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Adult Cleaner Wrasse Outperform Capuchin Monkeys, Chimpanzees and Orang-utans in a Complex Foraging Task Derived from Cleaner – Client Reef Fish Cooperation

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Abstract

The insight that animals’ cognitive abilities are linked to their evolutionary history, and hence their ecology, provides the framework for the comparative approach. Despite primates renowned dietary complexity and social cognition, including cooperative abilities, we here demonstrate that cleaner wrasse outperform three primate species, capuchin monkeys, chimpanzees and orang-utans, in a foraging task involving a choice between two actions, both of which yield identical immediate rewards, but only one of which yields an additional delayed reward. The foraging task decisions involve partner choice in cleaners: they must serve visiting client reef fish before resident clients to access both; otherwise the former switch to a different cleaner. Wild caught adult, but not juvenile, cleaners learned to solve the task quickly and rehearsed the task when it was reversed. The majority of primates failed to perform above chance after 100 trials, which is in sharp contrast to previous studies showing that primates easily learn to choose an action that yields immediate double rewards compared to an alternative action. In conclusion, the adult cleaners’ ability to choose a superior action with initially neutral consequences is likely due to repeated exposure in nature, which leads to specific learned optimal foraging decision rules.

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Introduction

The ecological approach to cognition proposes that a species’ ability to solve a particular problem is tightly linked to its evolutionary history and, hence, to the ecological conditions under which it was selected [1–3]. A classic example is the tight link between spatial memory abilities and the dependency on food caching in corvids [4]. The ecological approach provides a general functional theoretical framework which allows for the integration of studies on any animal species, including invertebrates, such as the demonstration of sophisticated spatial orientation skills of bees [5], and the ability of jumping spiders to plan where to go to in order to attack prey [6]. The ecological approach has led to a great diversification of animals studied, and in particular to the appreciation that animal clades that lack particularly large and complexly structured brains may provide examples of impressive cognitive abilities. This is in particular true for fishes [7], which have provided some excellent examples for complex social strategies. Male cichlids (Astatotilapia burtoni) use transitive inference to predict fighting abilities of competitors [8] and sticklebacks (Pungitius pungitius) employ so-called hill climbing social learning strategies [9], in which they compare their own foraging success with the success of observed individuals to update foraging decisions. Another example involves the foraging decisions of cleaner wrasse, Labroides dimidiatus. These cleaner fish occupy small territories (so-called ‘cleaning stations’) in which they interact with a variety of reef fish species (so-called ‘clients’) from which they remove ectoparasites, but also mucus and scales [10]. Conflict occurs because cleaners prefer to eat mucus over ectoparasites [11], where eating the former constitutes cheating [for a review of cleaners’ decision rules, see [12,13]]. Cleaners adjust levels of cooperation to the strategic options available to clients to react to cheating by cleaners. Predatory clients typically receive the highest service quality, whereas non-predatory resident clients, who lack choice options, punish cleaners for cheating. Visiting clients who have access to alternative cleaning stations receive faster service than resident clients that have access to only one cleaning station. This is because visiting clients represent an ephemeral food source: they may swim off and visit another cleaner for their next inspection if not inspected immediately. In contrast, resident clients must wait for inspection because of a lack of alternatives. Furthermore, cleaners pay attention to the presence of potential clients and are more cooperative to current clients if that allows them to access bystanders [14]. Thus, cleaner wrasse show high adaptation to the specifics of an interaction in their foraging
decisions, which are at the same time linked to interspecific social behavior. The precision with which cleaners adapt current service quality to current conditions may be predicted by their ecology: cleaners have over 2000 interactions per day with a great variety of clients and fully depend on cleaning for their diet [15], thus their performance during the interactions has a major impact on their fitness. However, the ecological approach is rather nonspecific with respect to the cognitive processes that underlie the performance. Hence, we cannot infer from the precision and flexibility in cleaner foraging decisions that they warrant much learning, memory or comprehension and hence, ultimately any adaptive changes in corresponding brain areas. In addition, we do not know whether reaching their food maximizing decisions involves widespread learning rules or whether rather specific abilities must be evolved or developed. Thus the question of interest is whether any (vertebrate) species could easily behave like a cleaner wrasse if it switched its diet to ectoparasites and mucus of fishes, or whether specific selection pressures on cleaner wrasses have caused specific abilities? And if specific abilities do exist in cleaner wrasses, what is the role of cognition?

