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Variable binding by synaptic strength change

Christian R. Huyck*

School of Computing Science, Middlesex University, The Burroughs, London, NW4 4BT, UK

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Variable binding is a difficult problem for neural networks. Two new mechanisms for binding by synaptic change are presented, and in both, bindings are erased and can be reused. The first is based on the commonly used learning mechanism of permanent change of synaptic weight, and the second on synaptic change which decays. Both are biologically motivated models. Simulations of binding on a paired association task are shown with the first mechanism succeeding with a 97.5% *F*-Score, and the second performing perfectly. Further simulations show that binding by decaying synaptic change copes with cross talk, and can be used for compositional semantics. It can be inferred that binding by permanent change accounts for these, but it faces the stability plasticity dilemma. Two other existing binding mechanisms, synchrony and active links, are compatible with these new mechanisms. All four mechanisms are compared and integrated in a Cell Assembly theory.

Keywords: variable binding; cell assembly; short-term potentiation; long-term potentiation; synchrony; stability plasticity dilemma

1. Introduction

Symbol systems have been enormously successful and it has been proposed that, at least at some level, humans are symbol processors (Newell 1990). Whether humans are symbol processors or not, they can effectively use rules, and symbolic systems, such as ACT (Anderson and Lebiere 1998), have been very successful as models of human cognition. This success is probably due to the rule based or at least rule-like behaviour of humans in a wide range of tasks such as natural language processing.

Unfortunately, symbolic systems also have problems with brittleness (Smolensky 1987). The symbols are not grounded (Harnad 1990) and it is difficult or impossible to learn new symbols that are not just some combination of existing symbols (Frixione, Spinelli, and Gaglio 1989).

These and other problems provided motivation for the rise of connectionism, particularly in the 1980s. Connectionist systems are particularly good at learning, and thus may be able to learn

*Email: c.huyck@mdx.ac.uk

51 new symbols. If the systems learn from an environment, the newly learned symbols might even
 52 have semantic content grounded in that environment.

53 However, early connectionist systems were criticised for their inability to perform symbolic
 54 processes (Lindsey 1988). In particular, they were criticised for their lack of compositional syntax
 55 and semantics (Fodor and Pylyshyn 1988).

56 Variable binding offers an answer to these criticisms. A good variable binding solution allows for
 57 the implementation of rules; connectionist primitives can be combined, and variables instantiated
 58 as constants. If this can be done so that the result has compositional syntax and semantics, the
 59 criticism will have been answered.

60 For a binding mechanism to be functional, it must be able to support a range of binding
 61 behaviours (Section 2.1). Binding by synchrony (Malsburg 1981) is a well explored mechanism
 62 that is functional, but it can only support a limited number of bindings. Similarly, binding by
 63 active links (van der Velde and de Kamps 2006) has also been explored and is functional. Both
 64 mechanisms are restricted to active bindings, that is, the bindings must be continuously supported
 65 by neural firing, and when that firing ceases so do the bindings. This may limit the effectiveness
 66 of a neural system, particularly as it relates to composition (Section 6.2).

67 After some background for reader orientation, binding by synaptic change is introduced. This
 68 comes in two forms, binding by short-term potentiation (STP) and binding by compensatory long-
 69 term potentiation (LTP). Simulations that indicate these mechanisms are functional are described,
 70 in particular showing bindings can be formed and erased, that bindings can overlap, that a large
 71 number of bindings can be supported simultaneously, and that they can provide compositional
 72 syntax and semantics. It is shown that the four binding mechanisms, two existing and the two
 73 novel synaptic change mechanisms, are not mutually exclusive, and one system could use all four
 74 mechanisms. Ramifications for memory formation speed and duration are also explored along
 75 with other issues in the discussion and conclusion.

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2. Background

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Humans behave as if they have compositional syntax and semantics, so if systems based solely on
 neural models are to duplicate human behaviour, they too must exhibit compositional syntax and
 semantics behaviour. One way for neural systems to exhibit compositional syntax and semantics
 is by variable binding.

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A good cognitive model should have compositional syntax and semantics (Fodor and Pylyshyn
 1988). Standard symbolic cognitive architectures have this compositionality, but it is more difficult
 for connectionist models to exhibit it.

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Compositional semantics means that the semantics of a complex thing includes the semantics
 of that thing's constituents. So sentence 1

90

91

Pat loves Jody.

Sentence 1

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Includes the semantics of *Pat*, *love*, and *Jody*. Compositional syntax means that the syntactic
 structure of complex things affects the underlying semantics. For example, the semantics of
 sentence 1 is different from the semantics of sentence 2.

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Jody loves Pat.

Sentence 2

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So the semantics of a sentence must be more than the sum of its parts.

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Variable binding can be used to solve these problems in a neural system by binding the semantics
 of constituents in a syntax sensitive way. Sentence 1 could be represented by a case frame (Fillmore
 1968) for *love* where the actor slot is bound to *Pat*, and the object slot to *Jody*.

100

101 **2.1. The variable binding problem**

102 The variable binding problem is a key neural network problem that involves combining representations.
 103 It is also called the binding problem (Malsburg 1986), and the dynamic binding problem
 104 (Shastri and Aijanagadde 1993).
 105

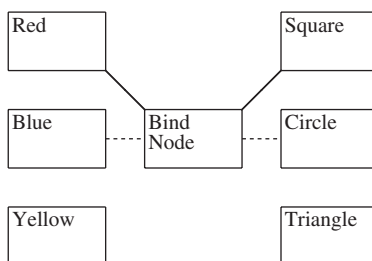
106 Perhaps the simplest variable binding problem is binding the features of an object. This is
 107 required when a new object is presented. If an object is composed of features, then when an object
 108 is presented, its features need to be bound together. One classic example is the *red-square* problem.
 109 If the system is presented with two objects, a *red-square* and a *blue-circle*, it can relatively easily
 110 activate the internal representation of all four of these features. The question is, how does the
 111 system know which pairs are bound.

112 A system can use a solution based on existing objects. For example, if there are two sets of
 113 100 features that can be bound, the problem can be solved by having 10,000 stored bindings, but
 114 this number will grow exponentially with the number of features, and the number of potential
 115 combinations. This solution is just a form of auto-associative memory that is open to the problem
 116 of exponential growth and thus combinatorial explosion. However, the features being bound into
 117 an object do not need to be a variant of an existing object, but can be a combination that is novel
 118 for the system.

119 Another example of this problem is binding parts into a whole, such as binding elements of a
 120 square lattice into rows or columns (Usher and Donnelly 1998). A third variant of this problem
 121 is the what-where problem. If a system can recognise multiple objects simultaneously and their
 122 locations, how does it know where each thing is and which things are in each location. This is an
 123 example of the above problem; in this case, location is one of the features, so one variant is the
 124 *left-square right-circle* problem.

125 Furthermore, unlike the standard associative memory task, binding features of an object has the
 126 associated difficulty of erasing the binding. After some time, *red* and *square* are no longer bound,
 127 and both may be bound to some other concept, for example *red-triangle*. This reuse problem is
 128 also a question of binding duration. As long as the binding persists, it can be used, but once it stops
 129 working, it can be reused for a new binding (Section 2.2). This paper is mainly concerned with
 130 bindings that are formed and then later erased so they can be reused. Figure 1 is an example of
 131 this. Here each box refers to a group of neurons with the outer six boxes referring to concepts (e.g.
 132 *Red* and *Circle*) and the centre box acting as a binding node. An initial binding of *Red* and *Square*,
 133 represented by the solid lines, is later replaced by the binding of *Blue* and *Circle*, represented by
 134 the dashed lines.

135 Another standard problematic example is filling in frames (Shastri and Aijanagadde 1993;
 136 Henderson 1994; Jackendoff 2002; van der Velde and de Kamps 2006). An example of this would
 137 be a verb frame (Fillmore 1968). For example the verb *move* might take an actor, object and
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 148
 149 Figure 1. Idealised binding with bind node: initial binding of red and square is later replaced by blue binding with
 150 circle.

151 location. In the sentence *Pat moved the ball to the door*. *Pat* would be the actor, *the ball* the
152 object, and *to the door* the location. When processing a sentence, the system would have to
153 fill in the details by binding these objects to these slots. Perhaps frames are a common task
154 for systems that use variable binding due to the compositional syntax and semantics prob-
155 lems mentioned by early critics of connectionist systems (Fodor and Pylyshyn 1988). Frames
156 are a flexible knowledge representation format (Schank and Abelson 1977); they are a rela-
157 tional structure where data is used to fill in structures with variables. The basic frames are
158 templates that need to be instantiated, and reused. Erasing the original's filler is one mech-
159 anism that can enable reuse. Moreover, if properly implemented, frames give compositional
160 semantics.

161 Rules are another important case where variable binding is needed. Firstly, rule based systems
162 are Turing complete (Hopcroft and Ullman 1979), so a neural implementation of rules would
163 be Turing complete. This is not particular surprising as others have shown other connectionist
164 systems to be Turing complete (Siegelmann and Sontag 1991). Secondly, rules are widely used
165 as a means of modelling human cognition (Laird, Newell, and Rosenbloom 1987; Anderson and
166 Lebiere 1998), so rules are important for cognitive modelling. An example rule would be *if X*
167 *gave Y to Z, then Z possesses Y*. Finally, sequences are important and can be implemented by
168 rules and by connectionist systems. For example, one system uses dynamic connections to learn
169 sequences (Feldman 1982). These learned sequences are then automatically forgotten by a process
170 of connection weight decay.

171 Unification is a more complex form of variable binding. This is done by symbolic systems
172 such as language processing systems (Shieber 1986) and logic programming. There are a range of
173 unification approaches, and complex structures such as directed acyclic graphs may be com-
174 bined (unified). It is a complex form of pattern matching. This can lead to a case where a
175 structure may be illegally combined with a subset of itself, known as the occurs check (Browne
176 and Sun 1999). Unification in neural systems may incorporate soft constraints making the sys-
177 tem more flexible (Hofstadter 1979; Kaplan, Weaver, and French 1990). For instance, there
178 may be a grammar rule that combines a noun phrase and a verb phrase and requires that they
179 agree in number; a soft constraint may allow the same rule to apply, in some circumstances,
180 when they do not agree in number, and this rule could be used to recognise ungrammatical
181 sentences.

182 A problem that is closely related to variable binding is Hetero-associative memory, which refers
183 to the association of an input with an output. This is roughly what Smolensky (Smolensky 1990)
184 refers to as variable binding, which differs from the term as used in this paper because hetero-
185 associative memories are permanent or extremely long-lasting. Perhaps this difference is the basis
186 of the term dynamic binding. To avoid confusion, in this paper, variable binding will only refer
187 to the case where a binding can be erased and reused.

188 Hetero-associative memory is a common and well understood form of memory (Willshaw,
189 Buneman, and Longuet-Higgins 1969). Here items are combined, and each is linked to that
190 combination. Presentation of one enables the system to retrieve the combined representation.
191 Of course restrictions can be placed on the inputs, and several features may be needed to
192 activate the full set of items (Furber, Bainbridge, Cumpstey, and Temple 2004). Standard neu-
193 ral models can account for this problem using standard Hebbian learning rules to implement
194 a form of LTP (Gerstner and van Hemmen 1992) for permanent synaptic change. How-
195 ever, this work is not easily extended to associative memories like semantic nets (Quillian
196 1967). The problem here is that one memory needs to be associated with another, yet the
197 two must remain separate. One excellent graph theoretic approach to this problem deals with
198 biological constraints on connectivity and activation (Valiant 2005). Both hetero-associative
199 memory and associative memory, though related, are different from variable binding (but see
200 Section 6.3).

