#### Marr's Theory of the Hippocampus: Part I

#### Computational Models of Neural Systems Lecture 3.3

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#### David Marr: 1945-1980



David Marr 1970 – Cambridge, England

#### Computational Models of Neural Systems

#### Marr and Computational Neuroscience

- In 1969-1970, Marr wrote three major papers on theories of the cortex:
  - A Theory of Cerebellar Cortex
  - A Theory for Cerebral Neocortex
  - Simple Memory: A Theory for Archicortex
- A fourth paper, on the input/output relations between cortex and hippocampus, was promised but never completed.
- Subsequently he went on to work in computational vision.
- His vision work includes a theory of lightness computation in retina, and the Marr-Poggio stereo algorithm.

#### Introduction to Marr's Archicortex Theory

- The hippocampus is in the "relatively simple and primitive" part of the cerebrum: the <u>archicortex</u>.
  - The *piriform* (olfactory) cortex is also part of archicortex.
- Why is archicortex considered simpler than neocortex?
  - Evolutionarily, it's an earlier part of the brain.
  - Fewer cell layers (3 vs. 6)
  - Other reasons? [connectivity?]
- Marr claims that neocortex can learn to classify inputs (category formation), whereas archicortex can only do associative recall.
  - Was this conclusion justified by the anatomy?

#### What Does Marr's Hippocampus Do?

- Stores patterns immediately and efficiently, without further analysis.
- Later the neocortex can pick out the important features and memorize those.
- It may take a while for cortex to decide which features are important.
  - Transfer is not immediate.
- Hippocampus is thus a kind of medium-term memory used to train the neocortex.

#### An Animal's Limited History

- If 10 fibers out of 1000 can be active at once, that gives C(1000,10) possible combinations =  $2.6 \times 10^{23}$ .
- Assume a new pattern every 1 ms.
  - Enough combinations to go for 10<sup>12</sup> years.
- So: assume patterns will not repeat during the lifetime of the animal.
- Very few of the many possible events (patterns) will actually be encountered.
- So events will be well-separated in pattern space, not close together.

#### **Numerical Contraints**

Marr defined a set of numerical constraints to determine the shape of simple memory theory:

- 1. Capacity requirements
- 2. Number of inputs
- 3. Number of outputs
- 4. Number of synapse states = 2 (binary synapses)
- 5. Number of synapses made on a cell
- 6. Pattern of connectivity
- 7. Level of activity (sparseness)
- 8. Size of retrieval cue

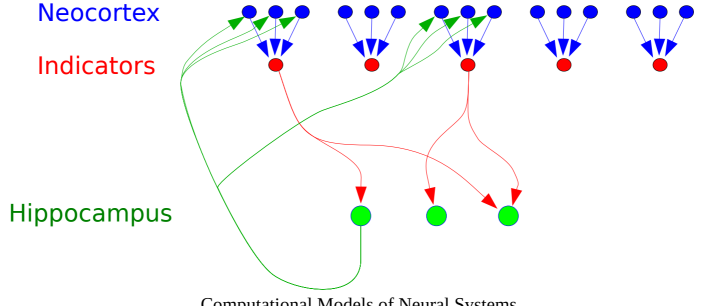
#### N1. Capacity Requirements

- A simple memory only needs to store one day's worth of experiences.
- They will be transferred to neocortex at night, during sleep.
- There are 86,400 seconds in a day.
- A reasonable upper bound on memories stored is:

#### 100,000 events per day

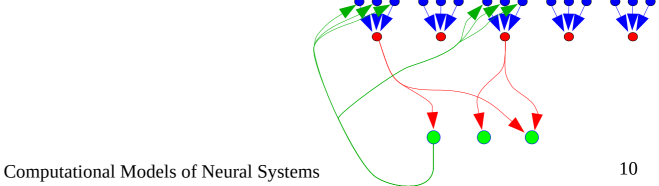
#### N2. Number of Inputs

- Too many cortical pyramids (10<sup>8</sup>): can't all have direct contact with the hippocampus.
- Solution: introduce indicator cells as markers of activity in each local cortical region, about 0.03 mm<sup>2</sup>.
- Indicator cells funnel activity into the hippocampal system.