Here, we provide the first test of the hypothesis that cleaner wrasse foraging decisions are the result of specific cognitive abilities. Our laboratory experiment involved two identical food sources – two plates differing in colour and patterns to allow discrimination, but providing exactly the same food - where one source (plate) was ephemeral and the other one permanent. This mimicked the simultaneous visit of a resident and a visitor to the cleaning station. Accordingly, the food maximizing solution involved eating from the ephemeral food source first and only then from the permanent one. The potential difficulty of the task is due to the fact that no matter which plate an individual chooses first, it will receive exactly the same immediate reward, and only then will it (possibly) have the chance to perform a second act that would yield an additional reward. Thus, the initial decision may not lead to reinforcement learning unless an animal is somehow able to integrate the future consequences into its immediate decision. Despite theoretical considerations indicating that the task is not trivial to solve, a previous study suggested that cleaners could quickly solve it, though individual learning was not investigated [16]. In order to test whether the ability to solve the task is linked to its ecological relevance and whether the solution by cleaners reflect specific learning rules, we subjected both adult and juvenile wild-caught cleaners as well as three primate species to the task. The comparison between adult and juvenile cleaners allowed us to address the potential role of individual experience. Client composition shifts during ontogeny, with adult cleaners interacting about three times more frequently with visitors than do juveniles (comparing data published in [17,18]). Thus, juveniles rarely experience the situation in which a visitor and a resident seek cleaning simultaneously. Therefore, if adult cleaners perform better than juveniles that would indicate that individual experience in the field helps to solve the abstract laboratory task.

An important aspect of the ecological approach is to test whether other species that do not engage in cleaning interactions are less able to solve the task. We decided to use primates – capuchin monkeys, chimpanzees and orang-utans – for the comparison for several reasons. First, the general circumstances of the cleaners’ decisions involve social interactions and foraging, which matches the two contexts that have been proposed to select for large brains in primates [19–22]. Second, primates, and in particular our three study species, have been shown to possess a large array of cognitive mechanisms in the context of social behavior and foraging. Specifically, all three species have a complex diet and have been classified as extractive foragers [21]. In addition, at least chimpanzees and capuchins hunt for meat and catch mobile insects and reptiles [23–25], and in doing so, encounter ephemeral food sources. Moreover, all three species are able to solve some cooperation tasks in the laboratory [26–32], and capuchins and chimpanzees do so in the wild [23–25,35].

Also, our task involved the ability to take not only immediate but also future consequences into consideration, an ability that primates have repeatedly demonstrated in foraging experiments (delayed rewards experiments: [34,35]; planning experiments: [36–39]). Finally, all three of our primate study species have large brains compared to other species, and large relative brain sizes (e.g., brain-to-body or neocortex-to-body ratios) even compared to other primates [40], again indicating high general cognitive abilities.

Although evidence suggests that the primates will excel in tasks that involve future consequences in the context of cooperation and foraging, the specifics of our task may favor cleaners. For example, cooperation and foraging are intertwined in cleaners in a way that is absent in primates; most importantly, cleaners cooperate with their food sources. In addition, primates encounter ephemeral food sources (e.g., insects, small vertebrates) unpredictably and opportunistically, and thus the ecological constraints are quite different from those of the fish, for whom the interaction with ephemeral sources is predictable. Based on this, we predicted that unlike the cleaner wrasse, the primates would not perceive the task as a social interaction but just as an optimal foraging task. Thus, our experiment offered us the opportunity to test the ecological intelligence hypothesis in a quite specific way. We expect that if ecology is the driving force that helps to solve the problem, then cleaners should individually learn to solve the tasks faster than any of the primate species. Conversely, if the general context and brain size (relative or absolute) prepare better for the task than rather specific ecological conditions do, then the primates should learn to solve the task faster than the cleaner wrasse. We also considered an additional way to test the role of learning for the cleaners’ decision making process, reversing the role of the two plates once an individual reached the learning criterion. The former permanent plate now became the ephemeral plate and vice versa. Although cleaners are able to discriminate between different client categories, including resident and visitor, and can even individually recognize clients [13], reversal of roles does not occur under natural conditions, i.e. a visitor individual/species never turns into a resident. Therefore, it appears to be highly unlikely that reversal learning could be aided by the adaptation of an innate program. For the primates we included this task only to see whether once the task has been solved, they understood its general principle. We predicted that if primates found the task initially difficult but solving it triggered a more general understanding, then their performance would greatly improve during the reversal.

