenness” [5, 13, 14].

Symmetry of the relation  $\eta$  is a powerful property. But, to better model real relationships we have relaxed this constraint throughout to allow directed (non-symmetric) connections. Under these conditions the preceding characterization of irreducible networks is no longer valid. For instance, the node  $f$  in Figure 3 is not an element of a chordless cycle.

In the case of non-symmetric networks we must ensure path connectivity through subsumed nodes. Let  $y$  subsume  $z$ , *i.e.*  $z.\eta \subseteq y.\eta$ . For all  $x$  such that  $z \in x.\eta$  we ensure that  $y \in x.\eta$ . This is always true when  $\eta$  is symmetric, but required the creation of the edge  $(f, c)$  in Figure 3. In this case, we can show [38] that if  $\mathcal{N}$  is irreducible then for all  $y$ , if there exists  $z \in y.\eta, z \neq y$ , there must exist a path

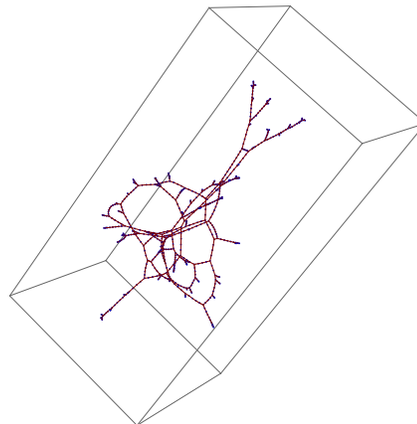
<sup>1</sup>It can be shown that worst case sequential behavior is  $O(n^2)$ , but actual performance is much better with a maximum of 6 iterations to reduce 1,000 node real networks.

<sup>2</sup>A graph with no chordless cycles is said to be *chordal*. There is an extensive literature on chordal graphs, *e.g.* [27]

through  $z$  terminating in a chordless cycle of length  $\geq 4$ . Even when the relation is non-symmetric, chordless cycles dominate the reduced representation.

Are chordless cycles really fundamental in the representation of biological information?

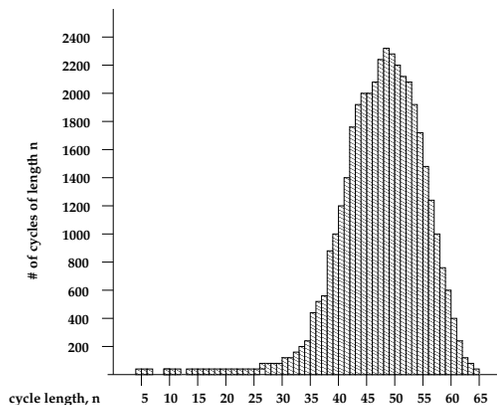
We can provide no definitive answer to that question. We can point out that protein polymer molecules composed of chordless cycles exist in every cell of our bodies [45]. One example is a 154 node phenylalaninic-glycine-repeat (nuclear pore protein),  $\mathcal{N}$ , which is shown in Figure 4. This is not at



**Figure 4: A 3D rendering of a 154 node protein polymer molecule.**

all like the dense network of Figure 1. Nevertheless, one can easily see the chordless loops, with various linear tendrils attached to them. When these are removed by  $\omega$ , there were 107 remaining elements involved in the chordless cycle structure.

If we count [10, 23] the numbers of chordless cycles of length  $n$  in the reduction of Figure 4, we obtain the distribution of Figure 5. The average cycle length is 44.9 in this



**Figure 5: The distribution of chordless cycles in a 107 element network.**

network; the modal length is 48; and the 4 longest chordless cycles have length 65. It is clear that the combinatorial possibilities based on cycle lengths alone would permit the representation of a considerable amount of information.

These are only indications that chordless cycle structures might be an important component of relational information

representation. The following sections will show how they can be involved with the retrieval process.

### 3. INFORMATION ACCESS

Information cannot be retrieved unless it is first identified and located within storage. Pointers and URLs identify storage locations. But, lacking these, one must search based on the “content” of the desired information. An early, and still common, mechanism is to attach key words or hash tags to the information, which are then used to create an easily searchable index [3, 17, 30, 31], in the case of hashed storage, to directly control the storage location [12] or to perform hash-joins in relational databases [42].

As access speeds have increased so dramatically, both of the preceding access techniques have often been superceded by simple linear search and comparison.

But, this still presumes that the “content identification” is based on specific terms, or tokens. If important information is based on relationships, as we have asserted in Section 1, then such token based search is rather limited.

Suppose we wish to retrieve within a relational network. We may assume that the search key,  $\mathcal{K}$ , is itself relational. It might look like Figure 6, but very much larger!