#### **Indicator Cells**

- Indicator cells funnel information into hippocampus.
- Don't we lose information?
  - Yes, but the loss is recoverable if the input patterns aren't too similar (low overlap).
- The return connections from hippocampus to cortex must be direct to all the cortical pyramids, not to the indicator cells.
- But that's okay because there are far fewer hippocampal axons than cortical axons (so there's room for all the wiring), and each axon can make many synapses.



#### How Many Input Fibers?

- Roughly 30 indicator cells per mm<sup>2</sup> of cortex.
- Roughly 1300 cm<sup>2</sup> in one hemisphere of human cortex, of which about 400 cm<sup>2</sup> needs direct access to simple memory. Thus,

# About 10<sup>6</sup> afferent fibers enter simple memory.

• This seems a reasonable number.

#### N3. Number of Outputs

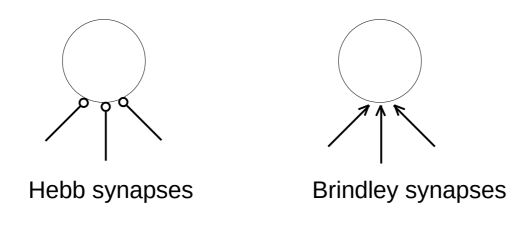
- Assume neocortical pyramidal cells have fewer than 10<sup>5</sup> afferent synapses.
- Assume only about 10<sup>4</sup> synaptic sites available on the pyramidal cell for receiving output from simple memory.
- Hence, if every hippocampal cell must contact every cortical cell, there can be at most 10<sup>4</sup> hippocampal cells in the memory. Too few!
  - If 100,000 memories stored, each memory could only have 10 cells active (based on the constraint that each cell participates in at most 100 memories.) Too few cells for accurate recall.
- Later this constraint was changed to permit 10<sup>5</sup> cells in the simple memory.

## N4. Binary Synapses

- Marr assumed a synapse is either on or off (1 or 0).
- Real-valued synapses aren't required for his associative memory model to work.
  - But they could increase the memory capacity.
- Assuming binary synapses simplifies the capacity analysis to follow.

## Types of Synapses

- Hebb synapses are binary: on or off.
- Brindley synapses have a fixed component in addition to the modifiable component.



- Synapses are switched to the on state by simultaneous activity in the pre- and post-synaptic cells.
- This is known as the Hebb learning rule.

#### N5. Number of Synapses

- The number of synapses onto a cell is assumed to be high, but bounded.
- Anatomy suggests no more than 60,000.
- In most calculations he uses a value of 10<sup>5</sup>.

#### N6. Pattern of Connectivity

- Some layers are subdivided into blocks, mirroring the structure of projections in cortex, and from cortex to hippocampus.
- Projections between such layers are only between corresponding blocks.
- Within blocks, the projection is random.

#### N7. Level of Activity

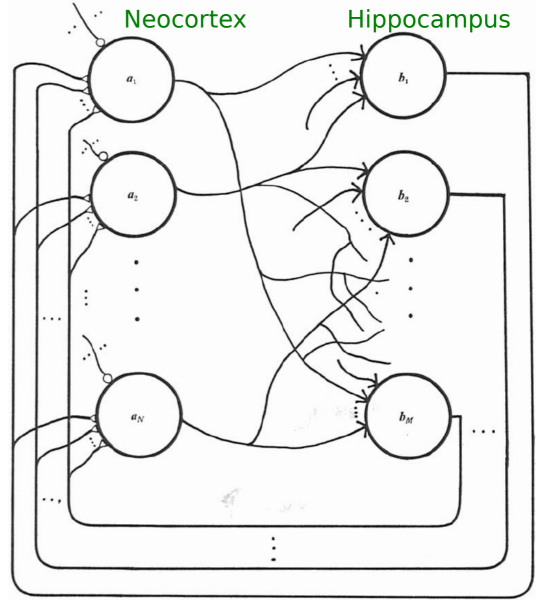
- Activity level (percentage of active units) should be low so that patterns will be sparse and many events can be stored.
- Inhibition is used to keep the number of active cells constant.
- Activity level must not be too low, because inhibition depends on an accurate sampling of the activity level.
- Assume at least 1 cell in 1000 is active.
- That is,  $\alpha > 0.001$ .

