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**THE ADAPTIVE ADVANTAGE OF SYMBOLIC THEFT OVER SENSORIMOTOR
TOIL: GROUNDING LANGUAGE IN PERCEPTUAL CATEGORIES**

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Abstract

Using neural nets to simulate learning and the genetic algorithm to simulate evolution in a toy world of mushrooms and mushroom-foragers, we place two ways of acquiring categories into direct competition with one another: In (1) "sensorimotor toil," new categories are acquired through real-time, feedback-corrected, trial and error experience in sorting them. In (2) "symbolic theft," new categories are acquired by hearsay from *propositions* – boolean combinations of symbols *describing* them. In competition, symbolic theft always beats sensorimotor toil. We hypothesize that this is the basis of the adaptive advantage of language. Entry-level categories must still be learned by toil, however, to avoid an infinite regress (the "symbol grounding problem"). Changes in the internal representations of categories must take place during the course of learning by toil. These changes can be analyzed in terms of the compression of within-category similarities and the expansion of between-category differences. These allow regions of similarity space to be separated, bounded and named, and then the names can be combined and recombined to describe new categories, grounded recursively in the old ones. Such compression/expansion effects, called "categorical perception" (CP), have previously been reported with categories acquired by sensorimotor toil; we show that they can also arise from symbolic theft alone. The picture of natural language and its origins that emerges from this analysis is that of a powerful hybrid symbolic/sensorimotor capacity, infinitely superior to its purely sensorimotor precursors, but still grounded in and dependent on them. It can spare us from untold time and effort learning things the hard way, through direct experience, but it remain anchored in and translatable into the language of experience.

THE ADAPTIVE ADVANTAGE OF SYMBOLIC THEFT OVER SENSORIMOTOR TOIL: GROUNDING LANGUAGE IN PERCEPTUAL CATEGORIES

1. Language Evolution: A Martian Perspective

Whatever the adaptive advantage of language was, it was indisputably triumphant. If all our linguistic capabilities were subtracted from the repertoire of our species today, very little would be left. Not only would all the fruits of science, technology and culture vanish, but our development and socialization would be arrested at the stage still occupied currently by the members of all other species, along with only the severely retarded members of our own. Buried somewhere among all those undeniable benefits that we would lose if we lost language there must be a clue to what language's original bonus was, the competitive edge that set us inexorably on our unique evolutionary path, distinct from all the nonspeaking species (Harnad, Steklis & Lancaster 1976; Steels 1997).

There has been no scarcity of conjectures as to what that competitive edge might have been: It helped us hunt; it helped us make tools; it helped us socialize. There is undoubtedly some merit in such speculations, but it is hard to imagine how to test them. Language is famously silent in the archeological and paleontological record, requiring interpreters to speak for it; but it is the validity of those very interpretations that is at issue here.

Perhaps we need to take a step back, and look at our linguistic capacity from the proverbial Martian anthropologist's perspective: Human beings clearly become capable of doing many things in their world, and from what they can *do*, it can also be inferred that they know a lot about that world. Without too much loss of generality, the Martian could describe that knowledge as being about the kinds of things there are in the world, and what to do with them. In other words, the knowledge is knowledge of *categories*: objects, events, states, properties and actions.

Where do those categories come from? A Martian anthropologist with a sufficiently long-range database could not fail to notice that some of our categories we already have at birth or soon

after, whereas others we acquire through our interactions with the world (Harnad, 1976). By analogy with the concept of wealth, the Martian might describe the categories acquired through the efforts of a lifetime to be those that are *earned* through honest toil, whereas those that we are born with and hence not required to earn he might be tempted to regard as ill-gotten gains -- unless his database was really very long-range, in which case he would notice that even our inborn categories had to be earned through honest toil: not our own individual toil, nor even that of our ancestors, but that of a more complicated, collective phenomenon that our (ingenious) Martian anthropologist might want to call *evolution*.

So, relieved that none of our categories were acquired other than through honest toil, our Martian might take a close look at precisely what we had done to earn those that we did not inherit. He would find that the way we earned our categories was through laborious, real-time trial and error, guided by corrective feedback from the consequences of sorting things correctly or incorrectly (Catania & Harnad, 1988). As in many cases the basis for sorting things correctly was far from obvious, he would note that our honest toil was underwritten by a substantial inborn gift, that of eventually being able to find the basis for sorting things correctly, somehow. A brilliant cognitive theorist, our Martian would immediately deduce that in our heads there must be a very powerful device for learning to detect those critical features of things (as projected onto our sensory surfaces) on the basis of which they can be categorized correctly (Harnad, 1996b). Hence he would not be surprised that this laborious process takes time and effort -- time and effort he would call "acquiring categories by *Sensorimotor Toil*" (henceforth *Toil*).

Our Martian moralist would be surprised, however, indeed shocked, that the vast majority of our categories turn out *not* to be learned by *Toil* after all, even after discounting the ones we are born with. At first the Martian thinks that these unearned categories simply appear spontaneously; but upon closer inspection of his data he deduces that we must in fact be *stealing* them from one another somehow. For whenever there is evidence that one of us has acquired a new category without first having performed the prerequisite hours, weeks or years of *Toil*, in the laborious real-time cycle of trial, error and feedback, there is always a relatively brief vocal episode between that individual and another one who has himself either previously earned that category through sensorimotor *Toil*, or has had a very brief vocal encounter with yet another individual who has himself either... and so on.

Without blinking, our Martian dubs this violation of his own planet's Protestant work ethic "the acquisition of categories by *Theft*," and immediately begins to search for the damage done to the victims of this heinous epistemic crime. To his surprise, however, he finds that (except in very rare cases, dubbed "plagiarism," in which the thief falsely claims to have acquired the stolen category through his own honest toil), category Theft seems to be largely a victimless crime.

Ever the brilliant cognitive theorist, our Martian would quickly discern that the mechanism underlying Theft must be related to the one underlying Toil, and that in principle it was all quite simple. The clue was in the vocal episode: All earthlings start with an initial repertoire of categories acquired by sensorimotor Toil (supplemented by some inborn ones); these categories are grounded by the internal mechanism that learns to detect their distinguishing features from their sensorimotor projections. These *grounded* categories are then assigned an arbitrary symbolic name (lately a vocal one, but long ago a gestural one, his database tells him [Steklis & Harnad, 1976]). This name resembles neither the members of the category, nor their features, nor is it part of any instrumental action that one might perform on the members of the category. It is an *arbitrary symbol*, of a kind with which our Martian theorist is already quite familiar with, from his knowledge of the eternal Platonic truths of logic and mathematics, valid everywhere in the Universe, which can all be encoded in formal symbolic notation (Harnad, 1990).

When our Martian analyses more closely the brief vocal interactions that always seem to mediate Theft, he finds that they can always be construed in the form of a *proposition* that has been heard by the thief. A proposition is just a series of symbols that can be interpreted as making a claim that can be either true or false. The Martian knows that propositions can always be interpreted as statements about category membership. He quickly deduces that propositions make it possible to acquire new categories in the form of recombinations of old ones, as long as all the symbols for the old categories are already grounded in Toil (individual or evolutionary). He accordingly conjectures that the adaptive advantage of language is specifically the advantage of Symbolic Theft over Sensorimotor Toil, a victimless crime that allows knowledge to be acquired without the risks or costs of direct trial and error experience.

Can the adaptive advantage of Symbolic Theft over Sensorimotor Toil be demonstrated without the benefit of the Martian Anthropologist's evolutionary database (in which he can review at leisure the videotape of the real-time origins of language)? We will try to demonstrate them in a computer simulated *toy* world considerably more impoverished than the one studied by the Martian. It will be a world consisting of mushrooms and mushroom foragers who must learn what to do with which kind of mushroom in order to survive and reproduce (Parisi, Cecconi & Nolfi, 1990; Cangelosi & Parisi, 1998). This artificial-life approach to modeling language evolution has itself evolved appreciably in the last decade (Cangelosi & Parisi, 2002; Kirby, 2000; Steels, 1997;) and is based on languages whose terms are grounded in the objects in the simulated world (Steels, 2002; Cangelosi, 2001; Steels & Kaplan, 1999).