Results

Initial learning tests

All six adult cleaner fish individuals learned to eat first from the ephemeral plate, which was smoothly withdrawn if the cleaner were to forage on the permanent plate first. Individuals took 3–10 sessions (of 10 trials each) to reach the criterion of significance with a median of 4.5 sessions. In contrast to the adult cleaners, only one juvenile cleaner and two out of four chimpanzees solved the task within 10 sessions, and all other subjects failed (Fig. 1). Thus, there was a significant difference in learning speed between the species/age classes (Kruskal-Wallis Test: df = 4, H = 18.4, p = 0.001). Post-hoc comparisons revealed that adult cleaners performed better than juvenile cleaners or any of the three primate species (Student
Figure 1. The number of trials required for individuals to learn to eat first from the ephemeral plate. Dots represent an individual. The y-axis indicates the number of trials required to learn the task. doi:10.1371/journal.pone.0049068.g001

Figure 2. The number of trials required for individuals to reverse their preference when the plates switched roles (e.g., the permanent tray became ephemeral and the reverse). Again, dots represent an individual, and the y-axis indicates the number of trials required to reverse the preference. doi:10.1371/journal.pone.0049068.g002

Discussion

A key conclusion from our experiment is that the sophisticated foraging decisions which cleaner wrasses demonstrate during interactions with client reef fish are not easily achieved by other species with larger and more complexly organized brains. The ability to choose between an ephemeral and a more permanent food source of otherwise identical quality is apparently far from simple as the vast majority of individuals from three primate species that otherwise excel in cognitive tasks failed to learn the task within 100 trials, as did juvenile cleaners. However adult cleaners consistently solved the task. Thus, our task differs from experiments that demonstrate extremely fast learning of solutions if individuals are placed into a key stimulus-response context, in which even invertebrates like bees may outperform primates, including humans [42–43].

Why the task may be difficult to solve

When confronted with a choice that directly yields two different amounts of food primates can easily discriminate outcomes with one reward from those with two [44–47] (for that matter, fish can do the same [48,49]), even in cases in which the quantity to be received is indicated symbolically (e.g., via tokens or Arabic numerals [50–52]). Thus there must be another explanation for the decrement in performance in the primates as compared to the adult wrasses. We consider several possibilities for why this task may be difficult to learn. First, assuming that both species saw the food source of otherwise identical quality is apparently far from simple as the vast majority of individuals from three primate species that otherwise excel in cognitive tasks failed to learn the task within 100 trials, as did juvenile cleaners. However adult cleaners consistently solved the task. Thus, our task differs from experiments that demonstrate extremely fast learning of solutions if individuals are placed into a key stimulus-response context, in which even invertebrates like bees may outperform primates, including humans [42–43].

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little novelty or information value left for the second plate, lowering the incentive as compared to the first plate/reward. Thus phenomena like blocking (e.g., little conditioning is occurring) or overshadowing (e.g., less conditioning is occurring to this weaker conditional stimulus) might explain why there seem to occur little learning about the second plate if the first plate already has been rewarded.

Second, it is possible that the fish experienced the removal of the plate as a stronger punishment than did the primates. Both the fish and the primates presumably reacted to the removal of the second plate, containing food, as a negative reinforcer (e.g., punishment). However, fish may have additionally experienced it as a social punishment; one indication that they indeed perceive the task as a cleaning situation is that they respond with tactile stimulation when the plate returns, a behaviour cleaners use to reconcile and to make clients stay longer under natural conditions [53] to encourage it to stay this time. Hence negative social reinforcement (or: social punishment) would make the task more aversive, and hence easier to learn, for the adult fish as compared to the primates and juvenile fish, both of which have far less experience with this situation.