#### N8. Size of Retrieval Cue

- Fraction of a previously stored event required to successfully retrieve the full event.
- Marr sets this to 1/10.
- This constitutes the minimum acceptable cue size.
- If the minimum cue size is increased, more memories could be stored with the same level of accuracy.

#### Marr's Two-Layer Model

- Event E is on cells  $a_1 \dots a_N$ (the cortical cells)
- Codon formation on  $b_1...b_M$ (evidence cells in HC)
- Inputs to the b<sub>j</sub> use Brindley synapses
- Codon formation is a type of competitive learning (anticipates Grossberg, Kohonen)
- Recurrent connections to the a<sub>i</sub> use Hebb synapses



#### **Simple Representations**

- Only a small number of afferent synapses are available at neocortical pyramids for the simple memory function; the rest are needed for cortical computation.
- In order to recall an event E from a subevent X:
  - Most of the work will have to be done within the simple memory itself.
  - Little work can be done by the feedback connections to cortex.
- No fancy transformation from **b** back to **a**.
- Thus, for subevent X to recall an event E, they should both activate the same set of b cells.

#### **Recalling An Event**

- How to tell if a partial input pattern is a cue for recalling a learned event, or a new event to be stored?
- Assume that events E to be stored are always much larger (more active units) than cues X used for recall.
- Smaller pattern means not enough dendritic activation to trigger synaptic modification, so only recall takes place.

#### **Codon Formation**

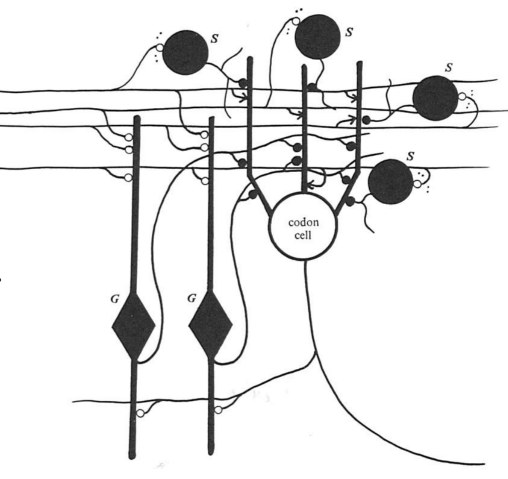
- Memory performance can be improved by orthogonalizing the set of key vectors.
  - The **b** cells do this. How?
- Project the vector space into a <u>higher dimensional</u> space.
- Each output dimension is a conjunction of a random *k*-tuple of input dimensions (so non-linear).
- In cerebellum this was assumed to use fixed wiring. In cortex it's done by a learning algorithm.
- Observation from McNaughton concerning rats:
  - Entorhinal cortex contains about 10<sup>5</sup> projection cells.
  - Dentate gyrus contains 10<sup>6</sup> granule cells.
  - Hence, EC projects to a higher dimensional space in DG.

#### **Codon Formation**

- For each input event E, different **b** cells will receive different amounts of activation.
- Activation level depends on which a cells connect to that b cell.
- We want the pattern size L to be roughly the same for all events.
- Solution: choose only the L most highly activated b cells as the simple representations for E.
- How to do this?
  - Adjust the thresholds of the **b** cells so that only L remain active.

#### Inhibition to Control Pattern Size

- S and G cells are inhibitory interneurons.
- S cells sample the input lines and supply feedforward inhibition to the codon cells.
- G cells' modifiable synapses track the number of patterns learned so far, and raise the inhibition accordingly. They sample the codon cell's output via an axon collateral.