2.2. *Properties of binding mechanisms*

Different binding mechanisms have different properties. This paper proposes that three properties are particularly important. These properties are:

- (1) Persistence of binding
- (2) Number of bindings supported
- (3) Speed to bind

Others have discussed the number of bindings property (e.g. Shastri 2006; van der Velde and de Kamps 2006), but persistence of binding and speed to bind are not typically discussed. This may be due to other work on binding being almost exclusively based on bindings being supported by neural firing (Section 6.2).

Persistence of binding has already been mentioned. Hetero-associative memories (Section 2.1), as typically modelled, persists forever. At the other extreme, binding via synchrony only persists as long as at least one of the bound items is firing, and binding by active links persists as long as the binding node is firing. This leaves a wide range of times that a binding might persist.

The number of bindings supported refers to how many entirely independent bindings, or distinct entities, can be supported simultaneously. One mechanism might be based on reusable binding nodes. Each node might be used to support one binding, and there are as many bindings as nodes. Figure 1 has one binding node that can support any of the nine possible bindings of one colour and one shape. A second, or third, node could be added to support another. The solution of forming a dedicated binding node for each possible binding is impractical because it would require an exponential number of nodes, so the nodes must be reusable. Therefore, in the case of verb frames (Filmore 1968), each slot of each verb might be a binding node. The slot fillers could be simple nouns, or they could consist of other verbs, in for example the case of sentential complements, to allow an arbitrary degree of complexity. Of course complex noun phrases would also need binding slots. With active links (van der Velde and de Kamps 2006) each binding node is represented by a circuit, and these can be combined to form verb frames. Binding via synchrony does not use nodes but has a limited number of bindings that a system can store (Sections 2.4.1 and 6.1).

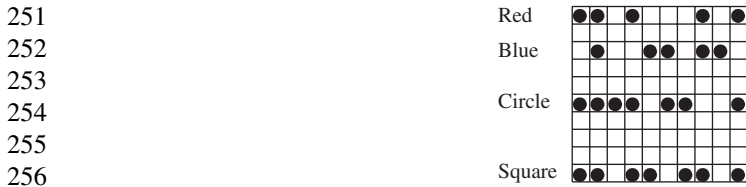
Finally, time to bind is an important consideration. How long must items be coactive before they can be bound? Binding via synchrony is very fast and can occur within tens of ms (Wennekers and Palm 2000). The binding via LTP mechanism proposed below (Sections 3.1 and 4.2) takes much longer.

2.3. *Cell assemblies and learning*

A cell assembly (CA) is the neural basis of a symbol (Hebb 1949). A CA is a subset of neurons that have high mutual synaptic strength enabling neurons in the CA to persistently fire after external stimulation ceases. In the simulations discussed in this paper, a small subset of all the neurons represents a symbol. If many of the neurons in the CA are firing, the symbol is active.

CAs give a sound answer to the neural representation of two types of memory, long-term memory and short-term (or working) memory. The firing of many neurons in a CA is the neural implementation of short-term memory; this high frequency and persistent firing makes the CA active.

The *red-square* problem can be restated in terms of CAs. There is a CA each for *red*, *blue*, *square*, and *circle*. When a *red-square* and a *blue-circle* are presented, all four base CAs are active. Figure 2 is an example of this problem. In this example, each cell represents a neuron with circles representing neurons that fire in a given period. The relevant rows are labelled with all neurons in a row representing the appropriate feature, and CAs are represented by orthogonal sets



257 Figure 2. Sample neural firing pattern for red-square and blue-circle.

258

259

260 of neurons. In this case some, but not all, of the relevant neurons are firing. Somehow the pairs
 261 must be bound, so that the system can ascertain, for example, the colour of the *square*, and this
 262 binding should only persist for a relatively small amount of time.

263 A CA is formed by a process of synaptic modification, and typically, this synaptic modi-
 264 fication is modelled as a form of LTP and long-term depression (LTD). CAs are long-term
 265 memories with Hebbian learning rules providing the link between long- and short-term mem-
 266 ories (Hebb 1949; O'Neill, Senior, Allen, Huxter, and Csicsvari 2008). When neurons co-fire,
 267 they become more likely to fire together because their mutual synapses are strengthened (Hebb
 268 1949), and eventually, this can lead to the formation of a CA. Hebbian learning is local; it
 269 occurs between two neurons that are connected and takes information based solely on these
 270 neurons. Typically the synaptic weight is increased when both the pre-synaptic and post-synaptic
 271 neurons fire. For all but the simplest forms of Hebbian learning, there is an associated form
 272 of forgetting that is, somewhat oddly, called anti-Hebbian learning. Here, if one neuron fires
 273 and the other does not, the synaptic weight is decreased (White, Levy, and Steward 1988),
 274 preventing the weight from growing without limit. There is significant biological evidence for
 275 Hebbian learning (Miyashita 1988; Brunel 1996; Messinger, Squire, Zola, and Albright 2005).
 276 Moreover, as this learning is based on pairs of neurons, biological experiments are relatively
 277 simple, so there is good reason to believe that some sort of Hebbian learning does occur in
 278 brains.

279 None the less, the precise mechanisms that are used by biological systems are not entirely
 280 clear. There are a range of Hebbian learning algorithms that follow the above definition, but
 281 differ from each other; none account for all biological data, and the biological data is far from
 282 complete.

283 The simplest rule merely increases the synaptic weight when both neurons co-fire. There is no
 284 anti-Hebbian rule, and the weight may be clipped at some value (Sompolinsky 1987) to prevent
 285 it growing without limit.

286 Timing is also important to learning. The Hebbian rule involves the firing of neurons at the
 287 same time. In a model that uses continuous time, the same time requires some degree of flex-
 288 ibility. Work on Spike Timing Dependent Plasticity (Gerstner and Kistler 2002) adds another
 289 dimension to the complexity of Hebbian rules. In these rules, precise timing dynamics are impor-
 290 tant with the order of neural firing affecting whether the change in synaptic weight is positive or
 291 negative.

292 The interaction between learning and firing leads to a complex dual dynamics (Hebb 1949).
 293 Once a CA is learned, it is hard to forget because any activation of it strengthens its intra-CA
 294 connections; this is a form of the stability plasticity dilemma (Carpenter and Grossberg 1988; Fusi,
 295 Drew, and Abbott 2005). Similarly, it is difficult to do anything with a CA until it has formed.

296 Hebbian learning rules are the most widely accepted model of the mechanism used by the
 297 brain to form CAs, the neural basis of concepts. Binding is not necessarily related to Hebbian
 298 learning, but if CAs, once formed, can be appropriately bound, then the resulting system can have
 299 compositional semantics and syntax. It then remains to ask what mechanisms can be used to bind
 300 CAs together?

2.4. Solutions to the problem

The mechanism that is most commonly used in neural simulations of variable binding is synchrony (Malsburg 1981). A lesser used mechanism is active links (van der Velde and de Kamps 2006), and both require neural firing to maintain the binding.

2.4.1. Binding via synchrony

Binding via synchrony requires neurons that are bound together to fire together. Therefore, if two neurons are bound, they might fire at times X , $X + 0.2$, $X + 0.5$, $X + 0.8$, and $X + 1$. For example, the neurons might fire at 0.1, 0.3, 0.6, 0.9, and 1.1; and then repeat the pattern at 1.5, 1.7, 2.0, 2.3, and 2.5. Of course there is some room for variation, and the binding usually applies to a much larger number of neurons than two.

A good example of this is SHRUTI, a non-neural connectionist mechanism (Shastri and Aijjanagadde 1993). In this model, different sets of concept nodes are bound together by firing at roughly the same time. Rules can be instantiated in the nodes, and these can continue to propagate the bindings to new items. SHRUTI has been used to develop, among other things, a syntactic parser (Henderson 1994). Here synchrony is used to bind slots and fillers. Unfortunately, the system only allows 10 bindings, so only relatively simple sentences can be processed.

There is significant evidence for synchronous firing in biological neural systems (Eckhorn et al. 1988; Abeles, Bergman, Margalit, and Vaddia 1993; Bevan and Wilson 1999). Some really convincing evidence that synchronous firing is used for biological binding is provided by a study that shows how binding is facilitated by a stimulus that is presented synchronously (Usher and Donnelly 1998).

There are several simulated neural models of binding via synchrony (e.g. Bienenstock and Malsburg 1987; Wennekers and Palm 2000). Networks of spiking neurons are used to segment a visual scene into different objects based on the firing timing of neurons associated with those objects (Knoblauch and Palm 2001); a scene with a triangle and a square is presented, and neurons associated with the square fire together and the triangle neurons fire together, but at different times from the square neurons. Spiking neurons are also used to parse simple text (Knoblauch, Markert, and Palm 2004) using binding via synchrony.

One major problem with binding via synchrony is the number of bindings that it supports (Section 2.2). The connectionist SHRUTI parser (Henderson 1994) is limited to 10 bindings, and Shastri and Aijjanagadde suggest that this limit is about 10 (Shastri and Aijjanagadde 1993). All bound items must fire in roughly the same pattern, but to handle variations within neural behaviour, this pattern must be somewhat flexible. Similarly, items that are bound differently must fire in a different pattern. For example, the neurons in *red* and *square* must fire in roughly the same pattern, while the neurons in *blue* must fire in a pattern that is different from *red*. As these firing patterns must occur in relatively brief time scales (~ 33 ms), and they must be relatively flexible, there are only a restricted number of bindings that can be maintained simultaneously. It is not entirely clear how many bindings biological neural systems allow, but as more bindings exist, there is an increased likelihood that closely related patterns will coalesce thus incorrectly combining sets of bound items.

2.4.2. Binding via active links

A more recent approach to the binding problem creates active neural circuits to support the binding (van der Velde and de Kamps 2006). Both primitives and binding nodes are represented by neural circuits, similar to CAs. The binding is selected by active primitives and is maintained by neural

351 firing in the binding node. Like binding by synchrony, the binding stops once firing stops in the
 352 binding node and stopping the binding circuit erases the binding. Binding can persist beyond
 353 firing in the primitives.

354 Effective simulations of natural language processing and vision have been demonstrated. This is
 355 a promising mechanism for variable binding. The active neural circuit solution is similar to an older
 356 connectionist solution called dynamic connections (Feldman 1982). Dynamic connections are
 357 used to store bindings that are activated by a pair of inputs, and then persist for a considerable
 358 period. The persistence automatically decays allowing the node to be reused later.

360 2.4.3. *Binding via LTP*

361 Another option is to bind by changing synaptic weights. An earlier version of the work presented
 362 in this paper used a fatiguing leaky integrate and fire (fLIF) neural model to implement rules to
 363 count from one number to another (Huyck and Belavkin 2006). A Hebbian learning rule is used
 364 to change synaptic weights permanently as a form of LTP.

365 Sougne provides an interesting blend between binding by changing synaptic weights and binding
 366 by synchrony (Sougne 2001). The changing synapses regulate synchrony by modifying delays
 367 on connections.

368 Unfortunately, a general binding solution based on LTP faces the stability plasticity dilemma
 369 (Carpenter and Grossberg 1988). The dilemma is how is it possible to add new knowledge without
 370 disrupting existing knowledge in a neural net (Lindsey 1988). With binding, base CAs would need
 371 to be stable, bindings would need to be plastic, and new CAs would still need to be formed. Thus
 372 any system that allowed a LTP based binding to be erased could have the problem of erasing the
 373 base CAs that are being bound.

376 2.4.4. *Other connectionist binding mechanisms*

377 One standard mechanism is to create a new binding element for each possible binding. As men-
 378 tioned earlier (Section 2.1), this has the problem of combinatorial explosion. This combinatorial
 379 explosion might be addressed by use of hierarchically allocated binding nodes (Hadley 2007)
 380 using prespecified roles. For natural language parsing, this requires millions of nodes, but the
 381 brain has billions of neurons, so this is plausible.