Finally, a more cognitive mechanism than associative learning that would allow subjects to solve the task is insight based on backwards induction. In backwards induction, one has to start with the desired endpoint and then figure out which steps lead to that endpoint. Evidence for backwards induction has been demonstrated in a chimpanzee, Julia, who had to open up to 10 Plexiglas boxes with specific tools inside in the right sequence to finally obtain food in the last box [36]. However, the primates in our study apparently failed to use backward induction, despite a large number of trials. Given the evidence for insight learning in our primate species, why did they fail to use this ability? One possibility again relates to reinforcement; Julia was not rewarded for each step of her process, while in our experiment the subjects were. As discussed above, it is possible that the receipt of intermediate rewards interferes with learning mechanisms in that it lowers the incentive value of the second reward [54].

We finally note that the apes’ unexpectedly low performance on the reversal task was likely due to frustration with the procedure. Apes – including some of these subjects – are typically very good at reversal learning tasks [41,42]. Moreover, within the primates, reversal learning performance is associated with brain size [55], and apes typically outperform capuchins [56]. However, our subjective impressions indicated that the apes found this task very frustrating. Despite there being only 10 trials in a session, we initially had to change the ITI from 5 minutes to 90 seconds in order to get them to complete a session (see Methodological considerations, below, for a more detailed discussion of this). Even with the 90 second ITI, by the later sessions, apes were hitting or grabbing the choice trays rather than choosing a reward, and often refused to participate. We believe it was this frustration with the task that caused the unexpectedly low performance on reversal learning. What is perhaps more notable is that the fish did so well. Their behavior is counter to that predicted by the primes’ association between reversal learning and brain size [55] and deserves far more attention as a potential area in which fish cognition equals that of the larger-brained primates (see Methodological considerations, below, for other areas in which fish cognition appears to equal that of far larger brained species).

Why adult cleaner wrasses may have been able to solve the task

We propose two non-mutually exclusive explanations for why adult cleaners learned to solve the task. First, the cleaners may have developed the decision rule to preferentially approach ephemeral food under natural conditions and then applied the same rule to this task. In contrast, the primates were born in captivity, where sufficient food is provided multiple times per day (at all facilities) and they rarely catch ephemeral food like invertebrates. Second, as discussed above, the cleaners may have perceived the task as a social interaction. In that case they would have perceived the removal of the ephemeral plate as the loss of a cooperation partner and hence as a negative reinforcer that reduced the likelihood that the subject would choose the permanent plate again on future trials. The aversion to losing any client would make the ephemeral plate more attractive to cleaners, whereas primates are not selected to experience either the negative reinforcement of a missed opportunity or social reinforcement for interacting with their foraging substrate. Thus, we consider it likely that cleaners, but not the primates, simultaneously experienced a positive and a negative reinforcer, which would explain why they learned to solve the task rather quickly as compared to the primates. If that was the case, a change in protocol for the primates that let them perceive the interaction as social (for example by replacing the trays with human partner) should yield much faster learning.

If our hypotheses are correct then one would also predict that even individuals of the closely related cleaner wrasse species L. bicolor should have problems solving the task. This is because adult bicolor individuals rove over large areas and typically approach the clients they want to interact with rather than having to wait for them at a cleaning station [57]. Thus, the distinction between residents and visitors is not crucial to them, and they can follow clients that are about to leave in order to prolong interactions. For bicolor individuals, it appears to be mainly important where an interaction takes place within their home range: they are more cooperative in their core area than in the periphery [58]. To explore this hypothesis, we additionally collected preliminary data on L. bicolor. We tested two individuals in May 2009 at the University of Neuchâtel following exactly the same protocol as we used for adult L. dimidiatus. One bicolor failed to learn the initial task within 200 trials. The other one learned the initial task in 70 trials but failed at the reversal: after a short period of random choices it redeveloped a preference for the initially ephemeral plate. Taken together, there is thus a significant difference in overall performance (in trials to complete the entire experiment) between the two species (Mann-Whitney-U-Test, m = 6, n = 2, U = 0, p<0.05). Clearly, more bicolor individuals should be tested (fortunately, they are very difficult to obtain from licensed commercial pet shops; three individuals were all we managed to obtain over a six week search period, with one not willing to participate in the experiment). Nevertheless, the preliminary results suggest that the ability of L. dimidiatus individuals to solve the task is linked to very specific ecological conditions that are not met in L. bicolor.