## **Threshold Setting**

- Two factors cause the activation levels of **b** cells to vary:
  - 1) Amount of activity in the **a** cells (not all patterns are of the same size, due to partial cues)
  - 2) Number of potentiated synapses from **a** cells onto the **b** cell. This value gradually increases as more patterns are stored.
    - More cells can become active as more weights are set.
- Solution:
  - S-cells driven by codon cell afferents compute an inhibition term based on the total activity in the a<sub>i</sub> fibers.
    Assumes no synapses have been modified.
  - 2) G-cells driven by codon cell axon collaterals use negative feedback to compensate for effects of weight increases.
- Together, S and G cells provide <u>subtractive inhibition</u> to maintain a pattern size of L over the **b** units.

#### **Recall From a Subevent**

- If subevent X is fully contained in E, the best retrieval strategy is to lower the codon threshold until roughly L of the b cells are active.
- But if X only partially overlaps with E, some spurious input units will have synapses onto codon units. A better strategy is for codon cells to take into account the fraction *f* of their A active synapses that have been <u>modified</u> by learning (meaning they are part of some previously-stored pattern).
- Unmodified synapses that are active during recall can only be a source of noise.
- Thus, a b cell should only fire if a sufficient proportion f of its active synapses have been <u>modified</u>, meaning they are part of at least one stored pattern — perhaps the correct one, E.

#### **Recall From a Subevent**

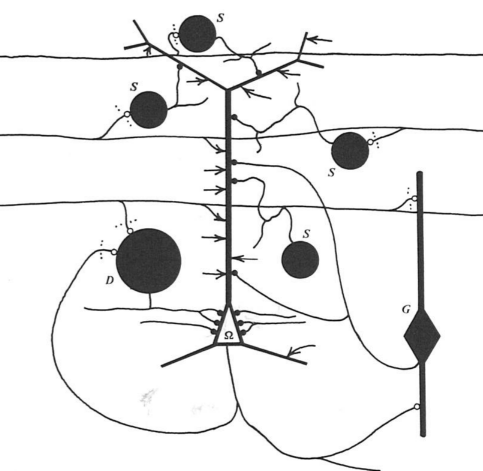
- A cell should only fire if it's being driven by enough modified synapses.
- A = number of active synapses.
- f = fraction of synapses that have been modified.
- The cell's <u>division</u> <u>threshold</u> is equal to *fA*.
- Let *S* be the summed activation of the cell:

$$S = \sum_{i} a_{i} w_{i}$$

• The cell should fire if S > fA, or S / (fA) > 1.

## **D-Cells**

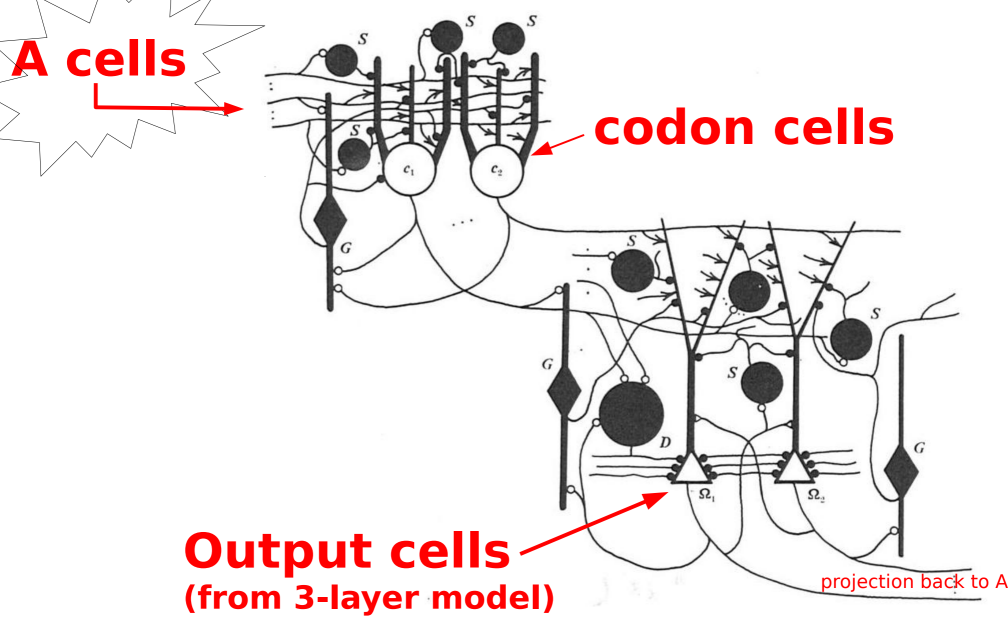
- D cells compute fA and pass it as an inhibitory input to the pyramidal cells.
- D cells apply their inhibition directly to the cell body, like basket cells in hippocampus.
- This type of inhibition causes a division instead of subtraction.
- McNaughton: division can be achieved by shunting inhibition, e.g., the chloridedependent GABA<sub>A</sub> channel.