Before we describe the simulation we must introduce some theoretical considerations that are too fallible to be attributed to our Martian theorist: One concerns a fundamental limitation on the acquisition of categories by Symbolic Theft (the symbol grounding problem) and the other concerns the mechanism underlying the acquisition of categories by Sensorimotor Toil (categorical perception).

1.1. The Symbol Grounding Problem

Just as the values of the tokens in a currency system cannot be based on still further tokens of currency in the system, on pain of infinite regress -- needing instead to be grounded in something like a gold standard or some other material resource that has face-value -- so the meanings of the tokens in a symbol system cannot be based on just further symbol-tokens in the system. This is called the symbol grounding problem (Harnad, 1990). Our candidate for the face-valid groundwork of meaning is *perceptual categories*. The meanings of symbols can always be cashed into further symbols, but ultimately they must be cashed into something in the world that the symbols denote. Whatever it is inside a symbol system that allows it to pick out the things its symbols are about, on the basis of sensorimotor interactions with them (Harnad, 1992; 1995), will ground those symbols; those grounded symbols can then be combined and recombined in higher-level symbolic transactions that inherit the meanings of the ground-level symbols. A simple example is "zebra," a higher-level symbol that can inherit its meaning from the symbols "striped" and "horse," provided "striped" and "horse" are either

ground-level symbols, or grounded recursively in ground-level symbols by this same means (Harnad 1996a; Cangelosi, Greco & Harnad, 2001).

The key to this hierarchical system of inheritance is the fact that most if not all symbolic expressions can be construed as propositions about set (i.e., category) membership. Our Martian had immediately intuited this: The simplest proposition "P," which merely asserts that the truth-value of P is true, is asserting that P belongs to the set of true propositions and not the set of false propositions. In the classical syllogism: "All men are Mortal. Socrates is a Man. Therefore Socrates is Mortal," it is again transparent that these are all propositions about category membership. It requires only a little more reflection to construe all the sentences in this paragraph in the same way, and even to redraw them as Venn Diagrams depicting set membership and set inclusion. Perceptual categories are the *gold standard* for this network of abstractions that leads, bottom-up, from "horse," "striped" and "zebra" all the way to "goodness," "truth" and "beauty."

1.2. Categorical Perception

Can perceptual categories bear the weight of grounding an entire symbolic edifice of abstraction? Some parts of the world that our senses must categorize and tag with a symbolic name do obligingly sort themselves into disjunct, discrete categories that admit of no overlap or confusion, so our senses can duly detect and distinguish them. For these happy categories it does look as if the perceptual groundwork can bear the burden. But in those parts of the world where there is anything approaching the "blooming, buzzing confusion" that William James wrote about, the world alone, and passive senses (or even active, moving, Gibsonian ones; Gibson, 1979) are not enough. Here even an active sensorimotor system needs help in detecting the *invariants* in the sensorimotor interaction with the world that *afford* the ability to sort the subtler, more confusable things into their proper categories. Neural networks are natural candidates for the mechanism that can learn to detect the invariants in the sensorimotor flux that will eventually allow things to be sorted correctly (Harnad, 1992; 1993). This is the process we have agreed to call Toil.

A sensorimotor system with human-scale category learning capacities must be a *plastic* (modifiable) one: Inside the system, the internal representations of categories must be able to

change in such a way as to sort themselves, reliably and correctly. It is perhaps an oversimplification to think of these internal representations as being embedded in a great, multidimensional *similarity space*, in which things position themselves in terms of their distances from one another, but this simplification is behind the many regularities that have been revealed by the psychophysical method of multidimensional scaling (Livingston & Andrews, 1995) which has been applied to category learning and representation in human subjects (Andrews, Livingston & Harnad, 1998). What has been found is that during the course of category learning by what we have called sensorimotor Toil, the structure of internal similarity space changes in such a way as to *compress* the perceived differences between members of the same category and *expand* the differences between members of different categories, with the effect of separating categories in similarity space that were highly interconfusable prior to the Toil (Andrews, Livingston & Harnad, 1998; Goldstone, 1994; Pevtsov & Harnad, 1997). This compression/separation in turn allows an all-or-none (*categorical*) boundary to be placed between the regions of similarity space occupied by members of different categories, thereby allowing them to be assigned distinct symbolic names.

These compression/separation effect has come to be called *categorical perception* (CP) (Harnad 1987) and has been observed with both inborn categories and learnt ones, in human subjects (Goldstone, 1994; Pevtsov & Harnad, 1997) as well as in animals and in neural nets (Cangelosi, Greco & Harnad, 2001; Harnad, Hanson & Lubin 1991; 1995; Nakisa & Plunkett, 1998; Tijsseling & Harnad, 1997). The neural nets offer the advantage that they give us an idea of what the functional role of CP might be, and what they suggest is that CP occurs in the service of categorization. It can be seen, for example, as changes in the *receptive fields* of hidden units in the supervised backpropagation nets that will be used in this study. What will be analyzed for the first time here is how the CP "warping" of similarity space that occurs when categories are acquired by sensorimotor Toil is transferred and further warped when categories are acquired by Theft. Categorical perception induced by language can thus be seen as an instance of the Whorfian Hypothesis (Whorf 1964), according to which our language influences the way the world looks to us.

2. The Mushroom World

Our simulations take place in a *mushroom world* (Cangelosi & Parisi, 1998; Harnad, 1987) in which little virtual *organisms* forage among the mushrooms, learning what to do with them (eat or don't eat, mark or don't mark, return or don't return). The foragers feed, reproduce and die. They must learn that mushrooms with feature A (i.e. those with black spots on their tops, as illustrated in Figure 1) are to be eaten; mushrooms with feature B (i.e. a dark stalk) are to have their location marked, and mushrooms with both features A and B (i.e. both black-spotted top and dark stalk) are to be eaten, marked and returned to. All mushrooms also have three irrelevant features, C, D and E, which the foragers must learn to ignore.

Apart from being able to move around in the environment and to learn to categorize the mushrooms they encounter, the foragers also have the ability to vocalize. When they approach a mushroom, they (innately) emit a call associated with what they are about to do to that mushroom (EAT, MARK). The correct action pattern (eat, mark), coupled (innately) with the correct call (EAT, MARK) are learned during the foragers' lifetime through supervised learning (Sensorimotor Toil). Under some conditions, the foragers also receive as input, over and above the features of the mushroom itself (+/-A, +/-B, +/-C, +/-D, +/-E), the call of another forager. This will be used to test the adaptive role of the Theft strategy. Note, however, that in the present simulations the *thief* steals only the knowledge, not the mushroom (cf. note 1 for simulations on environments with shared resources).

The foragers' world is a 2-dimensional (2D) grid of 400 cells (20x20). The environment contains 40 randomly located mushrooms, 10 per category. Mushrooms are grouped in four categories according to the presence/absence of features A and B: 00, A0, 0B, and AB (Figure 1). In each world there are 40 mushrooms: 10 instances of each of the four categories. Our ecological *interpretation* of the *marking* behavior is that it has two functions: Both the inedible 0B and the edible AB mushrooms have a toxin that is painful when inhaled, but digging into the earth (*marking*) immediately after exposure blocks all negative effects. There is also a delayed contingency on the AB mushrooms only, which is that wherever they appear, many more mushrooms of the same kind will soon grow in their place. So with AB mushrooms it is adaptive to remember to return to the marked spots.