A comparison between juvenile and adult cleaners

There are various potential explanations for why juveniles failed to solve the task while adults managed. One possibility is that maturation processes in the brain preclude juveniles from solving the problem at hand. Second, there were small differences in the experimental protocol due to different research sites and in turn testing possibilities: juveniles – kept on Lizard Island for the period of the experiment - experienced longer time intervals between subsequent trials as compared to the adults which where housed and tested in Neuchâtel, Switzerland. However, in an earlier experiment adult cleaner fish that were trained on a similar task (i.e. “one plate remains until inspected while the other does not”,
Methodological considerations

We note that the primates consistently performed poorly despite the fact that we ultimately adapted the methods to be as appropriate as possible for each species (within the constraints of using trays with different 'behaviours' to present identical foods). In particular, the capuchin monkeys received several different methodological modifications as we attempted to optimize a protocol which allowed them to eventually solve the initial task. Variables that seemed to have helped them included a barrier between the plates and much shorter time intervals between trials (Table 1). These shorter inter-trial intervals may have reduced the cost of an incorrect choice for the primates, potentially making it less likely for them to learn the task. However, we note several things which argue against this possibility. First, in the initial phase of pilot testing with capuchin monkeys, all subjects received ten sessions of ten trials each (100 total trials, a trial number which allowed all adult cleaners to learn the task) with 15 minute ITIs between each trial. No subjects' choices differed from chance (binomial test, all ps > 0.05 both for individual sessions and when sessions are combined). Four additional subjects received an additional four session (40 total trials, for a cumulative total of 140 trials) with 15 minute ITIs between each trial, and again, no subject's choices differed from chance (all ps > 0.05). Finally, in the last phase of testing, in which subjects learned the test, ITIs were reduced to 5 minutes (still with 10 trials per session, so an additional 100 trials total). We note that there were several other factors that changed between these tests. First, of course, there is an experience effect; however even with 100 trials with 15 minute, no capuchin learned within 100 trials, yet all of the fish did so. Additionally, in the first phase only, the ephemeral tray was pulled back, but not removed from sight as it was in later phases. While this may have confused the primates, they are accustomed to food rewards being visible, but unavailable from their daily life. In fact, being able to see the food that they could not access could arguably have increased the magnitude of the negative reinforcement for choosing the other plate, possibly supporting learning. Finally, there was no divider between the trays in the first phase of testing, which may have made it more difficult for the primates to discriminate between the choices. However, subjects had to reach through one of two discrete doors, actively pushing open the door in the process, so it is difficult to see how they could not discriminate between the options.

The capuchins were all tested prior to the tests with the apes, and as much as possible we used the final capuchin protocol for both ape species. The choice tray featured a divider, and non-chosen options were immediately removed from sight. One thing that we could not do similarly was the five minute ITI. In pilot testing, the chimpanzees and orang-utans reacted with extreme frustration to a 5 minute delay, leaving the testing area and refusing to return. Thus we shortened the ITI to 90 seconds to encourage subjects to participate. If, as we feared, the capuchins' behaviour was positively influenced by the shorter ITI, then a shorter one yet for the apes should have made the task easier. Additionally, while this might have resulted in less cost per choice, subjects still only received 10 trials per session, so there were very few chances to receive treats during testing (in most cognitive tests, subjects receive at least two to three times this many trials in a session). Overall, this meant that the details of the procedure were optimized during the course of the study for the primates, but not for the fish. Therefore, we consider it unlikely that cleaners outperformed primates due to advantages with respect to methodological details like the color or shape of trays, the food, or the inter-trial interval.