#### **Dual Thresholds**

- Cells have two separate thresholds:
  - The <u>absolute threshold</u> T, controlled by inhibition from S and G cells, should be close to the pattern size L, but must be reduced when given a partial cue.
  - The <u>division threshold</u> *fA*, controlled by inhibition from D cells.
- Marr's calculations show that both types of thresholding are necessary for best performance of the memory.
- How to set these thresholds? No procedure is given.
  - Willshaw & Buckingham try several methods, e.g., staircase strategy: start with small f and large T. Gradually reduce T until enough cells are active, then raise f slightly and repeat.

#### 3 Layer Model: A Simple Memory With Output Cells



#### Inadequacy of the Simple Model

- Assume that  $N = 10^6 a_i$  afferents.
- Assume each neocortical pyramid can accept 10<sup>4</sup> synapses from the b<sub>i</sub> cells.
- Assume upper bound of 200 learned events per cell, due to limitation on number of afferent synapses. (Marr derived this from looking at Purkinje cells in cerebellum.)
  - Use 100 events/cell as a conservative value.
- If capacity  $n = 10^5$  events, and each **b** cell participates in 100 of them, then activity  $\alpha = 10^{-3}$ . With  $10^4$  **b** cells, only 10 can be active per event.
  - Too few for reliable representation. Threshold setting would be too difficult with such a small sample size.

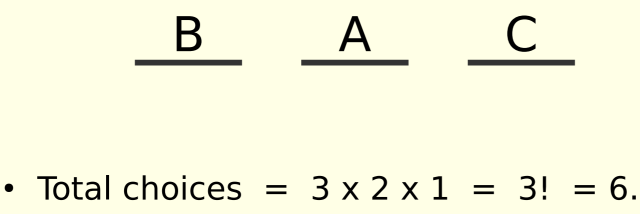
#### What's Wrong With This Argument?

- The simple model is inadequate because the activity level is too low: only 10 active units per stored event.
- But this is because Marr assumes only 10<sup>4</sup> evidence (codon) cells. Why?
  - Limited room for afferent synapses back to the cortical cells.
- This is based on the notion that every evidence (codon) cell must connect back to *every* cortical cell.
- Later in the paper he relaxes this restriction and switches to 10<sup>5</sup> evidence cells.

#### **Combinatorics 1: Permutations**

• How many ways to order 3 items: A, B, C?

- Three choices for the first slot.
- Two choices left for the second.
- One choice left for the third.



#### **Combinatorics 2: Choices**

• How many ways to choose 2 items from a set of 5?

In formal notation, what is the value of

$$\binom{5}{2} = C(5,2)$$

?

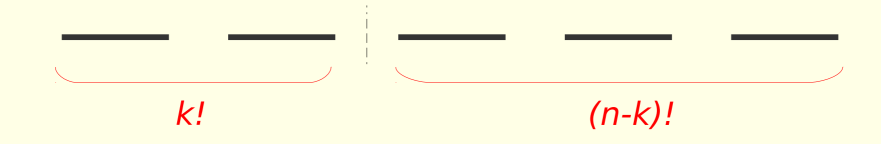
- Five choices for first item. Four choices for the second.
- Permutations of the chosen item are equivalent: combination B,E is the same as combination E,B
- So total ways to choose two items is  $(5 \times 4)/(2!) = 10$ .
- Since  $5! = 5 \times 4 \times 3 \times 2 \times 1$ , we can get  $5 \times 4$  from 5!/3!