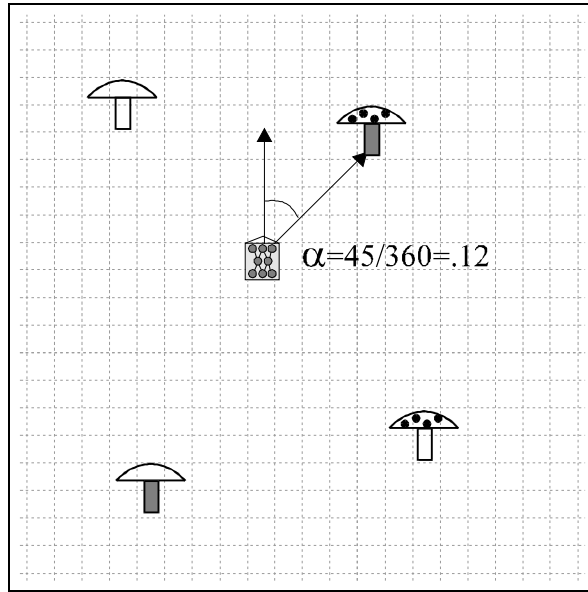


Figure 1: 2D world with one forager and the four samples of mushrooms. Mushroom feature A is the presence of black dots on the top; feature B is a dark stalk. Mushroom position corresponds to the normalized relative angle between forager's orientation and the closest mushroom.

Feature A is the black-spotted top and feature B is the dark stalk. Mushroom position is encoded as the normalized relative angle between the direction the forager is facing and the direction of the closest mushroom. In this simulation, the foraging is done by only one forager at a time. As it moves, the forager perceives only the closest mushroom. For each mushroom, the input to the forager consists of the 5 +/- features plus its location relative to the forager, expressed as the angle α , between its position and the direction the forager is facing. The angle is then normalized to the interval [0, 1]. The five visual features A, B, C, D, E are encoded in a binary localist representation consisting of five units each of which encodes the presence/absence of one feature. An A0 mushroom would be encoded as 10***, with 1 standing for the presence of feature A, 0 for the absence of feature B and *** being either 0 or 1 for the 3 irrelevant features, C, D, and E. 0B mushrooms are encoded as 01***, and AB as 11***. The calls that can be produced in the presence of the mushroom are also encoded in a localist binary system. There are 3 units for each of the three calls: 1** EAT, *1* MARK and **1 RETURN, so EAT+MARK+RETURN would be 111. Like the Calls, the three actions of eating, marking and returning are encoded with localist units.

3. The Neural Network and Genetic Algorithm

The forager's neural network processes the sensory information about the closest mushroom and activates the output units corresponding to the movement, action and call patterns. The net has a feedforward architecture (Figure 2) with 8 input, 5 hidden and 8 output units. The first input unit encodes the angle to the closest mushroom. Five input units encode the visual features and three input units encode incoming calls (if any). Two output units encode the four possible movements (one step forward, turn 90 degrees right, turn 90 degrees left, or stay in place) in binary. Three action units encode the action patterns eat, mark, and return, and three call units encode the corresponding three calls, EAT, MARK, and RETURN.

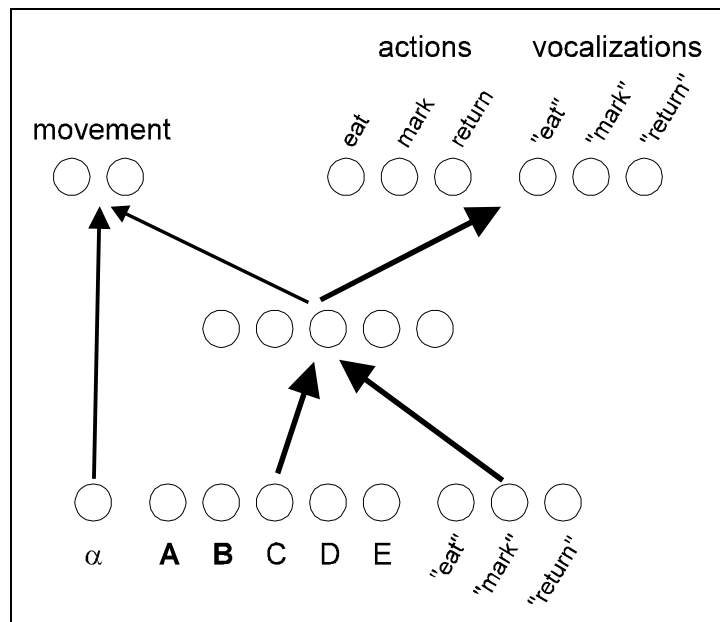


Figure 2 - Neural network architecture.

A forager's lifetime lasts for 2000 actions (100 actions in 20 epochs). Each epoch consists of sampling a different distribution of 40 mushrooms. Each action consists of two spreads of activation in the neural network, one for the action (movement and action/call) and one for an imitation task. The forager first produces a movement and an action/call output using the input information from the physical features of the mushroom. The forager's neural network then undergoes a cycle of learning based on the backpropagation algorithm (Rumelhart, Hinton & Williams, 1986).

The net's action and call outputs are compared with what they should have been; this difference is then backpropagated so as to weaken incorrect connections and strengthen correct ones. In this way the forager learns to categorize the mushrooms by performing the correct action and call. In the second spread of activation the forager also learns to imitate the call. It receives as input only the correct call for that kind of mushroom, which it must imitate in its call output units. This learning is likewise supervised by backpropagation.

The population of foragers is also subject to selection and reproduction as generated by the genetic algorithm (Goldberg, 1989). The population size is 100 foragers and remains constant across generations. The initial population consists of 100 neural nets with a random weight matrix. During the forager's lifetime its individual fitness is computed according to a formula that assigns points for each time a forager reaches a mushroom and performs the right action on it (eat/mark/return) according to features A and B. At the beginning of its life, a forager does not become much fitter from the first mushrooms it encounters because it takes some time to learn to categorize them correctly. As errors decrease, the forager's fitness increases. At the end of their life-cycles, the 20 foragers with the highest fitness in each generation are selected and allowed to reproduce by engendering 5 offspring each. The new population of 100 (20x5) newborns is subject to random mutation of their initial connection weights for the motor behavior, as well as for the actions and calls (thick arrows in Figure 2); in other words, there is neither any Lamarckian inheritance of learned weights nor any Baldwinian evolution of initial weights to set them closer to the final stage of the learning of 00, A0, 0B and AB categories. This selection cycle is repeated until the final generation.

4. Grounding *Eat* and *Mark* Directly Through Toil

Two experimental conditions were compared: Toil and Theft. Foragers live for two life-stages of 2000 actions each. The first life-stage is identical for both populations: they all learn, through sensorimotor Toil, to eat mushrooms with feature A and to mark mushrooms with feature B. (AB mushrooms are accordingly both eaten and marked.) Return is not taught during the first life-stage. The input is always the mushroom's position and features, as shown in Table 1. In the second life-stage, foragers in the Toil condition go on to learn to return to

AB mushrooms in the same way they had learned to eat and mark them through honest toil: trial and error supervised by the consequences of returning or not returning (Catania & Harnad 1988). In contrast, foragers in the Theft condition learn to return on the basis of hearing the vocalization of the mushrooms' *names*.

Table 1 - Input and backpropagation for Toil and Theft learning and for imitation learning

Condition	Feature Input	Call Input	Behavior Backprop	Call Backprop
TOIL EAT-MARK	YES	NO	YES	YES
TOIL RETURN	YES	NO	YES	YES
THEFT RETURN	NO	YES	YES	YES
IMITATION	NO	YES	NO	YES

We ran ten replications for each of the two conditions. In the first 200 generations, the foragers only live for the first life-stage. From generation 200 to generation 210 they live on for a second life-stage and must learn the return behavior. The first 200 generations are necessary to evolve and stabilize the ability to explore the world and to approach mushrooms. After the foragers are able to move in the 2D environment and to approach mushrooms, they learn the basic categories plus their names, EAT and MARK. The average fitness of the ten replications is shown in Figure 3. The populations that evolve in these 10 runs are the same ones that are then used in the Toil and Theft conditions from generations 200 to 210.