Finally, note that the primates acquired food by reaching out and grasping it, while fish swam to different foods and took them directly into their mouths. This was due to differences in body plan between fish and primates. Fish have to move between compartments with their whole body, but from where they were located could easily see both rewards simultaneously. Due to the size of the primates and caging constraints, it was impossible to house them such that they could simultaneously see both rewards and be housed in a third room. This would be particularly problematic for our study if they could not immediately view the ephemeral reward being removed when they chose the permanent reward first. Moreover, this procedure would have meant that the primates had far longer time intervals to both access the first reward and between the first and second. Additionally, primates typically make choices by grasping with their hands. Of course, when comparing species with very different body plans and abilities, identical procedures may be difficult or impossible, both for practical reasons (e.g., the presence or absence of hands) and differences in experience or ways of interacting with the world. In particular in cases such as ours, in which a species performs differently than expected, we encourage the use of multiple procedures in an effort to optimize the design for the species, even if this results in some methodological differences.

In conclusion, our results provide the first evidence that cleaners' sophisticated behavior in cleaning interactions is due to selection for specific rule learning that require experience and/or maturation. All three primate species have a complex diet and are known to cooperate, but still they were outclassed by adult cleaners in this foraging task. Although we cannot entirely rule out differences in procedure that resulted from the comparison between very different species, a possible mechanism underlying the fishes' response is that they perceive the leaving of a food source as a negative reinforcer, and therefore choose the ephemeral food source first before approaching the permanent one. This implies that the specificity of the cleaners' ability to give priority to ephemeral food sources lies not in a sophisticated cognitive process but in the ability to identify relevant stimuli. Nevertheless, recent research on fishes has yielded evidence for various supposedly more complex cognitive abilities (reviewed by [61]). As mentioned before, nine-spined sticklebacks use social learning rules that compare own success relative to the success of potential models [62,63], and male cichlids may use transitive inference to assess the strength of potential rivals [64]. But there
are many other examples. Various coral reef fishes have spawning traditions [65,66]. Fathead minnows show the ability to generalise between predators [67]. Groupers signal their intention to hunt to moray eels in the absence of prey [68]. Guppies' performance in relative quantity judgments adheres to that of humans tested in non verbal numerical tasks [69], and also mosquitofish can use numbers like primates [70]. On the neurophysiological level there is recent evidence that the reward structure of fish brains is similar to that of mammals: hedonistic rewards like receiving tactile stimulation may yield fitness advantages [71]. With respect to cleaner wrasse, we note that they express many abilities in the context of cleaning interactions, including adjustment of service quality to the presence of a co-inspecting partner [72] and to the presence of an audience [14], the use of predatory clients as social tools against chasing non predatory clients [73] and the ability to remember the ‘when’ and ‘what’ of interactions [74]. These phenomena are not specific to cleaner wrasse – client interactions, and they have been linked to more complex cognitive processes like social awareness [75] and an understanding of other individuals as agents [76] in studies on primates. It will therefore be of interest to use cleaner wrasse to test in how far such higher cognitive processes might be present or absent in a ‘lower’ vertebrate that is nevertheless under similar selective pressures of a complex social environment.

### Methods

Experiments on adult cleaner wrasse were carried out in March and April 2009 at the University of Neuchâtel, Switzerland, while juvenile cleaner wrasse were tested at Lizard Island Research Station, Australia, in July and August 2010. Experiments on capuchins (August to December 2009) and chimpanzees (August to December 2010) were carried out at the Language Research Center, Georgia State University, USA, and orang-utans were tested at Zoo Atlanta, USA (August to December 2010).