$$\begin{vmatrix} 5 \\ 2 \end{vmatrix} = \frac{5!}{3!} / 2! = \frac{5!}{3! \cdot 2!}$$

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## Choices (continued)

- How many ways to choose k=2 items from n=5?
- Allocate 5 slots giving n! = 120 permutations:



- All permutations of the k chosen items are equivalent, so divide by k! = 2.
- All permutations of the (n-k) unchosen items are equivalent, so divide by (n-k)! = 6.

$$\binom{n}{k} = \frac{n!}{k! \cdot (n-k)!}$$

#### **Review of Probability**

- Suppose a coin has a probability *z* of coming up heads.
- The probability of tails is (1-z).
- What are the chances of seeing *h* heads in a row?

#### $z^h$

- What are the chances of seeing exactly *h* heads in a row, followed by exactly *t* tails?  $z^{h} \cdot (1-z)^{t}$
- What about seeing exactly *h* heads total in *N* tosses?

$$\left. \begin{array}{c} N \\ h \end{array} \right| \cdot z^h \cdot (1-z)^{(N-h)}$$

#### **Binomial Distribution**

 How many heads should we expect in N=100 tosses of a biased (z=0.2) coin?

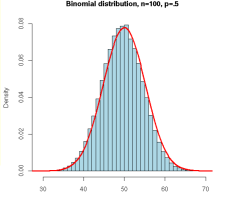
- Expected value is E < h > = Nz = 20.

 What is the probability of a particular sequence of tosses containing exactly h heads?

$$P\left[\langle t_{1,}t_{2,}\ldots,t_{N}\rangle\right] = z^{h} \cdot (1-z)^{N-h}$$

The probability of getting exactly *h* heads in any order follows a binomial distribution:

Binomial 
$$(N;z)[h] = {\binom{N}{h}} \cdot z^h \cdot (1-z)^{N-h}$$



#### Marr's Notation

- $P_i$  Population of cells.
- $N_i$  Number of cells in population  $P_i$
- $L_i$  Number of active cells for a pattern in  $P_i$
- $\alpha_i$  Fraction of active cells:  $L_i/N_i$
- $R_i$  Threshold of cells in  $P_i$
- $S_i$  Number of afferent synapses of a cell in  $P_i$
- $Z_i$  Contact probability: likelihood of synapse from cell in  $P_{i-1}$  to  $P_i$
- $\Pi_i$  Probability that a particular synapse in  $P_i$  has been modified
- $E\langle x \rangle$  Expected (mean) value of x
  - *n* Number of stored memories

#### Response to an Input Event

- Assume afferents to P<sub>i</sub> distribute uniformly with probability Z<sub>i</sub>.
- $L_{i-1} =$  number of active afferents.
- What is the expected pattern size in this population?

$$E\langle L_i \rangle = N_i \sum_{r=R_i}^{L_{i-1}} \binom{L_{i-1}}{r} \cdot (Z_i)^r \cdot (1-Z_i)^{L_{i-1}-r}$$

• What do the terms in this formula mean?

## **Response to an Input Event** $E \langle L_i \rangle = N_i \sum_{r=R_i}^{L_{i-1}} \binom{L_{i-1}}{r} \cdot (Z_i)^r \cdot (1-Z_i)^{L_{i-1}-r}$ probability a unit has EXACTLY *r* active input fibers probability a unit has AT LEAST *R*<sub>i</sub> active input fibers (so is active)

- One term of the summation is the probability that a cell will receive an input of size exactly r, given L<sub>i-1</sub> active fibers in the preceding layer.
- *r* is number of active fibers; *R*<sub>*i*</sub> is the threshold.
- Must have  $r \ge R_i$  in order for the layer *i* cell to fire. Also,  $r \le L_{i-1}$ , the pattern size for layer *i*-1.
- Large R<sub>i</sub> keeps us on the tail of the binomial distribution.
- The value of  $\alpha_i = L_i / N_i$  will be small.