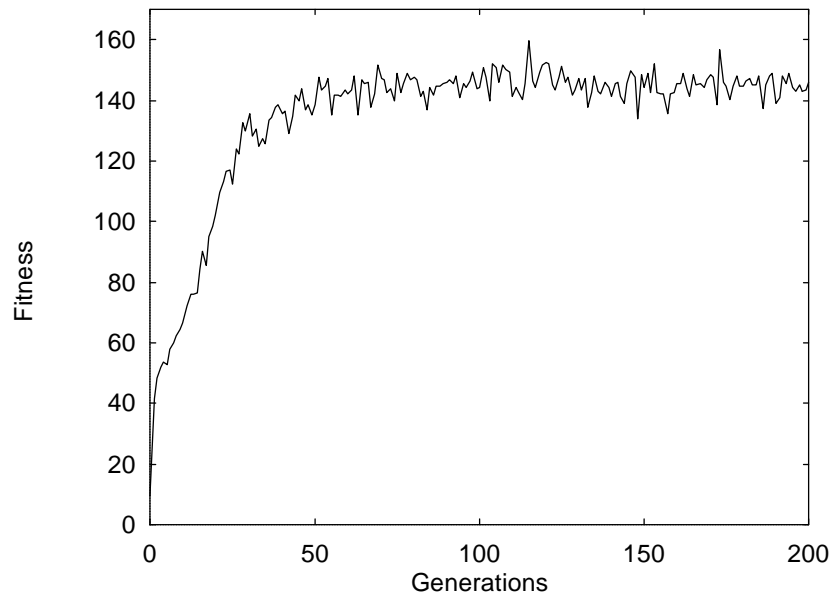


Figure 3 - Average fitness of the best 20 individuals in ten replications. Foragers lived one life-stage and only eating and marking was taught.

In the next runs, the second life-stage differs for the Toil and Theft groups: The Toil group learns to return and to vocalize RETURN on the basis of the feature input alone, as in the previous life-stage. Their input and supervision conditions are shown in Table 1. In the Theft condition the foragers rely on other foragers' calls to learn to return. They do not receive the feature input, only the vocalization input.

Our hypothesis is that the Theft strategy is more adaptive (i.e. results in greater fitness and more mushroom collection) than the Toil strategy. To test this, we compare foragers' behavior for the two conditions statistically. For our purposes we count the number of AB mushrooms that are correctly returned to. The average of the best 20 foragers in all 10 replications is 54.7 AB mushrooms for Theft and 44.1 for Toil. That is, Thieves successfully return to more AB mushrooms than do Toilers. This means that learning to return from the grounded names EAT and MARK is more adaptive than learning it through direct toil based on sampling the physical features of the mushrooms. To compare the two conditions, we performed a repeated measures analysis of variance (MANOVA) on the 10 seeds. The dependent variables were the number of AB mushrooms collected at generation 210 averaged over the 20 fittest individuals in all 10 generations. The independent variable was Theft vs. Toil. The difference between the

two conditions was significant [$F(1,9)=136.7$ $p<0.0001$]. Means and standard deviations are shown in Figure 4.

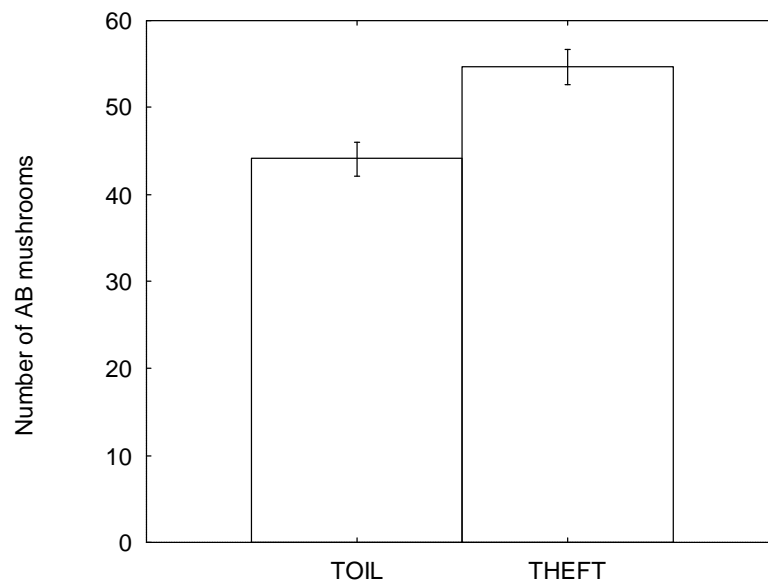


Figure 4 - Mean number of AB mushrooms correctly returned to in Toil and Theft simulations

5. Theft vs Toil: Simulating Direct Competition

A direct way to study the adaptive advantage of Theft over Toil is to see how they fare in competition against one another. We ran 10 competitive simulations, using the genotypes of generation 200 from the previous 10 runs. From generation 201 to 220, the 100 foragers of each population are randomly divided into 50 Thieves and 50 Toilers for the learning to return. There is no real on-line competition in our simulations because in each run, only one individual is tested in its world. The number of AB mushrooms to which a forager is able to return will strongly affect its fitness. Direct competition occurs only at the end of the life cycle, in the selection of the fittest 20 to reproduce. Direct competition for variable mushroom resources in shared environments has been studied separately in other simulations (note 1); in the present ecology, the assumption is that mushrooms are abundant and that the only fitness challenge is to emerge among the top 20 eaters/markers of the generation. Figure 5 shows the proportion of Thieves in the overall population of the 10 replications of Theft vs Toil (from generation

200 to 220). Even though Thieves are only 50% of the population at generation 201, they gradually come to outnumber Toilers, so that in less than 10 generations the whole population consists of Thieves.

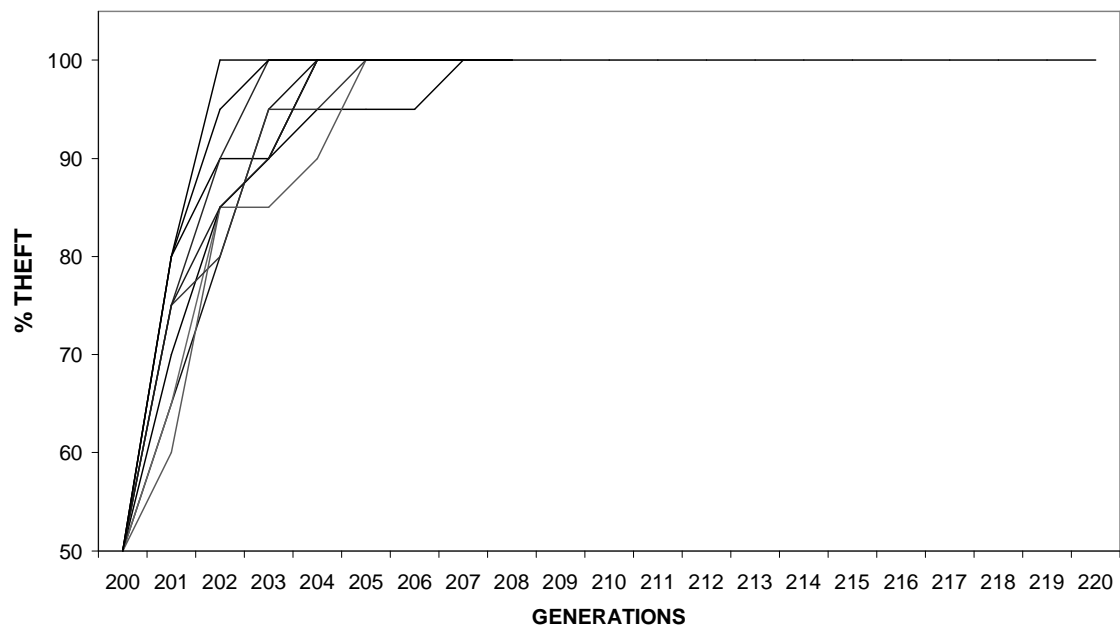


Figure 5 - Percentage of Thieves in the 10 competitive simulations.

6. What Changes During Learning? Analysis of Internal Representations

In this section we compare the changes in the foragers' hidden-unit representations for the mushrooms to determine what it is that *changes* internally during Toil and Theft. The activations of the 5 hidden units are recorded during a test cycle in which the forager is exposed to all the mushrooms as input. We will report the analysis of a single case study using the network of the fittest individual in seed 8. These results are representative of the learning dynamics in all nets that successfully learned to categorize mushrooms.