382 Another connectionist mechanisms for binding is to merely combine the bound representations,
 383 but this leads to systems that have problems with compositional syntax. An example is Tensor
 384 Product binding (Smolensky 1990) which forms a type of cross product of the variables that are
 385 being bound.

386 While some work has been done on binding via synaptic change in neural systems, most
 387 neural binding work has been done using synchronous firing. Some non-neural connectionist
 388 work is relevant to the problem. However, the possibility of binding via synaptic change is an
 389 under-explored area.

393 3. **Binding via LTP and STP**

394 There is strong evidence that distinct features that co-occur in a particular object cause synchronous
 395 neural firing (Eckhorn et al. 1988; Abeles et al. 1993; Usher and Donnelly 1998). While this appears
 396 to be solid evidence for binding via synchrony, it is not conclusive proof. Synchronous firing may
 397 simply be an emergent property of the neural representation of the new object as it is an emergent
 398 property of standard long-term CAs (Wennekers and Palm 2000). Assuming there is binding by
 399 synchrony, it still has a problem with capacity and a problem with duration of binding.
 400

401 It is not entirely clear how many bindings can be maintained by a network at any given time,
402 but each binding must have its own unique pattern of synchrony (Section 2.4.1). Natural language
403 processing may require many bindings as do other tasks such as object recognition. Since CAs
404 cross brain areas (Pulvermuller 1999), orthogonalising domains (e.g. vision and language) is not
405 a viable solution; that is, the brain can not be partitioned into areas where bindings are distinct so
406 that binding frequencies can simultaneously support multiple distinct bindings.

407 Also, the synchronous binding only persists as long as the CAs are active. Once they stop, the
408 binding is lost. While it is not entirely clear how long memories persist, there is a wide range of
409 times over which a binding might persist.

410 Even if binding via synchrony occurs in the brain, this does not mean that there are not other
411 types of binding. A different mechanism for binding, as is shown below, is change in synaptic
412 weights. There are at least two variants of known biological synaptic weight change, LTP and STP.
413

414 3.1. *Binding via LTP*

415
416 One possible solution to the binding problem is permanent synaptic change; biologically this is
417 LTP and LTD. Objects are bound using synaptic weight change, and these weight changes remain
418 until future learning erases them.

419 For LTP to be able to solve the variable binding problem, the binding must be able to be erased.
420 The mechanism then faces the stability plasticity dilemma (Carpenter and Grossberg 1988). If the
421 same mechanism is used to form the initial memories and to do the binding, something else must
422 prevent the initial memories from being erased when the bindings are erased.
423

424 3.2. *Binding via STP*

425
426 Most simulation work that involves learning relies on LTP. However there is another type of learn-
427 ing, STP, and there is extensive evidence that STP occurs in biological neural systems (Buonomano
428 1999; Hempel, Hartman, Wang, Turrigiano, and Nelson 2000). It is still a type of Hebbian learn-
429 ing, based on the firing behaviour of the neurons a synapse connects, so that co-firing increases
430 the synaptic weight. However, unlike LTP, the change is not permanent.

431 Some have proposed that STP provides support for LTP (Kaplan, Sontag, and Chown 1991).
432 That is, in the initial stage of CA formation, short-term connection strength adds activation to the
433 nascent CA that supports the co-firing that provides impetus for LTP. More recently, short-term
434 connection strength has been proposed as another basis of working memory (Fusi 2008; Mongillo,
435 Barak, and Tsodyks 2008). This contradicts the basic idea of active CAs as the basis of working
436 memory, but the two proposals may be compatible.
437

438 Another use for STP is for binding. In this case, the base memories are bound using STP. As
439 the STP is automatically erased, so is the associated binding. This paper is the first to describe
440 the use of STP in simulations of binding.

441 Note that the four binding mechanisms, synchrony, active links, compensatory LTP and STP,
442 are not mutually exclusive. Section 5.4 shows synchronous firing behaviour alongside binding
443 via LTP and STP, and describes how all four mechanisms could be combined in a single system.
444

445 4. *Simulating binding with LTP and STP*

446
447 To show that the STP and compensatory LTP binding mechanisms function, simulations of a
448 simple paired association task, similar to the *red-square* problem (Section 2.1), are described.
449 These and all the simulations described in this paper, use the same basic fLIF neural model.
450

4.1. *fLIF model*

The neural model that is used for the simulations described in this paper is an extension of the standard leaky integrate and fire (LIF) model which is in turn an extension of the integrate and fire (IF) model. A similar model (Chacron, Pakdaman, and Longtin 2003) has been shown to account for inter-spike intervals under various input conditions better than the standard LIF model. The IF model, commonly called the McCulloch Pitts neuron (McCulloch and Pitts 1943), has a long standing history and is quite simple. Roughly, neurons are connected by uni-directional synapses. A neuron integrates activity from the synapses connected to it, and if the activity surpasses a threshold, the neuron fires sending activity to the neurons it connects to. Connections may be excitatory or inhibitory; excitatory connections adding activity from the post-synaptic neuron and inhibitory connections subtract activity. LIF models are more biologically faithful than simple IF models (Churchland and Sejnowski 1992). In the IF model, if a neuron does not fire, it loses all its activity. In the LIF model, a neuron retains a portion of that activity making it easier to fire later. Typically, the neuron loses all its activity when it fires (Maass and Bishop 2001). All of these models are less complex and less accurate than Hodgkin Huxley models (Hodgkin and Huxley 1952) and other compartmental models (Dayan and Abbott 2005) which are extremely faithful to biology, breaking each neuron into several compartments and modelling interactions on a fine time grain (<1 ms).

The simulator runs in discrete steps with every neuron being modified in each step, and activity being collected in the next. The network of neurons can be broken into a series of subnets. Each neuron has two variables associated with it, and an array of synapses, and each subnet has four constants associated with all its neurons.

The two variables associated with each neuron i are fatigue F_i and activation A_i . As neurons fire, activation is passed to neuron i and is accumulated in A_i .

The first constant is the firing threshold, θ . A neuron i fires if

$$A_i - F_i \geq \theta \quad (1)$$

If the neuron fires, it loses all its activation. If sufficient activation is provided from neurons sending spikes to it, it may fire in the next time step.

If a neuron does not fire, some of its activation leaks away. This leak, or decay, is the second constant D where $D > 1$. Ignoring external input and assuming i did not fire at $t - 1$, activation of neuron i at time t is

$$A_i^t = \frac{A_i^{t-1}}{D}. \quad (2)$$

When neuron i fires, it sends activation (or inhibition) along its synapses to other neurons according to the strength of each synapse, so neuron j receives activation according to synaptic strength w_{ij} . The neuron is an integrator, so it accumulates activity from the synapses connected to it. Therefore, given P_j , the prior activation of neuron j , either 0 or Equation (2), the activation at time $t + 1$ is

$$A_j^{t+1} = P_j + \sum_{i \in V_i} w_{ij}, \quad (3)$$

where V_i is the set of all neurons that fired at time t .

These equations describe an LIF model (Maass and Bishop 2001). The fatigue variable is incremented by the third constant F_c in a cycle when the neuron fires, and is decremented by the fourth constant F_r in a cycle when the neuron does not fire. This makes it more difficult for neurons to fire the longer they are firing. Fatigue is a property of biological neurons (Kaplan et al. 1991).

501 The model has a loose link with time in biological neurons. The model does not incorporate
 502 conductance delays or refractory periods, and these behaviours all happen in under 10 ms, so
 503 each given cycle can be considered to be roughly 10 ms. Consequently, each neuron emits at
 504 most one spike per 10 ms. of simulated time, and the timing precision is at most 10 ms. This is a
 505 shortcoming of the model, but enables efficient simulation of hundreds of thousands of neurons
 506 on a standard PC.

507 The model also has some degree of topological faithfulness. The Hopfield Net (Hopfield 1982)
 508 has been a popular system for modelling brain function (Amit 1989), but it requires neurons to be
 509 well connected and connections to be bi-directional. Neither constraint is biologically accurate.
 510 However, one key point that these and other attractor nets (e.g. Rumelhart and McClelland 1982;
 511 Ackley, Hinton, and Sejnowski 1985) show is that attractor states are important; an attractor state
 512 is where roughly the same neurons and only those neurons fire in each cycle. This is a key point
 513 of CAs (Section 2.3).

514 The system uses neurons that are either inhibitory or excitatory but not both. While there is
 515 some debate over the biological behaviour, this follows the strict constraint of Dale's Law (Eccles
 516 1986). In the simulations described in this section, the ratio is 4 excitatory to 1 inhibitory neuron
 517 as is claimed in the mammalian cortex (Braitenberg 1989).

518 The connectivity of the network, and subnets is also important. Like the mammalian brain,
 519 excitatory neurons are likely to connect to neurons that are nearby. The network is broken into
 520 a series of rectangular subnets. As distance is relevant, the topology of each subnet is toroidal
 521 (the top is adjacent to the bottom, and sides are adjacent to each other, like folding a piece of
 522 paper into a donut) to avoid edge problems. In the simulations described in this section, excitatory
 523 neurons also have one long distance axon with several synapses. So a neuron connects to nearby
 524 neurons and to neurons in one other area of the subnet. These connections are assigned randomly,
 525 so each new subnet is extremely unlikely to have the same topology as another subnet with the
 526 same number of neurons. Equation (4) is used for connectivity.

$$527 \quad 528 \quad r < \frac{1}{(N * 8)} \longrightarrow \text{connect.} \quad (4)$$

530 It is initially called for each neuron with N (distance) of one for three adjacent neurons. It is
 531 subsequently called recursively on all four adjacent neurons with distance increasing one on each
 532 recursive call, and the recursion is stopped at distance 5. r is a random number between 0 and 1.
 533 The long-distance axon uses the same process though starts with distance 2. Inhibitory neurons
 534 are connected randomly within a subnet. This makes it easier for localised CAs to inhibit each
 535 other. There are approximately 60 synapses leaving a neuron to other neurons in the subnet, for
 536 both inhibitory and excitatory neurons.
 537

538 539 **4.2. Simulating binding by compensatory LTP** 540

541 The first set of simulations being reported in this paper involve binding via permanent changes of
 542 synaptic strength. This involves a compensatory Hebbian learning mechanism (Huyck 2004) that
 543 makes permanent changes to increase a synapse's strength, akin to LTP, and permanent changes
 544 that decrease the strength, akin to LTD. The simulation also makes use of spontaneous neural
 545 activation, a known biological phenomenon (Amit and Brunel 1997), to support erasing bindings.

546 The gross topology is shown in Figure 3. There are three subnets called the *letter* subnet, the
 547 *number* subnet, and the *binding* subnet. The *letter* and *number* subnets are trained to contain
 548 10 CAs each. Both nets consist of 1600 neurons and the *binding* subnet has 400. The binding
 549 subnet has spontaneous neural firing (see below) to enable erasing. As the base subnets do not
 550 have spontaneous firing, their CAs, once learned, are much more stable.

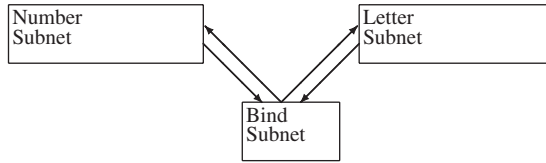


Figure 3. Topology of intra-subnet connections in the compensatory LTP binding simulation: each neuron in the base subnets connect to the bind subnet, and each neuron in the bind subnet connects to the base subnets.

In addition to the intra-subnet connection, each *bind* neuron has 15 connections to both the other subnets. The neurons of the base subnets, *letter* and *number*, have 16 connections to the *bind* subnet and all inter-subnetwork connections are randomly assigned. The initial weights are initialised to a number close to 0.