### Subjects and housing conditions

**Cleaner wrasse.** Six adult wild caught *Labroides dimidiatus* (5.5–7.6 cm total length; TL) of unknown sex were purchased from a licensed pet shop. All adults were individually housed in aquaria measuring 60×30×30 cm³ in size and filled with

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**Table 1. A summary of information about subjects and experimental protocol.**

|                | Adult wrasse | Juv. wrasse | Chimpanzees | Orang-utans | Capuchins |
|----------------|--------------|-------------|-------------|-------------|-----------|
| **General**    |              |             |             |             |           |
| N individuals  | 6            | 7           | 4           | 4           | 8         |
| Date           | 3–4/09       | 7–8/10      | 8/10–4/11   | 8/10–4/11   | 8–12/09   |
| Location       | Neuchâtel, CH| Lizard Island, AUS | LRC GSU, USA | Zoo Atlanta, USA | LRC GSU, USA |
| **Experiments**|              |             |             |             |           |
| Time between trials | 15 min | 30 min | 90 sec | 1. 90 sec | 1. 15 min |
| Subject order  | varied       | varied      | varied      | varied      | varied    |
| Plate color    | red-yellow   | green-grey  | blue        | blue        | green-blue violet-yellow blue-yellow |
| Food type      | mashed prawn | mashed prawn | banana      | cheerios cereal | apple    |
| Plate preference test | yes | yes | yes | yes | 1. no |
| Food already on plate/tray | yes | yes | yes | no | yes |
| Removed plate/tray | out of view | out of view | out of view | out of view | 1. visible |
| **Initial learning test** |              |             |             |             |           |
| Maximum N sessions | 10 | 10 | 10 | 10 | 10 |
| Trials per session | 10 | 10 | 10 | 10 | 10 |
| N sessions per day | 2 | 1 | 1 | 1 | 1 |
| N test days per week | 7 | 7 | 3 | 5 | 3 |
| **Reverse learning test** |              |             |             |             |           |
| Maximum N sessions | 10 | - | 10 | 10 | 10 |
| Trials per session | 10 | - | 10 | 10 | 10 |
| **modifications** |              |             |             |             |           |
|                    |              |             |             |             |           |
|                    |              |             |             |             |           |
|                    |              |             |             |             |           |
|                    |              |             |             |             |           |

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experiments. Individuals were kept for 10 days prior to commencing prawn flesh or a mixture of mashed prawn flesh and fish flakes. to ensure sufficient food intake independent of their performance trained to feed off grey Plexiglas plates, and fed ad libitum every day to ensure sufficient food intake independent of their performance during experiments. All individuals were initially fed mashed prawn flesh or a mixture of mashed prawn flesh and fish flakes. Individuals were kept for 10 days prior to commencing experiments.

Primates. All primates were captive born. The eight brown adult capuchin monkeys (5 males, 3 females, age range 5–20 years, median age of 10 years) were from two separately housed social groups at the Language Research Center of Georgia State University, USA. The four chimpanzees (2 males, 2 females, age range 25–40 years) were also from the Language Research Center, whereas the four orang-utans (3 males, 1 female, age range 7–33 years) were from Zoo Atlanta, USA. All primates lived in stable social groups consisting of adult male(s) and female(s) and any attendant offspring. They were separated from these groups only for behavioral and cognitive testing. Details regarding housing conditions are provided in Information S1. Subjects were fed a diet according to their species-specific needs, but generally consisting of primate chow and fresh fruits and vegetables. They also received enrichment-foods several times per day; consequently, animals were never food or water deprived for testing purposes. Running water was available ad libitum at all times. Subjects could choose not to participate at any time by walking away from the experimenter.

General procedure

The experimental design was based on a study by Bshary & Grutter [16]. Some variation occurred in the way the three primate species were tested (Table 1). Both fish and monkeys had to make an initial choice between two visually distinct food plates/trays, both offering the same food in equal quantity: 0.0001–0.001 g pieces of prawn (highly preferred food) for the fish, 0.5×0.5×0.5 cm³ dried apple pieces for the capuchins, cheerios for the orang-utans and 5 mm slices of banana for the chimpanzees. Plates and trays varied in size, shape and color between species in species-appropriate ways (see Table 1). All plates and trays were attached to handles so that they could be moved towards subjects but also be retracted rapidly. The “ephemeral” plate would be removed immediately if not chosen first by the subject, whereas the “permanent” tray would remain until the subject had taken the food item. Which plate was which this required reaching outside of the compartment. The experimental compartment of the aquarium. After approximately 60 s, the client plates were placed at equal heights at the opposite end of the aquarium, i.e. the experimental compartment. After about 10 s, the door was opened and the cleaner could enter the experimental compartment at will.