We first used Principal Components Analysis (PCA) to display the network's internal states in two dimensions, thereby reducing the 5 activations to 2 factor scores. PCA, however, has the limitation that the different conditions cannot be compared directly because of differences in

scale. For each PCA, factor scores are normalized to a distribution with a mean of 0 and a standard deviation of 1. Hence this analysis can only be used to compare internal representations within each condition, not between conditions.

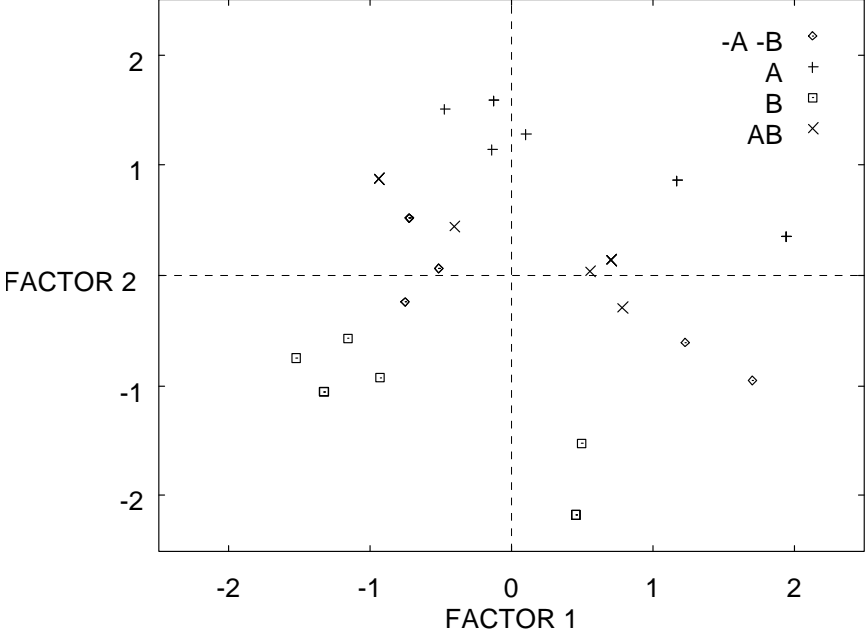


Figure 6 - Similarity space for network with random weights. Factors are obtained after PCA on the activation values of the five hidden units.

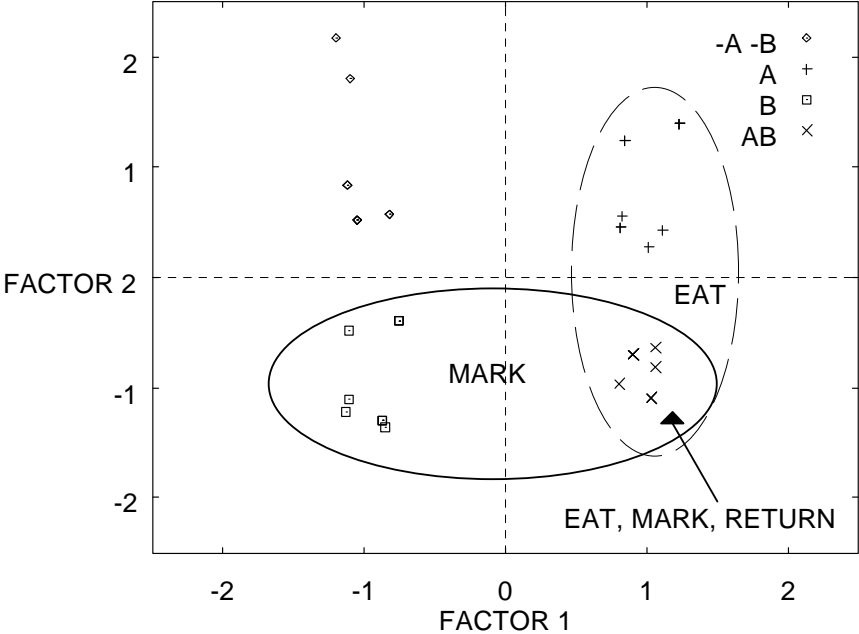


Figure 7 - Similarity space for network that learned to eat, mark, and return by Toil.

Figure 6 and 7 show the effect of category learning (Toil) on the distances between the internal representations of the mushrooms in hidden unit similarity space. In Figure 6, prior to Toil, the four kinds of mushroom are not clearly distinguishable. During the course of learning the actions/calls eat-mark-return, the representations form four separable clusters. We will now show how these representations can be used to analyze the effects of Toil and Theft learning on similarity space directly.

7. Categorical Perception Effects

The change in our networks' hidden-unit representations during the course of category learning can be analyzed and understood in terms of learned categorical perception (CP) effects (Harnad 1987, Goldstone 1994; Andrews et al., 1998), i.e. the compression of within-category distances and the expansion of between-category distances. CP has already been demonstrated to occur with Toil learning (Harnad et al., 1991; 1995; Goldstone et al., 1996; Csato et al., submitted); we will now extend this to an examination of what happens to the internal representations with Theft learning.

To overcome the limitations of the previous principal component analysis, we record the Euclidean distances between and within categories using the coordinates of the five hidden unit activations directly. At the end of each simulation, the 5 fittest foragers in each population are tested by giving them 40-mushroom samples as input. The hidden unit activations for each kind of mushroom are saved for three input conditions: (1) Features-only (only the 5-bit feature input); (2) Calls-only (only the 3-bit call input) and (3) Features+Calls (both types of input). The within-category distances are calculated as the mean squared Euclidean distances between each individual mushroom's coordinates and its category mean. There are four means, one for 00, A0, 0B, and AB respectively. These parameters reflect the within-category similarity amongst the members of each category: the lower the average within-category distance for a category, the more similar the hidden-unit representations of its members. Between-category distances are calculated as the distances between the category means. These reflect the dissimilarity between the members of different categories: the higher the average between-

category distance for a pair of categories, the greater the difference between the hidden-unit representations of their respective members.

Four learning conditions are used to analyze within-category and between-category distances for CP effects: (1) *Pre-learning*, for random-weight nets before learning; (2) *No-return*, for nets that were only taught to eat and to call EAT, and to mark and to call MARK, (3) *Toil*, for nets that also learned to return and to call RETURN with feature input, (4) and *Theft* for learning to return from calls alone. In every replication one mean was obtained for each of the 10 between- and within-category distances (4 within measures for each category, plus 6 between measures for all the possible pairings of the 4 categories) by averaging the distances derived from the 5 fittest foragers. These 10 mean distances were collected for each of the three input conditions. Because we have 10 replications, the 10 means for each distance can be used as dependent variables in two separate analyses of variance, one for within-category, the other for between-category distances. Our MANOVA for the within-category distances had two independent variables: LEARNING CONDITIONS with 3 levels (Pre, No, Toil) and CATEGORY TYPE with 4 levels (Eat, Mark, Return, Do-nothing) (note 2).

We used a repeated measure MANOVA because all levels of CATEGORY TYPE and LEARNING CONDITIONS involve repeated measures in the same set of nets. (We excluded the Theft condition in which the within-category distance is 0 because all ten samples of mushrooms use the same call input.) The chart of the average within-category distances in the 4x3 conditions is shown in Figure 8.

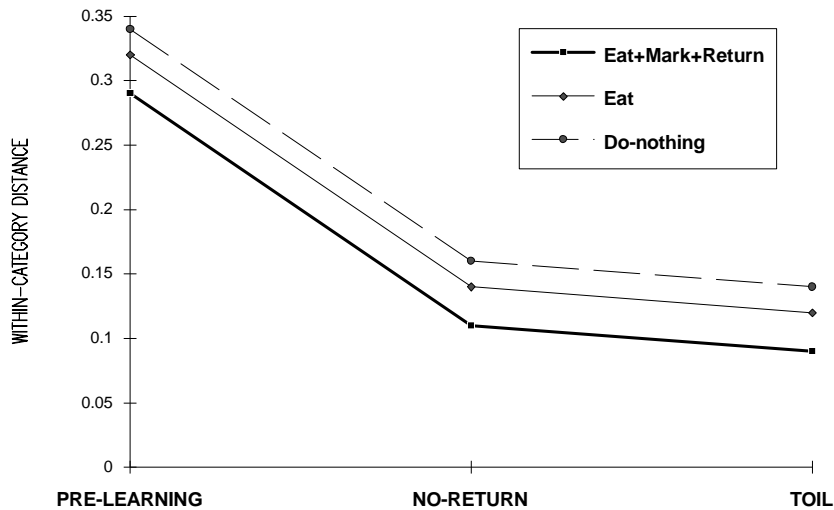


Figure 8 - Average within-category distances in the three conditions. The curve for Mark is not shown because it coincides with the curve for Eat.