The compensatory learning mechanism is another type of Hebbian learning. It forces the total synaptic strength leaving a neuron towards the desired weight, W_B . Elsewhere (Huyck 2007), this learning mechanism has been used to learn hierarchical categories where categories share neurons. Compensatory learning is biologically plausible because the overall activation a neuron can emit is limited. Since a neuron is a biological cell, it has limited resources, and synaptic strength may well be one such resource.

The compensatory rule modifies the correlatory learning rules to include a goal total synaptic weight W_B . Equation (5) is the compensatory increase rule and Equation (6) is the compensatory decreasing rule; that is, Equation (5) is a Hebbian rule and Equation (6) an anti-Hebbian rule. W_B is a constant which represents the desired total synaptic strength of the pre-synaptic neuron, and W_i is the current total synaptic strength. R is the learning rate, which is 0.1. P is a constant and must be greater than 1. The larger it is, the less variance the total synaptic weight has from W_B . P , W_B , and R are constants associated with a particular subnet. When the two neurons co-fire there is an increase in synaptic weight corresponding to Equation (5). If the pre-synaptic neuron fires and the post-synaptic neuron does not fire, the weight is decreased according to Equation (6).

$$\Delta_+ w_{ij} = (1 - w_{ij}) * R * P^{(W_B - W_i)}, \quad (5)$$

$$\Delta_- w_{ij} = w_{ij} * -R * P^{(W_i - W_B)}. \quad (6)$$

Compensatory learning is important in the erasing process described below.

A summary of the value of the constants used in the first simulation can be found in Table 1. These values were determined by exploration of the parameter space via simulation. The parameter space, including topology, is practically infinite. This particular location is almost certainly not optimal, but does show solid results. An understanding of the dynamics of CA activation and formation is essential to select these parameters; this includes knowledge of various tradeoffs between parameters such as reducing firing threshold is similar to increasing synaptic strength. To a lesser extent, biological constraints also help in directing the search. For instance, excitatory synaptic weight is in the range of 0–1, and it is known that several neurons are needed to cause another

Table 1. Network constants.

Name	Symbol	Base net	Bind net
Threshold	θ	4	7
Decay	D	1.5	5
Fatigue	F_c	1.0	1.0
Fatigue recovery	F_r	2.0	2.0
Saturation base	W_B	21	28
Compensatory base	P	1.3	1.3

601 to fire (Abeles 1991) so the threshold θ is much greater than that. In one study of anaesthetised
 602 guinea pigs, simulated models accounted for spiking behaviour when decay was roughly $D = 1.25$
 603 (Lansky, Sanda, and He 2006).

604 During the entire run, there is spontaneous activation in the binding net. Spontaneous neural
 605 firing is a property of biological neurons (Abeles et al. 1993; Amit and Brunel 1997; Bevan and
 606 Wilson 1999), and it has been proposed as a mechanism for weakening and even erasing memories
 607 (Huyck and Bowles 2004).

608 In this simulation, some neurons may be spontaneously activated. This is modelled by the
 609 selection of a random number $0 \leq r < 1$ for each neuron in each cycle. If the $r < 0.03$ the neuron
 610 is spontaneously active. Therefore, roughly 3% of neurons in the bind subnet fire spontaneously
 611 each cycle.

612 The simulation first learns the base *number* and *letter* CAs, then one of each is randomly
 613 selected to be bound. This is a simple paired association task similar to the task performed in
 614 earlier connectionist simulations (Feldman 1982) and those done in psychological experiments
 615 (e.g. Sakai and Miyashita 1991). Once bound, the binding is tested, followed by a test for an
 616 unbound *letter* and *number*. The binding is then erased by spontaneous activation; and the tests
 617 are rerun. For measurement, this binding, testing, erasing, and retesting process is repeated 10
 618 times on each of 10 different networks.

619 The base CAs are learned by merely presenting components of them. As both the base nets
 620 consist of 1600 neurons, they can be divided into 10 orthogonal CAs of 160 neurons each. Fifty
 621 randomly selected neurons of a particular CA are selected and presented for 10 cycles. This is
 622 akin to clamping, but these neurons are given $\theta * (1 + \text{random})$ units of activation. After fatigue
 623 has accumulated they may not fire. After the 10 cycles of activation, the network is allowed to
 624 run for 40 more cycles. It is then reset with all activation and fatigue zeroed. Then a new CA is
 625 presented. Each set of 50 cycles of activation, run-on, and short-term variable resetting is called
 626 an epoch.

627 Each base CA is presented in a rotation so that all CAs are presented once every 1000 cycles.
 628 The complete training phase is 20,000 cycles so that each base CA is presented 20 times. Note
 629 that spontaneous activation in the *bind* net continues throughout this time.

630 Figure 4 shows the CA formation process. A network is created with synaptic weights near 0.
 631 It is then trained, and at the 45th cycle of each training epoch, the number of neurons in the
 632 presented CA is measured. This is averaged over the presentation of each of the 20 base CAs, and
 633 over 10 networks. The number of neurons outside the desired CA firing was also measured, but
 634 was always zero. This shows a rapid increase in persistence, neurons firing towards the end of
 635 each training epoch, followed by a gradual increase after the 5000th cycle. Note, the maximum
 636 number of neurons that could be firing is 160, but fewer are firing due to fatigue. By cycle 20,000,
 637 the base CAs are quite persistent.

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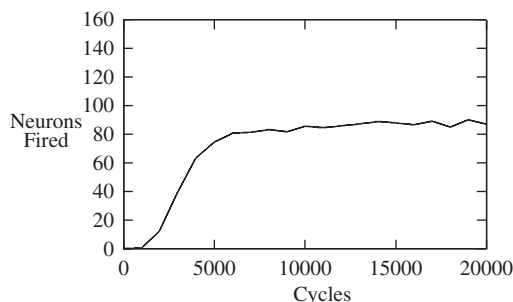


Figure 4. Neurons firing per cycle indicating CA formation.

651 After the training phase, the epoch duration is lengthened to 1000 cycles for the binding phase. A
 652 randomly selected *letter* CA and a randomly selected *number* CA are presented simultaneously.
 653 In a system that accepted visual input, both items would be presented simultaneously as in a
 654 paired association task. In this simulation, 50 neurons from both CAs are selected at random
 655 and presented for 10 cycles. As the CAs are already formed, these almost always persist for the
 656 duration of the binding epoch.

657 As ever, the *bind* subnet is spontaneously activated during this phase. Throughout this period
 658 the synaptic weights between the subnets gradually increase. When binding is successful, neurons
 659 in the *bind* subnet fire due to input from the active *number* and *letter* CA. This in turn causes the
 660 inter-subnet synapses to increase. In essence, a new CA is being formed and it includes neurons
 661 from all three subnetworks.

662 It is crucial that two CAs in the base subnets are simultaneously active. This is similar to the
 663 mechanism used for node activation by dynamic connections (Feldman 1982). Along with the
 664 spontaneously active *bind* neurons, these base neurons provide sufficient activation to fire some
 665 of the neurons in the *bind* subnet. Firing these base neurons causes the mutual synaptic strength
 666 between them and the base neurons to increase leading to further neural firing in the *bind* subnet.
 667 By the end of the binding epoch, a CA has been formed that includes the binding neurons, and
 668 this composite CA can be reactivated at any time over a significant period of time.

669 In the second epoch, the bound *number* is presented, and in the third, the bound *letter* is
 670 presented. When successful, this leads to activation of the binding CA and the opposite base CA.
 671 This further reinforces the inter-subnet synaptic strengths, improving the binding.

672 In the fourth epoch a randomly selected unbound *number* is presented, and an unbound *letter*
 673 is presented in the fifth. The correct result here is that no neurons in the opposite subnetwork fire.

674 The synaptic strength from the binding subnet that supports the binding is being reduced during
 675 the test unbound phase, but four further epochs of no base presentation are run to allow the binding
 676 to be sufficiently erased. The synapses from the binding subnet that support the binding move
 677 rapidly towards zero due to the application of compensatory learning rules (Equations (5) and
 678 (6)) caused by spontaneous firing.

679 Synapses from the *bind* subnet to the base subnets are erased during the period of no presenta-
 680 tion. During this period, neurons in the *bind* subnet fire, but no neurons in the base subnets fire.
 681 Consequently, the weights are reduced towards 0.

682 However, the synapses to the *bind* subnet from the base subnets are not changed during the
 683 testing of unbound items or during the period of no presentation. Instead, these synapses are
 684 reduced by the compensatory learning mechanism during the next two test epochs (epochs seven
 685 and eight).

686 The synaptic weights from neurons in the base subnets to the *bind* subnet do not change between
 687 the last binding test, and the first *bind* retest. Why then does the presentation of the here to fore
 688 bound item not cause the *bind* subnet to activate as it had done during the presentation in the
 689 second and third epochs?

690 Firstly, there are fewer neurons firing in the just bound item. This is due to the loss of intra-
 691 subnet synaptic strength during the binding. Secondly, there is little initial feedback from the *bind*
 692 node since its neurons no longer have much synaptic weight to the recently bound item. During
 693 this initial phase, the synaptic weights in the just bound item are changing. The weights to the
 694 *bind* node are being reduced while the weights within the just bound item are increasing. There
 695 is only a small part of the parameter space where this difficult task can be solved (Section 4.4).

696 Finally, there are four tests to assure that the binding has been erased. The formerly bound
 697 *number* and *letter* CAs are presented, followed by the formerly tested unbound *number* and *letter*.

698 For each network, this series of tests was run 10 times. It was run on a total of 10 networks. When
 699 the testing epoch length was 1000 cycles, 192/200 or, 96%, of the binding tests were successful,
 700 and 595/600, or 99.2%, tests of unbound CAs were successful. These measurements can be

701 combined using a standard $F-2 (2 * Bound * Unbound)/(Bound + Unbound)$. The F -score is
 702 97.5%.

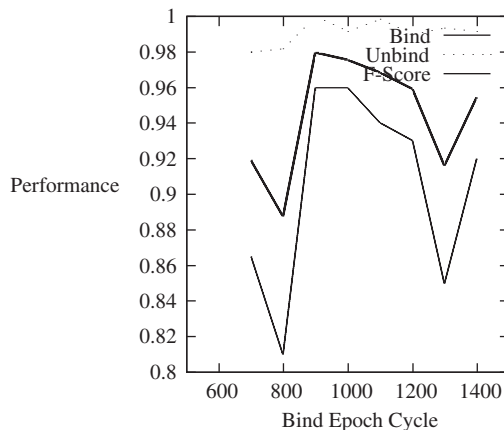
703 The length of the binding period is important. Substantial variations from the binding period
 704 of 1000 cycles causes decreased performance. Figure 5 shows this. Performance is best around
 705 1000 cycles, and trails off when it is shorter or longer.

706 It is important that the base CAs must be formed and solid before binding occurs. They need to
 707 be solid so that they can fully participate in the binding process. This solidity is supported by a low
 708 firing threshold ($\theta = 4$) and a low decay rate ($D = 1.5$); together these enable rapid recruiting of
 709 new neurons in a few presentations and high activity in the formed CA.

710 During base training, the binding area should not form a CA. That is, no single CA should be able
 711 to recruit many neurons from the binding area. Instead, two base CAs are needed to recruit neurons
 712 in the binding area. Consequently, little activity should be retained in the binding area ($D = 5$),
 713 and it should be difficult to fire a neuron in the binding area ($\theta = 7$). As the binding area needs
 714 to be quickly recruited when two CAs are active, the total synaptic strength is high ($W_B = 28$)
 715 so that the connections to the other areas and within can be quickly formed. This differentiation
 716 between systems (binding vs. bound) may be supported neurally by different neural types or
 717 neural pathways. It is a hallmark of neural processing, that different neurons behave differently.