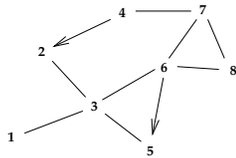


Figure 6: A structural search key,  $\mathcal{K}$ .

Reduction of  $\mathcal{K}$  by  $\omega$  yields its trace  $\mathcal{K}\omega$ , as shown in Figure 7. It is a simple cycle on the nodes  $\langle 4, 3, 6, 7, 4 \rangle$ .

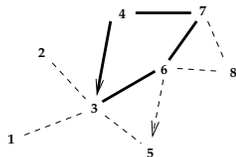


Figure 7: Its reduction,  $\mathcal{K}\omega$ .

Comparing Figure 7 with the trace of Figure 3, we see the obvious subgraph matching:  $3 \leftrightarrow m$ ,  $4 \leftrightarrow g$ ,  $6 \leftrightarrow p$  and  $7 \leftrightarrow k$ . In this small example there is only one possible matching chordless cycle. In the kinds of large networks we envision this is not so simple!

Imagine a 1,000 node search key which reduces to a key trace  $\mathcal{K}\omega$ , of say, 600 nodes. To find a matched subgraph in a reduced network of several million nodes is a combinatorially impossible task. All subgraph matching algorithms, known to the author, assume some form of node and/or edge labeling to initiate the search location and to control the expanding search [8, 24, 43]. Various forms of auxiliary indices provide direct access to graph elements matching those labels. However, while the retrieval of information from a graph structured data store implicitly assumes a labeled graph, we have focused in this paper on the relationships embodied in the network itself. There are no data labels.

Imagine “sliding” an unlabeled 600 irreducible subgraph around over an irreducible million node network to find a match, if one exists.

A method that we are beginning to explore involves the network providing information about its own irreducible structure. To do this we label each edge with the length of the “shortest” chordless cycle of which it is a member. This is not an artificial data label, but an element of the network representation itself. Thus, in the irreducible network of Figure 3, the edge  $(q, p)$  would be labeled with a 7 because it is a member of the chordless 7-cycle  $\langle q, p, k, g, c, d, i, m, q \rangle$ . This same edge is a member of longer cycles such as the 9-cycle,  $\langle q, p, u, z, A, B, x, t, n, q \rangle$ , but we only label with respect to the shortest.

The edge  $(m, q)$  would be labeled with a 4, as would the edge  $(m, p)$ . Each edge in the reduced search key,  $\mathcal{K}\omega$ , of Figure 7 must be labeled with a 4.

Each edge label of the search key must be exactly equal to the corresponding edge label in the larger network. So, based solely on the edge labeling, the only possible matches for this reduced key of Figure 7 in the reduced network of Figure 3 are the 4-cycles  $\langle g, m, p, k, g \rangle$ ,  $\langle n, q, m, i, n \rangle$  and  $\langle w, A, B, x, w \rangle$ .

Edge labeling with respect to shortest chordless cycle length eliminates fruitless search expansion, but it can’t indicate likely places to begin the comparison. The central retrieval problem is identifying likely nodes within the larger network to initiate the search comparison.

We believe that the solution lies with the imbedded “triangles”. Even though they are not chordless cycles of length  $\geq 4$  of which an irreducible network is comprised, they do exist, provided each edge is a member of a distinct chordless cycle of length  $\geq 4$ . In the reduced network of Figure 3 there is one embedded triangle, that is  $\langle q, p, m, q \rangle$ , with associated cycle labels  $\langle 7, 4, 4 \rangle$ . There is no embedded triangle in the reduced search key of Figure 7. This is unusual; but it is because the search key is “too small”. Our observation is that most reduced networks of 20 nodes, or more, have at least one embedded triangle. The network of Figure 4 has five, with associated cycle lengths of  $\langle 8, 6, 4 \rangle$ ,  $\langle 12, 5, 5 \rangle$ ,  $\langle 12, 7, 4 \rangle$ ,  $\langle 14, 10, 7 \rangle$  and  $\langle 16, 8, 6 \rangle$ . If the reduced search key contains any embedded triangle, it must match one of these.

In our JMC approach to information access, we 1) associate with each edge (functional relationship) in  $\eta$  the length of the shortest chordless cycle to which it belongs; and 2) create an external index of shortest length triples corresponding to embedded triangles. Assuming the search key is sufficiently well specified to include an embedded triangle, we first search this index of triples to identify one, or more, plausible search regions within the multi-million node reduced network constituting the data store.

This access mechanism is now being implemented, so its actual efficiency is still unknown. We believe it has considerable promise. More uncertain is whether the edge labeling and triangle index can be maintained in real time as the network changes dynamically [34]. Our hope is that if the network changes are *continuous* [32], then there exist effective update procedures.