The two main effects are statistically significant ($F(2,18)=917.6$ and $p<0.00001$ for LEARNING and $F(3,27)=18.8$ and $p<0.00001$ for CATEGORY TYPE); the interaction is not significant. Using the post-hoc Duncan test with a significance threshold of $p<.01$ to compare the means for each independent variable, all the comparisons in the LEARNING condition were significant. That is, within-category distances decrease significantly from Pre-learning to No-return to Toil. The biggest decrease is between the (random) Pre-learning and all the post-learning nets (Figure 8). In the four levels of CATEGORY TYPE, all means differ from each other except the Eat and Mark within-distances. That is, the within-category distance for Eat and Mark is the same, whereas the within distance of Do-nothing is the biggest and that of Return the smallest.

MANOVA for the between-category distances had two repeated variables: LEARNING CONDITIONS with 4 levels (Pre, No, Toil, Theft) and CATEGORY COMPARISONS with 4 levels (Eat Versus Mark, Eat vs Return, Eat vs Do-nothing, Return vs Do-nothing). The Mark vs Return and Mark vs Do-nothing comparisons are not included in the analysis because their means are very similar to the parallel comparisons Eat vs Return and Eat vs Do-nothing, respectively (Table 2). We then go on to generalize the results for the Eat vs Mark

comparisons. The between-category distances for the 4x4 repeated measure design are shown in Table 2 and Figure 9.

Table 2 - Table of means for the MANOVA of within-category distances

COMPARISON	PRE	NO-RET	TOIL	THEFT
EAT ↔ MARK	.57	1.47	1.47	1.42
RETURN ↔ EAT	.42	1.01	1.10	1.25
RETURN ↔ MARK	.39	1.01	1.12	1.25
EAT ↔ Do-nothing	.42	1.04	1.02	.93
MARK ↔ Do-nothing	.45	1.04	1.02	.95
RETURN ↔ Do-nothing	.54	1.42	1.52	1.61

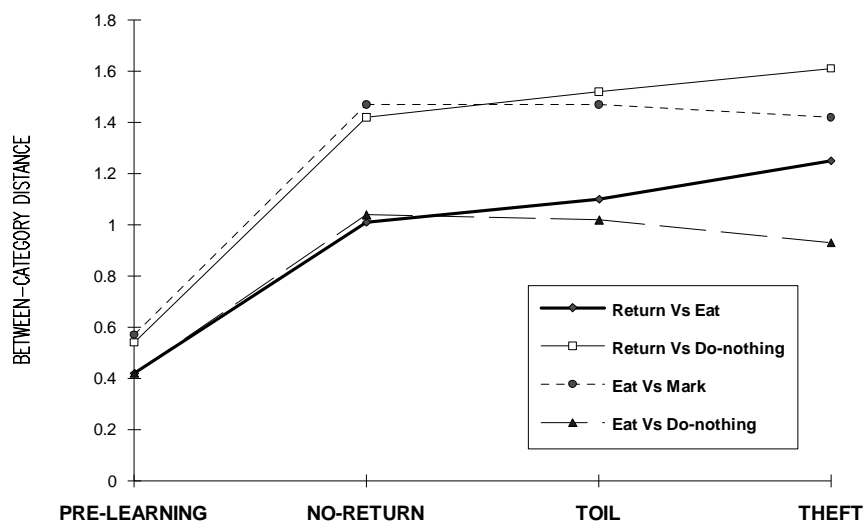


Figure 9 - Between-category distances in the four conditions. Return vs Mark and Mark vs Do-nothing are not shown because they are congruent with Return vs Eat and Eat vs Do-nothing respectively.

The two main effects are significant ($F(3,27)=3771.6$ and $p<0.00001$ for LEARNING and $F(3,27)=868.6$ and $p<0.00001$ for COMPARISONS) as is their interaction ($F(9,81)=75.7$ and

$p < .00001$). Duncan tests revealed, first, a significant difference in the distance between the Pre-learning nets and all the post-learning nets. (This expected effect only shows that any kind of systematic learning will increase between-category distances compared to random initial distances.) Comparing Toil vs Theft specifically, we see that all distances between Return and the other three categories are greater in the Theft nets. Learning Return by Theft has the effect of separating this category more from the others. The mean differences were all significant for Return vs Eat, Return vs Mark, and Return vs Do-nothing, 1.25, 1.25 and 1.25, respectively, in the Theft nets, and 1.10, 1.12, and 1.52 in the Toil nets. The Theft learning of Return caused the between-category distances not involving Return to decrease. [A last effect is that in all learning conditions the Eat vs Mark and Return vs Do-nothing distances are greater than the other pairs because the Hamming distances of their I/O codes are maximal (e.g. features A and B for Eat Vs Mark have the input contrast: 10 Vs 01).]

Figure 10 shows the change in the distances between the internal representations of the A (Eat only), B (Mark only), A&B (Eat & Mark & Return), and not-A¬-B (neither Mark nor Eat nor Return) Mushrooms. Prior to Toil, the circles, proportional to the within-category distances, are large, and the rectangle, proportional to the between-category distances is small. After Toil learning, the within-category differences shrink and the between-category distances expand.

Figure 11 then traces the between-category expansion to Theft Learning: The thin dashed rectangle is proportional to the between-category distances before learning (random). The thick dashed line is what they look like after Toil learning of Eat and Mark without Return; the thin continuous line is identical to Figure 9, that is, Toil learning of Eat and Mark, with Return, and the thick continuous line is for Theft learning of Return. Note the increased separation between A&B and not-A¬-B induced by Theft alone.