718 None the less, a similar system could probably be developed with all subnets having similar
 719 or even identical parameters. The difference in threshold could be removed with a correspond-
 720 ing change in synaptic weights. The total synaptic strength would see a corresponding reduction,
 721 though it would still be different between subnets. This could probably be compensated by chang-
 722 ing the number of neurons. Changing the decay rate would be more difficult because the bind
 723 subnet can not retain much activity. A plausible solution would be to include an inhibitory system
 724 for the subnet that would inhibit neurons in the bind subnet on each cycle and thus eliminate the
 725 effect of a small amount of activity over many cycles. This has not been implemented, but it is
 726 likely that such a system could be developed.

727 This simulation fits into a rather small part of the parameter space. This is largely due to the
 728 rather precise way that the synapses from the base CAs to the binding subnet are erased. There
 729 is no spontaneous activation in the base subnets so the connections remain the same during the
 730 erase epochs. However, during the binding epoch, the synapses between neurons in the base CAs
 731 being bound have their strength taken by the synapses to the binding net. The loss of the feedback
 732 from the binding net after the erase epochs is enough to prevent the activation of the binding net
 733 when the bound base CA ignites.



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750 Figure 5. Effect of binding period on binding.

751 Since this is so precise, minor changes to parameters cause a rapid decrease in performance.
 752 Changing the number of synapses from each base neuron to the bind neurons from 16 to 15 gives
 753 Bound/Unbound/ F -Score results of 87%/95.5%/91.1%, and changing the number from 16 to
 754 17 gives $B/U/F$ results of 78.5%/94.8%/85.9%. Similarly, changing the base nets' desired total
 755 synaptic strength (W_B) from 21 to 20 gives $B/U/F$ results of 45%/99.2%/61.9%, and changing it
 756 from 21 to 22 gives results of 80.5%/89.8%/84.9%. Changing parameters individually is a form
 757 of gradient descent search; while gradient descent is not the best way to find an optimal place in
 758 the space, it can help to find local minima.

759 This is a particularly difficult binding simulation because there is no spontaneous activation in
 760 the base nets to facilitate erasing the binding. However, the lack of this spontaneous activation
 761 allows those CAs to persist indefinitely. Additionally, binding still works quite effectively.

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4.3. Simulating binding by STP

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Another option to implement variable binding by synaptic modification is to change the basic
 mechanism of synaptic change. LTP and LTD require the synaptic weight to remain unchanged
 until there is another application of one of the rules. Since synaptic change is caused by neural
 firing, the synaptic weights will remain unchanged until the appropriate neurons fire.

Another option is to have the weights automatically revert to zero over time. A rule that did this
 would be akin to STP. Note that the rule is still Hebbian in nature, changing the synaptic weight
 based solely on the firing behaviour of the two neurons that a synapse connects, but in this case,
 the weight also changes towards 0 when there is no firing.

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The binding via STP simulations reported below are identical to the binding via compensatory
 LTP simulations (Section 4.2) except the *bind* subnet is removed, neurons are replaced by neurons
 that learn via both LTP and STP, and the binding epochs are 50 cycles. The *bind* subnet was
 provided to localise erasing of bindings; with STP the bindings are automatically erased at the
 neural level.

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For STP, the simulation uses a new type of model neuron, termed a fast-bind neuron. The basic
 properties remain the same (Section 4.1), but some of the synapses leaving these neurons change
 their weights based on a different mechanism that accounts for STP.

The learning rule for fast-bind synapses that was used in these simulations is the simplest type
 of Hebbian learning. For each fast-bind synapse, if the pre-synaptic neuron fires in the same cycle
 as the post-synaptic neuron, the strength increases by the learning constant, which is 0.1. The
 weight is clipped at 1.

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The rule for reducing synaptic weight is equally simple. If the neuron does not fire in a cycle, all
 fast-bind synapses leaving it have their weight decreased by a constant k (in this case $k = 0.004$
 which was selected to assure the binding persisted for roughly 250 cycles after last use). Therefore,
 a maximally weighted synapse, will return to 0 after 250 cycles of inactivity. Similarly, a minimally
 weighted synapse will go to 1 after 10 cycles of pre and post-synaptic co-firing.

The topology of the *number* and *letter* subnets is the same as in the LTP simulations, with 80%
 excitatory and 20% inhibitory, and inhibitory neurons have no fast-bind synapses. Each neuron
 has two fast-bind synapses to neurons in each CA in the opposite subnet, and those neurons are
 randomly selected.

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The constants of the *letter* and *number* nets are the same as those in the LTP experiment; these
 are shown in Table 1. The training length is the same, 20,000 cycles, and the procedure is the
 same. The testing patterns are the same: binding epoch, two bind test epochs, two unbound test
 epochs, four epochs with no presentation, then two more tests of the formerly bound CAs, and
 two tests of the unbound CAs.

When the epoch lengths are 50 cycles, the system performs perfectly over 10 bindings on each of
 10 nets. That is, all 100 bindings were successful, and all associated 100 erasings were successful.

801 The Bound/Unbound/*F*-Score results are 100%/100%/100%. The bindings only need 10 cycles
802 to be fully established, and as they are given 50, they are firmly established. Similarly, only 250
803 cycles are needed for the bindings to be fully erased. As there are two unbound test epochs, and
804 four non-presentation epochs after the binding, there are 300 cycles of erasing, so erasing is also
805 perfect.

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808 **4.4. Performance of LTP vs. STP**

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810 It has been significantly simpler to use binding by STP than to use binding by compensatory LTP.
811 The portion of the parameter space that has been explored, where binding via compensatory LTP
812 functions acceptably, is quite small. This has required the use of relatively precise topologies,
813 precise training and use regimes, and spontaneous activation has been used only in the *Bind*
814 subnet to support erasing. On the other hand, binding by STP works in a much larger range of
815 conditions, and no exploration was done as the parameters for the LTP experiment were used.
816 The manipulation of learning and forgetting weights allows for a corresponding manipulation of
817 bind and unbind times (Section 5.2). Consequently, the next section discusses simulations using
818 binding by STP to account for crosstalk and compositionality.

819

820 Compensatory LTP should be able to account for these phenomena, but complex training
821 regimes may be needed, so at this juncture it seems unwise to describe further LTP simulations.
822 The basic problem with binding by compensatory LTP along with erasing by spontaneous acti-
823 vation is that it faces the stability plasticity dilemma. Some memories are stable, the items being
824 bound, and some are not, the bindings. It is difficult for the same mechanism to account for both.
825 Formation of bindings is slow and they persist for a long time, just like CAs, so it may be better
826 to view binding by compensatory LTP as a form of associative memory. However, this provides a
827 new way of addressing the stability plasticity dilemma that is more fully discussed in Section 6.3.

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829 The above simulations use fLIF neurons, but binding by compensatory LTP and STP should
830 both be applicable to other neural systems. Spiking models are particularly appropriate (e.g. Maass
831 and Bishop 2001). The rules may force breaking of the constraints of some attractor nets (e.g.
832 Hopfield Net connections would no longer be bidirectional), but this is not incompatible from a
833 simulation perspective (Amit 1989). Continuous value output neural models (e.g. Rumelhart and
834 McClelland 1982) should also be compatible with binding via STP. It is not entirely clear how
835 spontaneous activation would be implemented in these models, but compensatory learning should
836 still work. It is also not clear how these mechanisms would apply to connectionist systems that do
837 not have a close relationship to biological neurons like multi-layer perceptrons (Rumelhart and
838 McClelland 1986).

839

840 The binding by compensatory LTP and binding by STP models that are presented in this paper
841 are examples of classes of learning algorithms. The compensatory LTP mechanism was chosen
842 because a compensatory mechanism eases recruitment of new neurons to a CA, binding, and
843 supports erasing. The STP mechanism was chosen because of its simplicity. Ultimately, it is
844 hoped that the neurobiological basis of neural learning will be sufficiently illuminated to say
845 which algorithms are used for memory formation and variable binding in the biological system.
846 Until then, an exploration of different binding algorithms and their use in large systems to simulate
847 complex behaviour may be a good way to explore alternative neural binding mechanisms.

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847 **5. Further evaluation of binding by STP**

849 In the binding by compensatory LTP simulation (Section 4.2), a binding node was used. In the
850 STP simulation (Section 4.3) no explicit binding node was used, but implicitly, each CA was a

binding node so that 20 bindings could be supported. This required that each CA was connected to each CA in the opposite subnet, and this would require a geometric growth in synapses as the number of base CAs grew linearly. The use of binding nodes can make growth of synapses grow linearly as the base CAs grow linearly with each base CA connecting to the binding node. Of course, it is also possible to have many binding nodes to support multiple bindings at a given time. How do multiple bindings interact and how many can be supported?

5.1. Crosstalk

In this section, a system that stores multiple bindings is described. Storing these bindings could lead to problems of cross talk, but none are seen. The simulation combines both STP and LTP on a single neuron with specific synapses devoted to each. The gross topology is similar to that of Figure 3, but in this experiment there are multiple binding nodes.

There are four CAs in the *letter* subnet, four in *number* and four in *bind*. The *letter* and *number* CAs consist of 160 neurons each and the *bind* CAs have 100. All excitatory neurons have synapses leaving them that are modified by the compensatory LTP rule and synapses that are modified by the STP rule. The intra-subnet connections are the same as in the experiment described in Section 4.2 and all of these are modified by compensatory LTP.

Each neuron also has connections outside of the subnet and these are governed by the STP rule. Each neuron in the *letter* and *number* subnets has two connections to a randomly selected neuron in each CA in the *bind* subnet, and each of the *bind* neurons had three connections to each CA in the other subnets. This means that each neuron received roughly the same number of fast bind inputs as those in Section 4.3.

As in Sections 4.2 and 4.3, the base CAs were trained for 20 epochs of 50 cycles each. This formed stable CAs, and there was no spontaneous activation. The constants were the same as those for the base subnets in Table 1 ($\theta = 4$, $D = 1.5$, $F_c = 1.0$, $F_r = 2.0$, $W_B = 21$, and $P = 1.3$).

Bindings were set by a single epoch of 50 cycles of presentation of one *letter*, one *bind*, and one *number* CA. Initially this was *A0*, *B1*, *C2*, and *D3* each with a unique binding node.

Testing followed immediately with the numbers being presented in order. At the end of 50 cycles, the net was reset and the next number presented. On 100 nets, 400 of 400 correct *bind* and *letter* CAs fired in cycle 49 and no other neurons in those subnets fired. As expected, a random one to one binding (e.g. *A1*, *B2*, *C3*, *D0* each with a unique binding node) faired as well.

This test means that bindings are set and then allowed to be maintained without activation for 150 cycles. With automatic synaptic reduction set at 0.004 ($k = 0.004$) for each cycle when the pre-synaptic neuron does not fire, the synaptic weights return to zero after 250 cycles of inactivity. The simulation is run with a 50 cycle rest after the last binding, for a total of 200 cycles between the last cycle of each binding and each test. On 100 nets, none of the *letter* CAs have neurons firing, though 21 of the 400 *bind* nodes have some firing. The simulation was run with a 100 cycle rest after the bindings are set, and indeed the weights have returned to 0 and no firing was found in the *bind* and *letter* subnets.

The bindings are not formed simultaneously. So simultaneous presentation of *red-square* and *blue-circle* to the visual channel could not readily form two separate bindings. An attentional mechanism might be used with one object being attended to first and bound, followed by the second. Alternately, a different mechanism, e.g. active links, could be used to solve this problem (Section 5.4).

One common problem with binding is the presentation of two overlapping bindings, e.g. a *red-triangle*, and a *red-square*. This has been called the problem of two (Jackendoff 2002). This has been solved by a separate binding node for each pair (van der Velde and de Kamps 2006); elsewhere, this binding has been modelled with a computer simulation of CAs (deVries 2004) to account for psychological evidence.

901 The simulation was modified so that *A0*, *B1*, *C0*, and *D1* were presented, each with a unique
902 binding node. When the *letter* was presented the correct *number* CA was highly active with no
903 incorrect neurons firing for each of the 400 presentations on 100 tests. This shows that the binding
904 by STP addresses the problem of two.