Counting Active Synapses  $N_{i-1}$  cells;  $L_{i-1}$  are active  $\alpha_{i-1} = L_{i-1}/N_{i-1}$  $S_i$  synapses; x are active

Number of active synapses x is binomially distributed.

$$P(x) = \begin{pmatrix} S_i \\ x \end{pmatrix} \cdot (\alpha_{i-1})^x \cdot (1 - \alpha_{i-1})^{S_i - x}$$

 $E\langle x\rangle = \alpha_{i-1}S_i$ 

#### **Constraint on Modifiable Synapses**

Activity  $\alpha_{i-1} = L_{i-1}/N_{i-1}$ .

Proportion of synapses active at each active cell of  $P_i$  is at least equal to

the mean  $\alpha_{i-1}$  because the active cells are on the tail of the distribution. The amount by which it exceeds this decreases as  $S_i \alpha_{i-1}$  grows. Probability that a (pre,post)-synaptic pair of cells is simultaneously active is  $\alpha_{i-1} \alpha_i$ .

After *n* events, probability that a particular synapse of  $P_i$  is facilitated is:

$$\Pi_i = 1 - (1 - \alpha_{i-1} \alpha_i)^n$$

If  $\alpha_{i-1}$  is small, then  $\alpha_{i-1} \alpha_i$  is smaller, so this gives roughly

 $\Pi_i \approx 1 - \exp\left(-n\alpha_{i-1}\alpha_i\right)$ 

because for small  $\epsilon$ ,  $(1-\epsilon)^n \approx \exp(-n\epsilon)$ 

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#### **Constraint on Modifiable Synapses**

- For modifiable synapses to be useful, not all should be modified after n events are stored.
  - Otherwise we could just make all of them fixed.
- Suppose we want at most 1 (1/e) of them to be modified, which is about 63%.

$$\Pi_{i} \leq 1 - (1/e)$$
  
= 1 - exp(-1)  
 $\approx 1 - \exp(-n\alpha_{i-1}\alpha_{i})$ 

• Thus we have computational constraint C1:

$$n\alpha_{i-1}\alpha_i \leq 1$$

#### **Condition for Full Representation**

- Activity in P<sub>i</sub> must provide an adequate representation of the input event.
- Weak criterion of adequacy: change in input fibers (active cells in P<sub>i-1</sub>) should produce a change in the cells that are firing in P<sub>i</sub>.
- Cells in  $\rm P_{_i}$  just above threshold  $\rightarrow$  losing one input will shut off the cell.

#### **Condition for Full Representation**

Probability *P* that an arbitrary input fiber doesn't contact any active cell of  $P_i$  (so  $P_i$  doesn't care if it's shut off) is:

$$(1-\epsilon)^{n} \approx \exp(-n\epsilon) \qquad P = (1-Z_{i})^{L_{i}} \qquad L_{i} = \alpha_{i}N_{i}$$
$$Z_{i} = S_{i}/N_{i-1}$$
$$P \approx \exp(-\alpha_{i}N_{i}\cdot S_{i}/N_{i-1})$$

Let's require  $P < e^{-20}$  (about  $2 \times 10^{-9}$ ). Then with a little bit of algebra we have computational constraint C2:

$$S_i \alpha_i N_i \geq 20 N_{i-1}$$

#### **Summary of Constraints**

• To store lots of memories, patterns must be sparse.

Constraint C1:  $n \alpha_i \alpha_{i-1} < 1$ 

- For the encoding to always distinguish between input patterns, outputs must change in response to any input change.
  - There must be enough units and synapses to assure this.

Constraint C2:  $S_i \alpha_i N_i \ge 20 N_{i-1}$ 

 Assumes output cells are just above threshold so losing 1 input fiber will turn them off. They must be on the tail of the binomial distribution for this to hold.

#### What's Next?

- Move to a larger, three-layer, block-structured model.
- Add recurrent connections.
- Derive conditions under which recurrent connections improve recall results.
- Map this model onto the circuitry of the hippocampus.