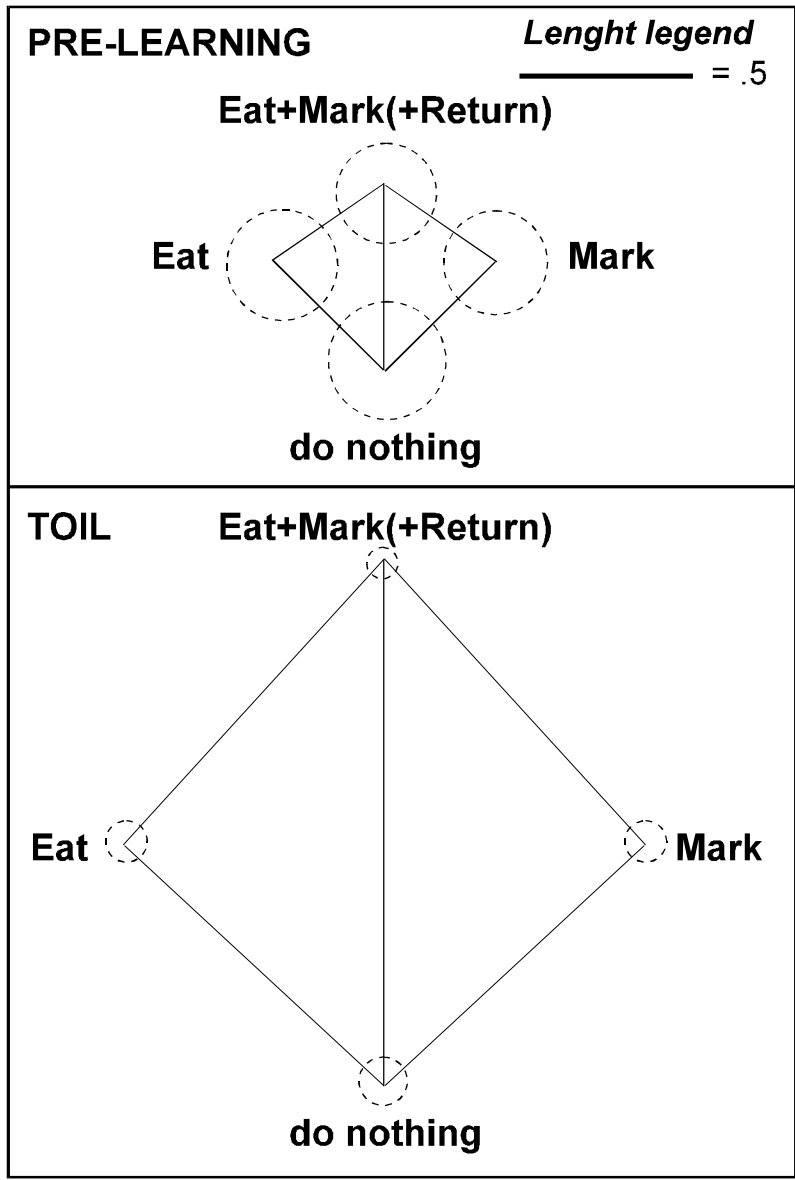


Figure 10 - 2D projections of between-category distances (quadrilateral sides) and within-category distances (circle radius) in the Pre-learning condition and after Toil learning of Eat, Mark, and Return. All distances except Eat vs Mark correspond to the actual Euclidean distances in 5 dimensional hidden unit space.

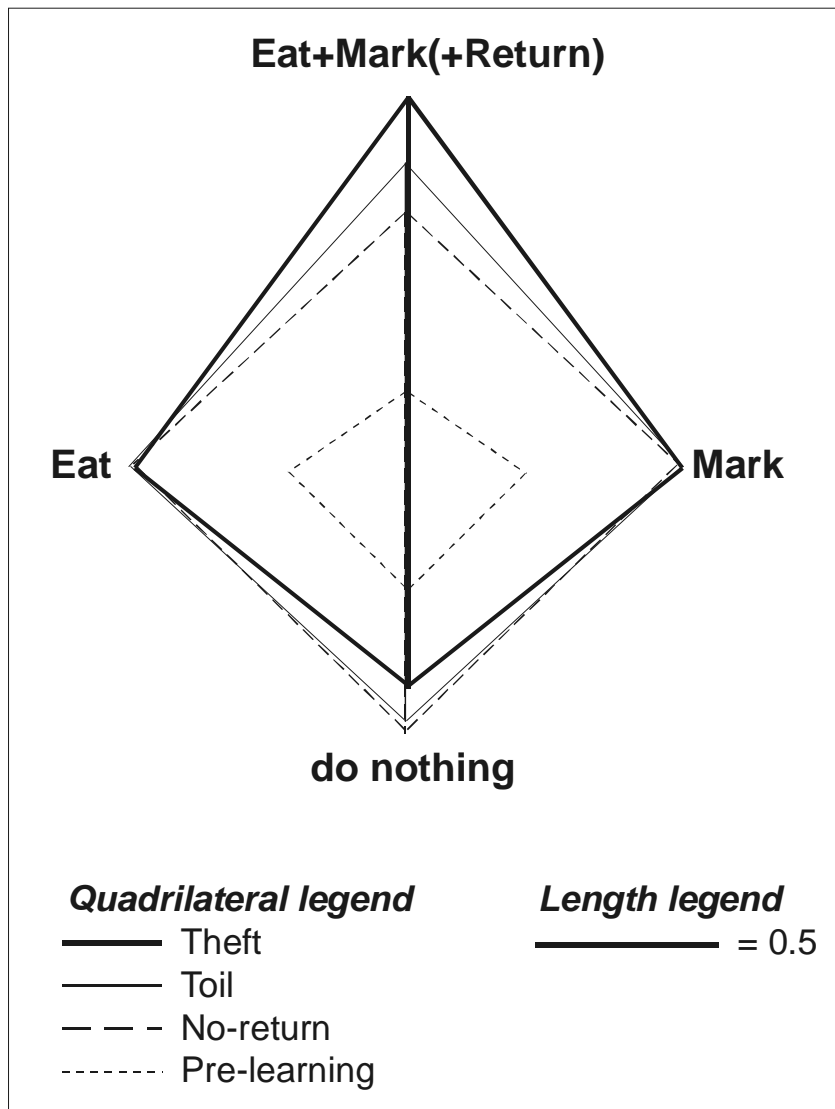


Figure 11 - 2D projections of the between-category distances (quadrilateral sides) in the four conditions. All distances, except Eat vs Mark, are comparable and reflect the actual Euclidean distances between categories (cf. length legend). Note that the distances between Return and all the other categories (Return vs Eat, Return vs Mark, Return vs Do-nothing) are the highest in the Theft condition.

8. Conclusions

We have shown that a strategy of acquiring new categories by Symbolic Theft completely outperforms a strategy of acquiring them by Sensorimotor Toil as long as it is grounded in categories acquired by Toil. The internal mechanism that makes both kinds of category

acquisition possible does so by deforming or “warping” internal similarity space so as to compress the internal representation of members of the same category and to separate those of different categories. The warping occurs primarily in the service of Toil, but Theft not only inherits the warped similarity space but can warp it further. This warping of similarity space in the service of sensorimotor and symbolic learning is called categorical perception and can be interpreted as a form of Whorfian effect (Whorf, 1964) in which language influences how the world looks to us.

From the standpoint of our Martian anthropologist, the influence would run roughly like this: All other species on this planet get their categories by toil alone, either cumulative, evolutionary toil or individual lifetime toil: Individuals encounter things, must learn by trial and error what to do with what, and to do so, they must form internal representations that reliably sort things into their proper categories. In the process of doing so, they keep learning to see the world differently, detecting the invariants, compressing the similarities and enhancing the differences that allow them to sort things the way they need to be sorted, guided by feedback from the consequences of sorting adaptively and maladaptively (as in the mushroom world).

That’s how it proceeded on our planet until one species discovered a better way: First acquire an entry-level set of categories the honest way, like everyone else, but then assign them arbitrary names. (Those names could start as nonarbitrary functional or imitative gestures at first, by-products of practical, collective social actions or even deliberate mimicry, but their nonarbitrary features would be irrelevant once they were used just to name; and vocal gestures would be least encumbered with other practical tasks, hence most readily available for arbitrary naming, especially across distances, out of eye-shot, and in the dark.) Once the entry-level categories had accompanying names, the whole world of combinatory possibilities opened up and a lively trade in new categories could begin (probably more in the spirit of barter than theft, and, within a kin-line, one of sharing categories along with other goods). In trading categories as they traded combinations of symbols, our species also traded “world-views,” for each category acquired by hearsay also brought with it some rearrangement of the internal representation of categories, a “warping” that was Whorfian, whether merely the subtle compression that results from learning that A is always conjoined with B, or the fundamental restructuring dictated by a radical scientific discovery.

Only our Martian knows the specific initial conditions in which the generative power of names and their boolean combinations made themselves felt biologically on our planet, but perhaps our simulations suggest how its benefits might have mushroomed, inducing a series of Baldwinian adaptations inclining ever our successful ancestors to name categories and to string names together so as to describe new categories to one another with ever more fervor and commitment.

Can results from a 3-bit toy world really cast light on the rich and complex phenomenon of the origin and adaptive value of natural language? This is really a question about whether such findings will “scale up” to human size in the real world. This scaling problem -- common to most fields of cognitive modeling where the tasks themselves tend not to be lifesize or to have face validity -- can only be solved by actually trying to scale our models upward, incorporating more and more of the real-world complexity and constraints into them. This is how our own research program will continue. In this paper, however, we wanted to enter our own toy candidate into the competition with the other toy models (tool-make, hunt-help, chit-chat, etc.; Knight et al., 2000) for the provenance of our species’ most powerful and remarkable trait. In other work, we have investigated categorical perception with continuous stimuli (Tijsseling & Harnad, 1997), the transfer of grounding to higher-order categories (Cangelosi, Greco & Harnad, 2001) and the emergence of syntax and compositional languages (e.g. Cangelosi, 2001).