905 Another test was done by presenting the *number*. When 0 was presented either *A*, *C*, or both
906 could ignite; and *B*, *D*, or both could ignite for 1. On 100 runs, when 2 or 3 were presented, no
907 letter neuron fired. Of the 200 positive tests, both of the bound *letter* CAs had over 100 neurons
908 fire 158 times, between 10 and 100 fired in one and the other was over 100 21 times, and in 21
909 tests fewer than 10 neurons fired in one while the other was near peak. This means that usually
910 both of the bound CAs ignited, but occasionally, due to competition, only one did.

911 As described in Section 4.1, each subnet is set up as a competitive subnetwork, with inhibitory
912 neurons that connect randomly within the subnet. In this case, each inhibitory neuron had 60
913 synapses. Fewer synapses lead to less competition, and more synapses to more competition. With
914 30 synapses on one hundred runs, both *letter* CAs fired on each of the 200 tests, though on two
915 tests less than 100 neurons fired in one CA. With 90 synapses on 100 runs on all 200 tests only
916 one was active and the other had less than ten neurons firing. Note that an inappropriate neuron
917 was never seen firing. Therefore, with ambiguous bindings, behaviour is dependent on the extent
918 of competition.

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920

921 5.2. Capacity

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923 In some sense, an exploration of the number of bindings that can be simultaneously supported
924 by STP is unnecessary. It is obvious that different orthogonal bindings can be independently
925 supported. For instance, filling in the topology for Figure 1 with values from the simulations
926 of Section 5.1 means that each orthogonal binding set can be represented by six base CAs of
927 160 neurons, and one binding CA of 100 neurons, or 1060 neurons. Therefore, the brain has a
928 capacity for billions of these orthogonal bindings, though it is extremely doubtful that the brain
929 has anything like that many orthogonal bindings.

930 Note that orthogonalising for synchrony is not the same as orthogonalising for STP binding
931 nodes. With STP, CAs can be involved with multiple bindings simultaneously without being
932 active, and there is no constraint on how many orthogonal bindings it can be in and be active.
933 With synchrony, if a CA is in multiple distinct bindings it has to fire in synchrony with all of them.

934 None the less it is interesting to see how many potentially overlapping bindings, as in the
935 experiments in Section 5.1, can be held simultaneously. Using the same method as in Section 5.1,
936 one binding can be set at a time, and parameters can be varied to expand from the four bindings
937 supported there. For simulations with extra CAs, an equal number of *letter*, *number*, and *bind* CAs
938 are added. Figure 6 shows a range of behaviour of simulations. The labels in the figure refer to
939 binding weight reduction k and bind durations with the 0.004/50 referring to the first simulations
940 of Section 5.1 that support four bindings. The other lines refer to different settings of k and bind
941 durations that allow more bindings to be supported.

942 First, the synaptic weight reduction parameter k can be reduced from 0.004. As it is reduced,
943 bindings will last longer and thus more can be set. In the 0.004/50 line of Figure 6 the maximum
944 theoretical duration of an inactive binding is 250 cycles as all of the synaptic weights will have
945 returned to 0. In the simulations, there are two synapses per neuron per CA, so several neurons
946 need to be active to cause firing and the bindings will not last for the full 250 cycles. More
947 synapses would cause this binding to persist longer, but could still not persist beyond 250 cycles.
948 The 0.001/50 line represents a synaptic weight reduction parameter of $k = 0.001$. This extends
949 the maximum duration to 1000 cycles, though again this may not be reached. Practically, this
950 performs entirely effectively to 650 cycles, and with 50 cycles to bind, this supports 13 bindings.

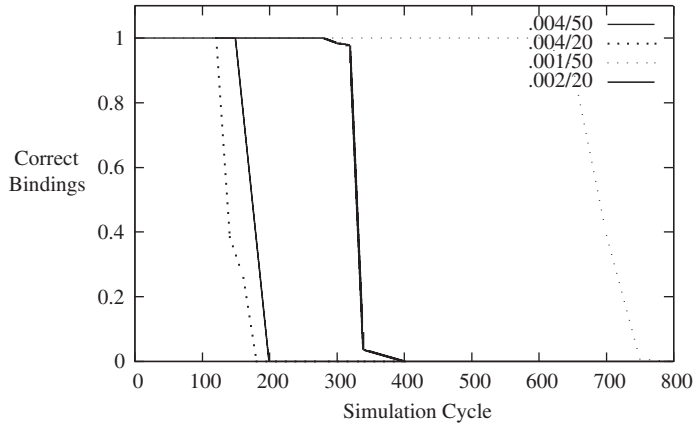


Figure 6. Duration of bindings via STP varying by reduction rate and time to bind.

Similarly, reducing bind time increases the bindings that can be maintained. With a learning weight of 0.1, 10 cycles are the minimum to fully bind. The 0.004/20 line in the figure represents a bind epoch of 20 cycles. This has the same maximum duration of 250 cycles, but more bindings can be supported over this time. There is theoretical limit of 10 bindings, but 7 are maintained perfectly.

Reduced bind time and smaller synaptic weight reduction combine multiplicatively. The 0.002/20 line in Figure 6 theoretically supports 20 bindings, four times two for the synaptic weight reduction parameter times 50/20 for the bind time. Practically it is supporting 15 perfectly effectively. A further set of simulations was run with 0.001/20 (not shown in figure). This shows 30 bindings being supported perfectly.

There is evidence that STP can last over 30 s (Varela, Sen, Gibson, Abbott, and Nelson 1997), which is 3000 cycles in the model. With 10 presentations to bind, 300 overlapping bindings can theoretically be supported simultaneously following the above binding setting mechanism.

5.3. Compositionality

The binding by STP mechanism supports frames, and thus supports compositional semantics. A simulation based on four subnetworks described in Figure 7 binds successfully over 98% of the time.

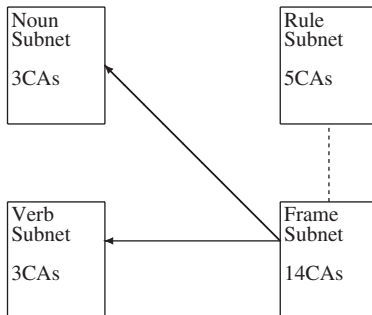


Figure 7. Gross topology of the simulation of binding with frames. The rule subnet inhibits the slots of the frames that are not active, and the slots are bound to the appropriate verbs and nouns via STP.

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1001 The four subnets are the *Verb*, *Noun*, *Rule*, and *Frame* subnets. The *Verb* and *Noun* subnet consist
 1002 of three CAs each of 160 neurons each representing a word; the *Rule* subnet of five CAs each of
 1003 800 neurons each representing a rule; and the *Frame* net consists of 14 CAs each of 100 neurons
 1004 which represent two frames each of seven slots. The constants were again the same as those for
 1005 the base subnets in Table 1 ($\theta = 4$, $D = 1.5$, $F_c = 1.0$, $F_r = 2.0$, $W_B = 21$, and $P = 1.3$).

1006 As in the earlier simulations, connectivity within each subnet was distance biased with 80–20
 1007 excitatory to inhibitory neurons. In the *Frame* subnet this was extended with synapses that learn
 1008 via the STP rule. Each frame consisted of seven slots, so the simulation has two frames. The base
 1009 slot was connected to the frame's other slots, and, as in the simulations from Section 5.1, each of
 1010 the neurons had two fast bind synapses to each of the appropriate CAs, along with the existing
 1011 synapses. The sentential complement slot had fast-bind synapses within the *Frame* subnet (see
 1012 below).

1013 Connectivity between the subnets was from the *Frame* subnet to the *Verb* and *Noun* subnets,
 1014 represented by the arrows in Figure 7; and from the *Rule* subnet to the *Frame* subnet, represented
 1015 by the dashed line. Each frame consisted of seven slots: *base*, *base verb*, *actor*, *object*, *location*,
 1016 *instrument*, and *sentential complement*. The *actor*, *object*, and *location* slots had connections
 1017 to each of the nouns, and the *base verb* slot had connections to each of the verbs. Each of the
 1018 excitatory neurons had two synapses to each of the appropriate CAs, and these synapses are
 1019 modified by the STP rule. The sentential complement slot was also connected to the base slot
 1020 of the other frame in the same fashion as the other slots were connected to nouns and verbs.
 1021 The instrument slot was not used in this simulation.

1022 The rules inhibited the frame slots that were incompatible. Each inhibitory neuron had 15
 1023 connections to each of those slots, and the rule CAs had 800 neurons to provide sufficient inhibition
 1024 to prevent those slots from igniting even when bound.

1025 As in the earlier simulations the net was trained by 20 presentations of 50 cycles for each of
 1026 the base CAs. As the rule CAs were 800 neurons, 400 neurons were presented during training
 1027 instead of 50 for CAs in the other nets.

1028 The relevant binding parameters are 20 cycles and $k = 0.004$. Binding was done by frames
 1029 that correlated to the sentences *Jody loves Pat.*, *Pat loves Jody.*, *Pat went to the store*, and *Jody*
 1030 *said Pat went to the store*. This was done by presenting the appropriate rule, slot and filler. For
 1031 example, the verb *love*, the first frame's base slot, the first frame's base verb slot, and the start VP
 1032 rule were presented for 20 cycles. For the first three sentences this was three presentations; for
 1033 *Jody loves Pat*:

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1035 (1a) the first frame's base and base verb, verb *love*, and the start VP rule;

1036 (1b) the first frame's base and actor slot, noun *Jody*, and add actor rule;

1037 (1c) the first frame's base and object slot, noun *Pat*, and add object rule;

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1040 The second sentence inverted the actor and object; the third used the verb *go* and replaced the
 1041 object rule and slot with location, and used the noun *store*. For the fourth sentence there was seven
 1042 presentations:

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1044 (4a) the first frame's base and base verb, verb *said*, and the start VP rule;

1045 (4b) the first frame's base and actor slot, noun *Jody*, and add actor rule;

1046 (4c) the first frame's base and *scomp*, and add *scomp* rule;

1047 (4d) the first frame's *scomp*, second frame's base, and add *scomp* rule;

1048 (4e) the second frame's base and base verb, verb *went* and the start VP rule;

1049 (4f) the second frame's base and actor slot, noun *Pat*, and add actor rule;

1050 (4g) the second frame's base and location slot, noun *store*, and add location rule.

1051 The complete test was done in three phases. The first phase bound the slots for *Jody loves Pat*.
 1052 into the first frame. There was then a period of erasing of 250 cycles. The second phase bound the
 1053 slots for *Pat loves Jody*. into the first frame and the slots for *Pat went to the store*. into the second
 1054 frame. There was then another period of erasing followed by the slots for *Jody said Pat went to*
 1055 *the store*. being bound into the first and second frame.

1056 Testing followed the phases before erasing. Testing was done by presenting the base frame slot
 1057 and the rule. The simulation was run on 10 different nets and each net did all three phases 10
 1058 times. The correct binding was considered to have occurred if more than 10 neurons in the correct
 1059 node were firing in the 19th cycle after presentation; no incorrect binding was considered to have
 1060 occurred if no other neuron in the appropriate net fired. For any given run, there were 14 possible
 1061 correct bindings, and 13 possible incorrect bindings (the sentential complement could not have
 1062 gotten the wrong base frame as both should be active). All of the correct bindings were formed and
 1063 in 1290 of the 1300 runs no incorrect bindings occurred. This gives a Bound/Unbound/*F*-Score
 1064 result of 100%/99.2%/99.6%. Note that the failures that occurred all occurred within one net
 1065 towards the end of the run, and were based on the base frame slot CAs recruiting each other via
 1066 LTP.