For capuchin monkeys, members of each social group, consisting of four subjects, were simultaneously tested in separated test chambers attached to their home enclosure. Monkeys had previously been trained to be separated from their social group and to individually enter these chambers, where virtually all testing was done. Dependent offspring were always allowed into the testing area with their mothers. Testing chambers measured 61×44.5×33 cm³ in size and were separated from each other by approximately 40 cm. The test chamber was backed by an opaque panel, allowing vocal, but no visual or tactile, access to their group. This allowed us to interact with subjects in a controlled manner with minimal distractions from the group. The sessions for the apes were organized in a similar way: subjects were all tested in a subsection of the indoor section of their home enclosures, while still in auditory and visible contact with the other group members (this is how all testing is done at these facilities). The order in which subjects were tested varied from day to day. As with the capuchins, dependent offspring were always allowed into the testing area with their mothers. Note that for all species, acquiring the food required accessing a separate area from where the subjects were initially located. For fish, this required swimming, while for the primates this required reaching outside of the compartment.

Presentation of plates

The position of the two plates was randomized, but with an equal number of presentations on each side within each 10 trial sessions. Randomization was constrained such that the same tray was never presented more than three consecutive times on the same side. (Note that capuchins were initially tested with the plates altering sides between sessions; see Table 1). The two plates were placed far enough apart that, following a choice of the permanent plate, the experimenter could remove the ephemeral plate before the subject could take the food. It proved impossible to put the trays far enough apart to stop the capuchins from grabbing both food items simultaneously, so we added trapdoors to allow access to only one at a time. These consisted of two doors attached to each other by a string that worked in a drawbridge-like fashion, pulling one door closed when the other was pushed open. No special constructions were required for the great apes, as the mesh structure of the cage prevented them from quickly grabbing both items simultaneously. The procedures were identical for the reversal learning phase, except that the plates’ behavior was reversed abruptly, i.e., the previous ephemeral plate now behaved like the permanent while the previous permanent plate became the ephemeral plate. There were differences between experimental groups concerning the number of sessions per day (one or two; all of ten trials), the number of testing days per week (every day for cleaners, but 5 days per week for orang-utans and 3 days per week for capuchins and chimpanzees), and the time interval between successive trials (15 or 30 min for the cleaners and generally shorter for the primates). See Table 1 for specific details.

Learning criterion

We based our significance criterion on Sign-Tests-Table (two-tailed). Significance was reached when a subject made correct choices on ≥8/10 trials on one session or ≥8/10 on two or ≥7/10 trials on three consecutive sessions. For capuchins in the initial sessions, the criterion for learning was ≥16/20 trials (e.g., over 2 sessions) because plate positions were constant during a session and hence a side bias would have led to the inaccurate assessment of significant “learning” in half of sessions (and significant “anti-learning” in the others). Once an individual had reached criterion,
we ran the reversal trials. We used the same criterion for the reversal test. We were primarily interested in relative performance rather than the question whether all subjects can learn to develop a food maximizing preference eventually if given sufficient opportunity. Adult cleaner wrasses were the first to be tested out of all the experimental groups. They formed the baseline for the others with respect to the questions we attempted to answer. As all of them learned to solve both the initial and reversal tasks within 100 trials, we fixed 100 trials as an upper limit for the other experimental groups. Because the reversal learning task required learning of the initial task, we decided to expose any primate that failed to learn the initial task within 100 trials to modified versions of the task. The modifications were designed to facilitate learning (see Table 1). Some of the modifications were included in the reversal learning task, such as re-baiting of the ephemeral plate for reversal learning task, such as re-baiting of the ephemeral plate for (see Table 1). Some of the modifications were included in the reversal learning task, such as re-baiting of the ephemeral plate for Puzzle 13: Cognition, Evolution, and Behavior, 2nd edition. Oxford: Oxford University Press. p. 720.

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