References

- Andrews, J., K. Livingston, and S. Harnad. 1998. Categorical perception effects induced by category learning. *Journal of Experimental Psychology: Human Learning and Cognition*. 24(3): 732-53
- Cangelosi, A. 2001. Evolution of communication and language using signals, symbols and words. *IEEE Transactions in Evolutionary Computation*, 5(2): 93-101
- Cangelosi, A., A. Greco, and S. Harnad. 2000. From robotic toil to symbolic theft: Grounding transfer from entry-level to higher-level categories. *Connection Science*, 12: 143-162
- Cangelosi, A., and D. Parisi. 1998. The emergence of a "language" in an evolving population of neural networks. *Connection Science*, 10: 83-97

- Cangelosi, A. and D. Parisi. 2002. *Simulating the evolution of language*. London: Springer-Verlag
- Catania, A.C. and S. Harnad. (Eds.). 1988. *The selection of behavior. The operant behaviorism of BF Skinner: Comments and consequences*. New York: Cambridge University Press.
- Csato, L., G. Kovacs, S. Harnad, R. Pevtzow, and A. Lorincz. Submitted. Category learning, categorization difficulty, and categorical perception: Computational and behavioral evidence. *Connection Science*.
- Gibson, J.J. 1979. *An ecological approach to visual perception*. Boston: Houghton Mifflin
- Goldberg, D.E. 1989. *Genetic algorithms in search, optimization, and machine learning*. Reading, MA: Addison-Wesley.
- Goldstone, R. 1994. Influences of categorization on perceptual discrimination. *Journal of Experimental Psychology: General*, 123:178-200
- Goldstone, R.L., M. Steyvers, M., and K. Larimer. 1996. Categorical perception of novel dimensions. In *Proceedings of the Eighteenth Annual Conference of the Cognitive Science Society*.
- Harnad, S. 1976. Induction, evolution and accountability. *Annals of the New York Academy of Sciences* 280: 58-60.
- Harnad, S. (Ed.) 1987. *Categorical Perception: The groundwork of cognition*. New York: Cambridge University Press.
- Harnad, S. 1990. The Symbol Grounding Problem. *Physica D* 42: 335-346.
- Harnad, S. 1992. Connecting Object to Symbol in Modeling Cognition. In A. Clark and R. Lutz (Eds.) *Connectionism in context* Springer Verlag, pp 75 - 90.
- Harnad, S. 1993. Grounding symbols in the analog world with neural nets. *Think* 2(1): 12-78 (Special issue on "Connectionism versus Symbolism," D.M.W. Powers & P.A. Flach, eds.).
- Harnad, S. 1995. Grounding symbolic capacity in robotic capacity. In L. Steels, and R. Brooks (Eds.) *The Artificial Life route to Artificial Intelligence: Building embodied situated Agents*. New Haven: Lawrence Erlbaum. pp. 277-286.
- Harnad, S. 1996a. The origin of words: A psychophysical hypothesis In B. Velichkovsky B and D. Rumbaugh (Eds.) *Communicating Meaning: Evolution and Development of Language*. NJ: Erlbaum: pp 27-44.

- Harnad, S. 1996b. Experimental analysis of naming behavior cannot explain naming capacity. *Journal of the Experimental Analysis of Behavior* 65: 262-264.
- Harnad, S., S.J. Hanson and J. Lubin. 1991. Categorical Perception and the evolution of supervised learning in neural nets. In DW Powers & L Reeker (Eds.), *Working Papers of the AAAI Spring Symposium on Machine Learning of Natural Language and Ontology*. pp. 65-74.
- Harnad, S., S.J. Hanson and J. Lubin. 1995. Learned categorical perception in neural nets: Implications for symbol grounding. In V. Honavar & L. Uhr (Eds.) *Symbol processors and connectionist Network models in artificial intelligence and cognitive modelling steps toward principled integration*. Academic Press. pp. 191-206.
- Harnad, S., H.D. Steklis and J.B Lancaster. (Eds.) 1976. Origins and evolution of language and speech. *Annals of the New York Academy of Sciences*, 280.
- Kirby, S. 2000. Syntax without natural selection: How compositionality emerges from vocabulary in a population of learners. In C. Knight, M. Studdert-Kennedy, J.R. Hurford (eds), *The evolutionary emergence of language: Social function and the origins of linguistic form*, Cambridge University Press, pp 303-323
- Knight, C., M. Studdert-Kennedy, and J.R. Hurfor. 2000. *The evolutionary emergence of language: Social function and the origins of linguistic form*, Cambridge University Press
- Livingston, K.R. and J.K. Andrews. 1995. On the interaction of prior knowledge and stimulus structure in category learning. *Quarterly Journal of Experimental Psychology Section A-Human Experimental Psychology*, 48: 208-236.
- Nakisa, R.C., and K. Plunkett. 1998 Evolution of a rapidly learned representation for speech. *Language and Cognitive Processes*, 13: 105-127
- Parisi, D., F. Cecconi, and S. Nolfi. 1990. Econets: neural networks that learn in an environment. *Network*, 1:149-168.
- Pevtsov, R. and S. Harnad. 1997. Warping similarity space in category learning by human subjects: The role of task difficulty. In M. Ramscar, U. Hahn, E. Cambouropoulos and H. Pain (Eds.), *Proceedings of SimCat 1997: Interdisciplinary Workshop on Similarity and Categorization*. Department of Artificial Intelligence, Edinburgh University, pp.189-195.
- Rumelhart, D.E, G.E. Hinton G.E. and R.J. Williams. 1986. Learning internal representations by error propagation. In D.E. Rumelhart and J.L McClelland (Eds.), *Parallel Distributed*

- Processing: Exploration in the microstructure of cognition*, Cambridge, MA: MIT Press, vol. 1
- Steels, L. 1997. The synthetic modeling of language origins, *Evolution of Communication*, 1: 1-34.
- Steels, L. 2002. Grounding symbols through evolutionary language games. In A. Cangelosi & D. Parisi, *Simulating the evolution of language*. London: Springer-Verlag
- Steels, L., and F. Kaplan 1999. Collective learning and semiotic dynamics. In D. Floreano et al. (Eds.), *Proceedings of ECAL99 European Conference on Artificial Life*, Berlin: Springer-Verlag, pp. 679-688
- Steklis, H.D. and S. Harnad. 1976. From hand to mouth: Some critical stages in the evolution of language. In: S. Harnad et al. 1976, pp. 445-455.
- Tijsseling, A. and S. Harnad. 1997. Warping similarity space in category learning by backprop nets. In M. Ramscar, U. Hahn, E. Cambouropoulos and H. Pain (Eds.), *Proceedings of SimCat 1997: Interdisciplinary Workshop on Similarity and Categorization*. Department of Artificial Intelligence, Edinburgh University, pp.263-269.
- Whorf, B.L. 1964. *Language, thought and reality*. Cambridge MA: MIT Press

Notes

- (1) In simulations conducted by Emma Smith and Gianni Valenti (unpublished data, BSc Theses, Department of Psychology, University of Southampton, 1997) we have shown that when the scarcity of the mushrooms is varied, Theft beats Toil when there are plenty of mushrooms for everyone, but when the mushrooms are scarce and vocalizing risks losing the mushroom to the Thief, Toil beats Theft and the foragers are mute. Further studies analyzing kinship showed that under conditions of scarcity vocalizing to relatives only beats vocalizing to everyone. Of course a mushroom world is too simple, and foraging categories are not the only ones that can benefit from Theft. The pattern may be different for categories related to danger, territory, mating, dominance, or instructing offspring.
- (2) We will use the names Eat, Mark, Return, and Do-nothing (i.e. non-A, non-B mushrooms) to refer to the four categories. Return categories could also be called Eat+Mark+Return because the Return category implies the co-occurrence of behaviors/calls Eat and Mark