1067 A sentence is represented by a verb frame that has slots that are dynamically filled. *Pat loves Jody*.
 1068 includes the semantics of *Pat*, *love* and *Jody*, and is different from *Jody loves Pat*. The simulation
 1069 shows the difference between these two sentences and shows that frames can be implemented by
 1070 STP. Similarly, the simulation of the semantic representation of *Pat went to the store*. shows that
 1071 extra slots can be added seamlessly, and that multiple sentences can be stored simultaneously.

1072 The phenomena is recursive. The simulation of the sentence *Jody said Pat went to the store*.
 1073 shows that verb frames can be slot fillers. There is no theoretical limitation to the depth from
 1074 a psycholinguistic standpoint. From a simulation standpoint, reactivation of bindings might be
 1075 necessary during parsing to support the bindings, but Section 5.2 shows how 300 bindings might
 1076 be stored without recourse to separation.

1077 CAs are associative structures but frames are relational. This difference is bridged, above, by
 1078 fast-bind connections. Initially, the frame is represented by the base slot, and the remaining slots
 1079 are inactive. As slots are filled, the base slot and the particular slots are coactive; STP causes them
 1080 to be bound so the base slot will activate the bound slots, but not the unbound slots. In the test,
 1081 the unbound slots are explicitly activated via external activation.

1082 A more sophisticated framing mechanism has been used in a natural language parser (Huyck
 1083 2009). This parser uses frames for both *Noun* and *Verb* phrases because both can have others as
 1084 components. Rules no longer suppress frames, but instead activate particular slots in combination
 1085 with existing activation. The parser is stackless and follows other psycholinguistic models (Lewis
 1086 and Vasishth 2005).

1087 When multiple rules are applicable because of simultaneous activation, competition via inhi-
 1088 bition selects the rule to apply. For instance, when parsing a simple sentence like *I saw*. two items
 1089 are active the *NP I* and the *VP saw*. Two rules are also applicable the *AddActor* rule and the
 1090 *AddObject* rule. The *VP* is more active since it has been more recently activated, so the *AddActor*
 1091 rule wins and is applied. Once a slot is bound, it is marked as bound (neurally) and cannot be
 1092 rebound. In more complex sentences, several frames can be simultaneously active. In *Pat said*
 1093 *go to the store yesterday*. The frames *VP1 said actor-Pat scomp VP2*, *VP2 go loc-to-store*, *PPI*
 1094 *to-store*, and *NP3 yesterday* are all simultaneously active; the *NP1 Pat* frame is inactive since
 1095 it can no longer be modified. The rule that adds *yesterday* as the time of *VP1* will activate the
 1096 appropriate slot in that frame and the binding will be complete; the other two frames *VP2* and
 1097 *PPI* are already bound. The rule causes the binding, but the binding persists after the rule ceases
 1098 firing.

1099 It is fair to note that during parsing of a sentence, multiple constituents may be simultaneously
 1100 active. Only the appropriate items must be used to fill the appropriate slots. Binding by STP has

1101 now been used in two parsers: a stack based parser (Huyck and Fan 2007), and a memory based
 1102 parser (Huyck 2009). In the stack based parser, the appropriate items are selected by activating
 1103 them off of the stack, while other items on the stack are dormant.

1104 In the memory based parser all active items are active, but binding sites are activated via rules.
 1105 The item being bound has particular neurons that are associated with it being bound, and these
 1106 are only activated by the rule. The slot that is being filled has connections to the neurons for all
 1107 possible fillers with synapses that learn via STP. As only one slot and one filler are activated by
 1108 a particular rule, only they are bound. Therefore, if a particular PP is being set as the instrument
 1109 of a particular verb, the PP's neurons for being bound are active while no other filler has those
 1110 associated neurons active; the verb's instrument slot is active and only that slot is active. The
 1111 binding is completed, and the PP has a feature (represented by neurons) set that shows it has been
 1112 bound. It may still remain active, but will no longer be used as a filler.

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1115 5.4. Combining binding mechanisms

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1117 Variable binding is a complex problem and is needed for a wide range of behaviour. Consequently,
 1118 a system that could use a range of binding mechanisms would be more flexible than one that was
 1119 limited to one mechanism. Fortunately, all four mechanisms, binding by compensatory LTP,
 1120 binding by STP, binding by active links, and binding by synchrony are compatible.

1121 The above binding by STP and by compensatory LTP experiments exhibit synchronous firing
 1122 behaviour. For example, Figure 8 shows the firing behaviour of neurons in one run of the binding
 1123 by compensatory LTP simulation described in Section 4.2. This shows a section of one binding
 1124 epoch. The x axis shows the number of neurons firing in a subnetwork, and the y axis shows
 1125 the cycle. Initially, the *number* and *letter* CAs are firing in different cycles. As the strength of
 1126 the binding node grows, its neurons fire more frequently, and all three subnets begin to fire in
 1127 synchrony; the firing is so closely correlated that the dotted *number* line disappears in the figure
 1128 as it is covered by the *letter* line. The number of neurons firing in the base CAs oscillates, while
 1129 the number firing in the binding CA oscillates while growing. This shows a strongly correlated
 1130 firing pattern between the CAs.

1131 Figure 9 shows that items bound by STP fire synchronously. Here one letter is bound to one
 1132 number as in the simulations in Section 5.1. The *number* is presented which leads to the activation
 1133 of the *bind* CA and then of the *letter* CA. The firing patterns quickly synchronise.

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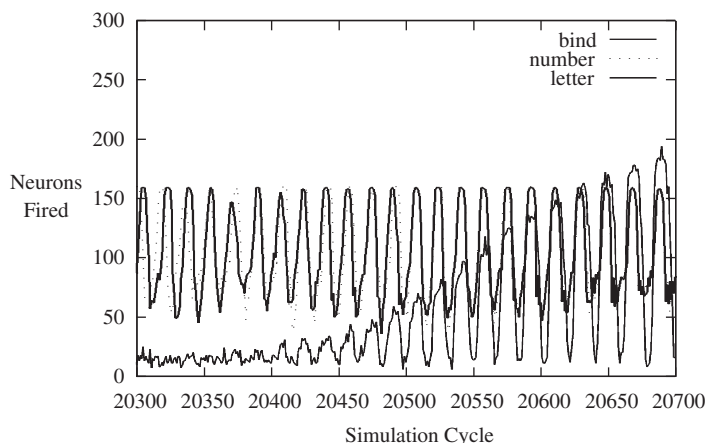


Figure 8. Firing of neurons showing synchronisation while binding.

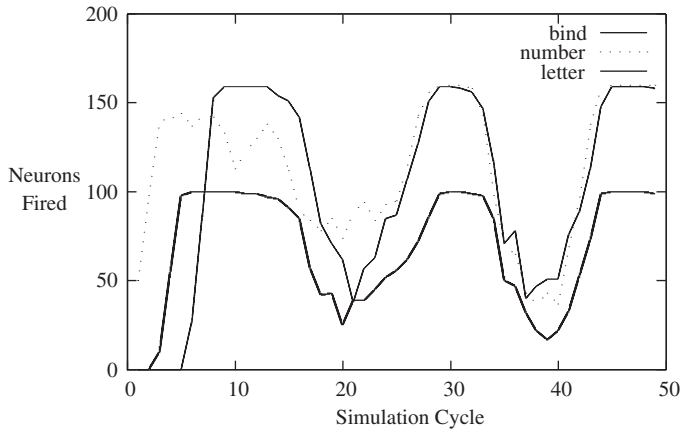


Figure 9. Firing of neurons showing synchronisation of items bound by STP.

The above fLIF neural model has been used to implement several systems including the Cell Assembly roBot version 1 (CABot1) agent (Huyck 2008). CABot1 is an agent in a video game that assists the user and is implemented entirely in fLIF neurons. It consists of vision subnets, planning subnets, an action subnet, a control subnet, and parsing subnets (Huyck and Fan 2007). The parsing subnets take the user's commands in natural language and parse them into semantic frames where the slots are filled via binding by STP. It is a stack-based system and the stack also binds by STP. The semantic result then leads to goals being set within the agent. Goals are context dependent, so a command like *Turn toward the pyramid.* needs to bind the goal to the location of the pyramid. This is done dynamically in a fashion similar to active links.

Similarly, a second parser has been developed that uses binding by STP for the stack and binding by compensatory LTP to fill the semantic frames. This indicates that these two variable binding mechanisms can be combined.

Referring back to associative memory (Section 2.1), both STP and synchrony have been proposed as mechanisms for supporting associative memory formation. There has been solid simulation work in the support of hetero-associative memory formation by synchrony (Shastri 2002; Gunay and Maida 2006). This avoids the stability plasticity dilemma by making bindings plastic and forgettable and hetero-associative memories permanent. It has also been proposed that short-term connection strength can be used to support long-term memory formation (Kaplan et al. 1991). Finally, there have been simulations that show active links also support long-term memory formation (van der Velde and de Kamps 2006).

6. Discussion

This paper has shown how two mechanisms for binding by synaptic change function. It has shown that one, STP, is capable of handling cross-talk and accounting for compositional semantics, and has inferred that the other mechanism, compensatory LTP, can too. Consequently, these new binding mechanisms can account for the problems described in Section 2.1. Elsewhere, it is shown how the earlier binding mechanisms, synchrony (Shastri and Aijanagadde 1993) and active links (van der Velde and de Kamps 2006), can solve these problems.

Since all four binding mechanisms are capable of binding, how do they differ? Below, each mechanism is evaluated on three important binding properties.

1201 **6.1. Binding properties**

1202
 1203 Both binding by compensatory LTP and binding by STP as described in this paper have values
 1204 associated with the properties of Section 2.2. Table 2 gives a qualitative overview of these values
 1205 and those associated with binding by synchrony and by active links. The first column refers to
 1206 the duration of the binding, the second to the number of different bindings that can be supported,
 1207 and the third to speed to bind.

1208 The persistence of binding for synchrony, and for active links is based solely on neural firing.
 1209 With synchrony, the binding persists while the bound items fire. With active links, the binding
 1210 persists while the binding node is firing.

1211 With binding by STP, the binding lasts as long as there is synaptic support for it. In the above
 1212 simulations using binding by STP, synaptic weights are reduced by 0.004 each cycle they are not
 1213 increased. Therefore, the weights are completely erased in 250 cycles, and may be effectively
 1214 erased in less; this equates to 2.5 s.

1215 The persistence of binding by compensatory LTP is more difficult to calculate. In Section 4.2,
 1216 6000 cycles (four erase epochs and two unbound test epochs) were used to erase the binding, or
 1217 60 s. Spontaneous activation in the *bind* subnet leads to the connections from the subnet being
 1218 erased. However, strong CAs can remain relatively stable under spontaneous activation due to the
 1219 relative stability of compensatory LTP. When there is spontaneous activation of a small number
 1220 of neurons, there are many more applications of anti-Hebbian learning than of Hebbian learning.
 1221 Therefore, the total synaptic weight, W_i , is significantly below the goal weight W_B . This means
 1222 that application of the anti-Hebbian rule changes the weights very little, and makes the original
 1223 weights surprisingly stable.

1224 The number of separate bindings differs between the four binding mechanisms. It is not clear
 1225 how many bindings can be supported by synchrony, but one simulation sets the limit at 10
 1226 (Henderson 1994). At the other extreme, binding by compensatory LTP supports a practically
 1227 unlimited set of bindings. In the first simulation, there is only one binding, but more could eas-
 1228 ily be modelled. Binding by LTP supports a number of bindings on the order of the number
 1229 of neurons. (As the number of synapses leaving a neuron is bounded by a constant, the bits per
 1230 synapse is constant, and these represent the memory of the system, memory is limited to $O(n)$ bits
 1231 where n is the number of neurons (Shannon 1948). Repeating the experiment from Section 4.2 on
 1232 orthogonal bindings would give $O(n)$ bindings.) There is no other practical limit for the number
 1233 of bindings except perhaps time to erase. The binding by STP mechanism that was used in the
 1234 above simulations also supports a practically unlimited number of binding nodes, though again
 1235 time is a factor. Section 5.2 shows that simultaneous support for 40 bindings is straight forward.
 1236 Of course, there can be multiple orthogonal sets of these bindings with, for instance, colour and
 1237 object, and verb and object, being bound. This would lead to a set of bindings on the order of the
 1238 number of neurons.