The approach described in this section seems quite appropriate to biological recall as well. We see “Mary” in the market. This is a visual experience involving many relationships. A rapid, almost instantaneous, search through our

memory yields a much larger relational information structure, including her name, her age and the names of her 3 children. This is often called semantic memory in the literature [16].

#### 4. RETRIEVAL OF RELATIONAL INFORMATION

We suppose that a reduced trace,  $\mathcal{T} = \mathcal{N}.\omega$ , or a portion of it, has been identified as in the preceding section. During the reduction process,  $\omega$ , used to consolidate the network information we need not store only the resulting trace. We can also represent various steps in the reduction process. Our own code, for example, records  $y.\beta$  for each retained node  $y$ . Thus, if the trace of Figure 3 was accessed given the key of Figure 7 then we would know that  $c.\beta = \{a, b, c\}$ . Moreover, we know the order in which  $a$  and  $b$  were subsumed by  $c$ .

From this, a very close approximation of the original network of Figure 1 can be recreated. But, there will still be some loss of information. For instance, the connection between  $f$  and  $b$  would be lost. More complete information could be stored with each subsuming node, though at some increased storage expense. Alternatively, the complete information network,  $\mathcal{N}$ , could be stored, with the trace network,  $\mathcal{T}$ , separately stored as an easily searchable index.

It is our belief that with information structures of this size, retrieval of close approximations, *i.e.* the trace itself, will be sufficient for most applications.

##### 4.1 Biological Recall

Biological recall is likely to be somewhat different from a largely deterministic computer recall. There is some evidence that biological organisms only store an abbreviated trace and that the memory, as we experience it, is somehow “recreated” [20]. This helps explain why human memories can be distorted in specific detail, yet correct in their overall structure.

In [38], the author describes a semi-random expansion process,  $\varepsilon$ , that fills in subsumed nodes to create a richer network  $\mathcal{N}'$ . In many respects,  $\varepsilon$  is an inverse operator to  $\omega$ . Given a trace,  $\mathcal{T}$ ,  $\mathcal{T}.\omega^{-1}$  defines the collection of all networks  $\{\mathcal{N}^*\}$  such that  $\mathcal{N}^*.\omega = \mathcal{T}$ .  $\varepsilon$  constructs a single network,  $\mathcal{N}'$ , within this collection. Thus, given an initial network,  $\mathcal{N}$ ,  $\varepsilon$  will “retrieve”  $\mathcal{N}'$  which may, or may not, be (isomorphic to)  $\mathcal{N}$ . However, we are assured that  $\mathcal{N}'.\omega = \mathcal{T} = \mathcal{N}.\omega$ , or  $\mathcal{N}.\omega.\varepsilon.\omega = \mathcal{N}.\omega = \mathcal{T}$ .

Such a semi-random “retrieval” process may be completely inappropriate in computer applications, but it seems to model biological recall rather well. Our memories often are confused with respect to detail, even when they are generally correct. It also supports the notion of “re-consolidation” which asserts that long-term memories are repeatedly rewritten, unless deliberately distorted in our (semi)conscious mind [28, 29, 44].

#### 5. SUMMARY

The representation of complex networks by a trace comprised of chordless cycles has a firm, well defined basis. We know that there exists an effective procedure  $\omega$  to reduce a network  $\mathcal{N}$  to its trace  $\mathcal{T}$  by local and easily parallelizable code, such as might be found in the visual pathway. It is also known that such chordless cycle assemblages exist as protein polymers in all the cells of the body, including synapses.

Clearly, there are significant advantages, in terms of many fewer nodes and links, to representing network data in this fashion. Readily, real information stores, whether computer generated or biological, may have differentiated (*i.e.* labeled) nodes and links. Nevertheless, this first treatment of the problem as a purely abstract, graph theoretic issue has value. It provides a theoretical basis for more applied work.

We have suggested that subsets of the stored information can be identified and retrieved by similarly reducing the search key. Then, in Section 3, using graph-theoretic properties arising from the fact that irreducible networks consisting of chordless cycles, we have sketched an access method that might support retrieval based solely on relational structure. In biological organisms, this or an associative memory approach may be used.

The next to last paragraph of Section 4.1 mentions a reconstruction process  $\varepsilon$  which functions as a kind of inverse operator to  $\omega$ . This may be the most important observation of the paper. We are just beginning to explore it. If there is a way, given a trace  $\mathcal{T}$ , to reconstruct a network  $\mathcal{N}'$  which closely approximates the original network  $\mathcal{N}$ , this could have profound implications for both computed and biological applications.

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