1239 For compensatory LTP, the values regarding time to bind are quite clear. The fLIF model
 1240 equates one cycle with 10 ms. Therefore, in Section 4.2, it takes roughly 1000 cycles to bind, so
 1241 roughly 10 s.

1242
 1243
 1244 Table 2. Binding property values by method.

	Persistence	Number	Speed
Synchrony	While firing	Few	Fast
Active links	While firing	Large	Fast
STP	Moderate	Large	Fast
LTP	Long	Large	Slow

1251 Compared with this, binding via STP is quite rapid. In the simulations in Sections 4.3 and 5
 1252 the learning rate is set to 0.1 and the weight is clipped at 1; so binding happens in 10 cycles, and
 1253 this equates to times about 100 ms. This contradicts the statement ‘it is unlikely that there exist
 1254 mechanisms that support widespread structural changes and growth of new links within’ hundreds
 1255 of ms (Shastri and Aijanagadde 1993). There is biological evidence of STP based on short bursts
 1256 of spikes that persist for seconds to minutes (Hempel et al. 2000).

1257 There is a vast range of evidence for synaptic changes of short duration (see Zucker and Regehr
 1258 (2002) for a review), and there are a wide range of behaviours, including different behaviours for
 1259 neurons in different portions of the brain (Castro-Alamancos and Connors 1997). Evidence shows
 1260 that short-term synaptic change can persist from under a second to over 30 (Varela et al. 1997).
 1261 It has been shown that as few as 10 spikes at 50 Hz can lead to STP of synapses (Tecuapetla,
 1262 Crillo-Reid, Bargas, and Galarraga 2007). In the simulations described in this paper, that would
 1263 be 10 sets of neural firings in alternating cycles. For all that is known to the contrary, it is possible
 1264 that the relevant form of rapid binding could be implemented by synaptic change. Bursts of 100 Hz
 1265 firings for as little as 300 ms. leads to STP that endures for tens of minutes (Schulz and Fitzgibbons
 1266 1997).

1267 It should also be noted that the time courses of the binding by STP and binding by compensatory
 1268 LTP are affected by the constants, topologies, and presentation mechanics. The above simulations
 1269 provide example time courses.

1270 Binding by synchrony can occur in tens of ms (Wennekers and Palm 2000). As active links
 1271 take only a few neural firings to form a binding, they too should occur on the order of tens of ms
 1272 (van der Velde and de Kamps 2006).

1273 A related property is the number of items per binding. The compensatory LTP mechanism
 1274 limits this, but binding by STP and binding by synchrony do not. Binding by active links allows
 1275 the developer to program this.

1276 **6.2. Maintaining binding by firing vs. by synapses**

1277 It has been stated that ‘the number of dynamic bindings expressed via some form of activity
 1278 (e.g. synchrony) will be comparable with the number of ignited (fired) CAs’. If bindings are
 1279 maintained by neural firing, this is the case, so it is the case for both binding by synchrony and
 1280 active links. However, if binding is done by synaptic modification, CAs do not need to be active
 1281 to remain bound; consequently, synaptic modification allows a much larger range of bindings to
 1282 be supported.

1283 If all of the bound items remain active, as in synchrony, or all of the binding nodes remain active,
 1284 as in active links, a large number of items are active. This can lead to problems of crosstalk. These
 1285 can be addressed programmatically, but it is clearly useful to be able to deactivate CAs and retain
 1286 bindings.

1287 Furthermore, maintaining a binding created by synaptic change, requires fewer neurons firing,
 1288 and neural firing is biologically expensive (Attwell and Laughlin 2001; Aiello and y Rita 2002).
 1289 Maintaining bindings by firing is thus biologically expensive. It costs a lot of energy.

1290 Therefore, binding by firing may be useful, but it comes at a cost. However, binding by synaptic
 1291 change has to pay much less.

1292 **6.3. Binding and memory**

1293 The three properties, speed to bind, number of bindings supported, and speed to unbind are also
 1294 issues of general memory formation. Recall that CAs give an explanation for short-term memory
 1295 (CA activation and persistence) and long-term memory (stable state CA formation based on LTP).
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1301 CA activation happens quickly (<20 ms), but does not last long (seconds). CAs form more slowly,
 1302 perhaps over days, but last much longer, perhaps years. CA activation and CA formation are akin
 1303 to speed to bind as all involve a memory formation. The cessation of a CA firing, and the loss of
 1304 a stable state are akin to a binding being erased as all involve the loss of memory.

1305 While there is some debate as to whether memories are lost or not, it is largely accepted that
 1306 as time passes, memories become less accessible (Klatzky 1980). Figure 10 shows the amount of
 1307 memory that can be accessed as time progresses by different neural memory processes. This figure
 1308 is meant to be a qualitative guide of the process indicating that as time passes fewer memories
 1309 from a particular time can be accessed. At the left of the figure, CA activation (CAA) does not last
 1310 long, but in a given period (say an hour) many memories can be used. On the right, CA formation
 1311 (CAF) shows that memories last a long time, but not many things (relative to the number of
 1312 CAs accessed) can be stored. Without binding, this leaves the middle ground empty; how can
 1313 something be forgotten after only a day? Binding fills in this middle area. Many items may be
 1314 bound by synchrony (BSyn) and by active links; there are fewer than the CAs that are active, and
 1315 they can persist longer as only one of the base CAs is needed to keep the binding. Binding by STP
 1316 (BSTP) probably occurs less frequently because it requires a modification of longer duration, but
 1317 it persists longer than binding by neural firing. Finally, binding by compensatory LTP (or any
 1318 LTP) has fewer items bound, but persists longer yet. Therefore, over a given hour, 1000 CAs
 1319 might activate, 100 sets of CAs might be bound via synchrony, 20 bound by STP, 10 bound by
 1320 LTP and two new CAs might be created. The active CAs would persist for 1 min, the synchronous
 1321 bindings for two, the bindings by STP for 5 min, the bindings by LTP for 2 h, one new CA might
 1322 last for a month and the other for 10 years.

1323 These memory mechanisms use and are influenced by the dual dynamics of CA activation and
 1324 CA formation. One good example of the complexity of these dual dynamics is the erasing of the
 1325 binding by compensatory LTP in Section 4.2. The weights from the bound letter to the *bind* subnet
 1326 are not changed during erasing. When the letter is presented after erasing, the synaptic weights
 1327 to the *bind* subnet are high, but they go down rapidly; there is a decline because the neurons in
 1328 the *letter* CA are firing and the *bind* neurons are not, and the decline is rapid because the total
 1329 synaptic strength is high. This rapid decline completes the erasing. The dynamics also have an
 1330 effect on the stability of existing CAs and formation of new CAs.

1331 Biological neural systems are always learning (Churchland and Sejnowski 1992), and there
 1332 is always spontaneous firing. Under these conditions, CAs must activate relatively frequently
 1333 to keep their mutual synaptic strength high. It does not seem reasonable that all CAs are acti-
 1334 vated relatively frequently. The relative stability of compensatory LTP bindings with spontaneous
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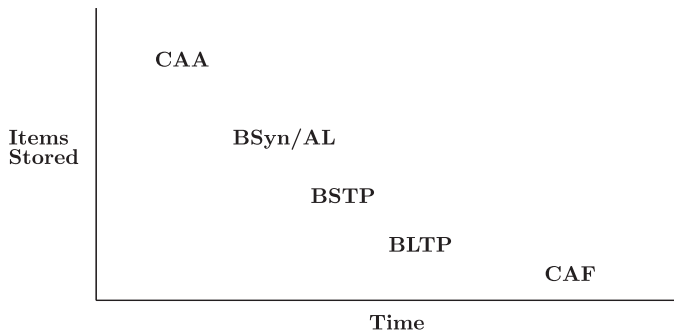


Figure 10. Memory hierarchy: different binding mechanisms provide a possible answer for the wide range of memory duration CAA, CA activation; BSyn, binding via synchrony; AL, binding via active links; BSTP, binding by STP; BLTP, binding by LTP; CAF, CA formation.

1351 activation provides some hope that this problem may be resolved, but it has not yet been since the
1352 system either is stable without spontaneous activation, or plastic with, but in neither case both.

1353 Binding by synchrony, active links, and STP have a lesser effect on CA stability and plasticity,
1354 but they still have an effect. They have less of an effect because they are not based on long-term
1355 synaptic change. They still have an effect because they cause the simultaneous firing of neurons
1356 in CAs, and this will lead to increased permanent synaptic weight between the bound CAs. This
1357 might lead to the CAs recruiting each other, so that they no longer can be independently active.

1358 Binding by compensatory LTP can now be looked at as an associative memory mechanism.
1359 CAs that are frequently bound may become more related but, perhaps due to topology, may not
1360 recruit each other. Other options for resolving stability problems include modified spontaneous
1361 activation mechanisms, subassemblies, and learning rules involving fatigue. In the simulations
1362 described in this paper, spontaneous activation is purely random; this might be modified to make
1363 neurons fire when they have not fired for a long time, and these neurons might co-fire based on
1364 their last activity. Subassemblies are merely sets of neurons that do not persist, but can be activated
1365 by spontaneous activation leading to synaptic support. Finally, if synaptic weights only changed
1366 significantly when neurons were fatigued, spontaneous activation would have little effect on them.
1367 These mechanisms are, of course, speculative.

1368 Binding by compensatory LTP, and to a lesser extent the other binding mechanisms, provides a
1369 window into the stability plasticity dilemma of associative memory. It is relatively easy to model
1370 the indefinite storage of memories as once stored all memories are stable. When the memory
1371 store is large, this may cause no obvious problems. However, access to all memories can not be
1372 retained, and access to psychological memories is lost on a range of scales. Perhaps binding by
1373 compensatory LTP will provide an answer to how memories can be forgotten after days or years.

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1377 **7. Conclusion**

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1379 Binding is an important problem because a solution to it allows a system to have compositional
1380 syntax and semantics. This composition is necessary for a system to model the full range of human
1381 behaviour. If the particular problems of binding features in an object, frames, and rules can be
1382 solved, then a system can be built that is compositional.

1383 This paper has introduced a new variable binding mechanism, binding by STP and made use
1384 of the relatively novel variable binding by compensatory LTP. Simulations have shown that these
1385 mechanisms, like synchrony and active links, can bind features in an object, and implement rules
1386 and frames. Simulations have shown that binding by STP also solves the problem of two and that
1387 binding by LTP should be able to.

1388 Binding by STP is fast to bind, persists beyond the activity of the bound CAs, is relatively
1389 easy to engineer, and works consistently. Binding by compensatory LTP works, but faces the
1390 stability plasticity dilemma. It is slower to bind and the bindings persist longer. Neither of these
1391 mechanisms faces a combinatorial explosion to bind items, and both can support a very large
1392 number of bindings.

1393 Binding via compensatory LTP and by STP can be used together and with the earlier defined
1394 binding mechanisms, binding via synchrony and binding by active links, to complement each
1395 other. They each have different behaviours on time to bind, time to erase, and capacity. Along
1396 with CA activation and CA formation, these binding mechanisms give a wide range of memory
1397 formation and retention behaviour.

1398 Together, these mechanisms allow for a sophisticated use of compositional syntax and semantics
1399 in a simulated neural system. This will support the development of complex symbol processing
1400 agents from simulated neurons bridging the gap between subsymbolic and symbolic systems.

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