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Insect-Inspired Sequential Inspection Strategy Enables an Artificial Network of Four Neurons to Estimate Numerosity

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SUMMARY

Varying levels of numerical cognition have been found in several animal species. Bees, in particular, have been argued to be able to count up to four items and solve complex numerical tasks. Here we present an exceedingly simple neural circuit that, when provided with the actual visual input that the bee is receiving while carrying out the task, can make reliable estimates on the number of items in the display. Thus we suggest that the elegance of numerical problem solving in bees might not lie in the formation of numerical concepts (such as "more," "less," or "zero"), but in the use of specific flight movements to scan targets, which streamlines the visual input and so renders the task of counting computationally inexpensive. Careful examination of the actual inspection strategies used by animals might reveal that animals often employ active scanning behaviors as shortcuts to simplify complex visual pattern discrimination tasks.

INTRODUCTION

Numerical cognition is traditionally considered a higher cognitive ability, perhaps because of its association with the most advanced human intellectual achievements. The symbolic, language-based mathematics we use, however, appears to be rooted in a predisposition to use quantitative information without symbolic representation, which exists in pre-verbal infants and in cultures that do not use symbols for counting (Dehaene, 2001; Feigenson et al., 2004). A growing body of experiments demonstrates that a wide range of animals possess a similar "number sense." Not only birds (Nieder, 2018; Pepperberg, 2006; Rugani, 2018) and mammals (Matsuzawa, 2009; Nieder, 2018) or other large-brained animals but also fish, frogs/toads, and even insects with miniature brains were shown to be able to make decisions based on numerosity (reviewed in Agrillo and Bisazza, 2014, 2018; Pahl et al., 2013; Rose, 2018; Skorupski et al., 2018). Bees, in particular, exhibit counting-like abilities and can be trained to search for food after a given number of landmarks (Chittka and Geiger, 1995; Dacke and Srinivasan, 2008; Menzel et al., 2010) or on the stimulus with a given number of items (Skorupski et al., 2018), and can use the number of items as the decision criteria in a match-to-sample task (Gross et al., 2009). Recently, honeybees have been argued to even understand numerical concepts of "less than," "greater than," and "zero" as a number (Howard et al., 2018).

However, how complex is numerical cognition in neurocomputational terms? Computer vision algorithms, often based on convolutional neural networks, are abundantly used for counting objects in images and offer a good starting point for addressing this question (e.g., Dijkstra et al., 2018 and references therein). Most of these algorithms rely on object detection and then count the detected instances, but because such methods explicitly make use of symbolic mathematics, they are not accessible for animal brains. However, computer vision has also proved that it is possible to reliably estimate object count without detecting and localizing individual object instances, using, for example, image density (Lempitsky and Zisserman, 2010; Rahnemoonfar and Sheppard, 2017). As for the size of the neural network necessary, Dehaene and Changeux (1993) proposed a formal model of only 480 neural units (plus 50 input units) to account for the elementary numerical abilities of infants and animals. This model is able to extract approximate numerosity from images, up to five items. It seems likely that the perception of numerosity is a basic attribute of visual systems (Burr and Ross, 2008) and emerges spontaneously when neural networks are trained to encode statistical properties of images (Stoianov and Zorzi, 2012).

Here we explore how serial processing reduces the size of the neural hardware required for basic counting. It seems likely that bees cannot extract complex visual pattern properties "at a glance" (Nityananda et al., 2014); they inspect pattern elements from up close (Guiraud et al., 2018; Ings et al., 2012) and one by one

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Figure 1. Simple Neural Model for Counting

The phasic "brightness" neuron extracts the change in brightness from the visual input. The working memory neurons in the second layer are recurrent and thus maintain exponentially decaying memory traces. The "brightness working memory" neuron receives strong input from the "brightness" neuron, and signals recent changes in brightness. The "counting working memory" neuron receives weak input from the "brightness" neuron, and so accumulates information about the changes in brightness over a longer period. Finally, the "evaluation" neuron subtracts the "counting working memory" from the "brightness working memory." Its response is inversely proportional to the number of brightness changes, and, with the right visual input, it provides an online evaluation of the numerosity of the stimulus.

(Skorupski et al., 2018). If such an inspection strategy is indeed universal, it has profound implications for the complexity of visual tasks. We present a simple abstract model of only four neural units that, when provided with the responses that known low-level visual neurons would produce during such a sequential scan, is able to match the bees' performance in a complex numerical ordering task (Howard et al., 2018; Skorupski et al., 2018). The output of this network is sufficient to distinguish between numerosities up to 4–6, produces an appropriate response to an empty set ("zero"), and reproduces Weber's law of number discriminability.

RESULTS

Our simple model (Figure 1) employs just four independent neural units (which we will refer to as neurons for simplicity). It is able to mimic the counting abilities of bees-provided that it receives sequential visual input of the countable items. The first neuron is a wide-field (60° visual angle) neuron that sums up the responses of a collection of phasic on-off narrow-field cells of the medulla (Arenz et al., 2017). Neurons with similar response properties have been found in the second and third visual ganglia (medulla and lobula) of insects (Douglass and Strausfeld, 2003; Hertel, 1980; Paulk et al., 2009a; Yang and Maddess, 1997). This phasic "brightness" neuron responds to changes in brightness within its receptive field. The model is not limited to this specific type of input but will provide comparable results when using input from a global brightness detector or an edge detector (Figure S1). The "brightness working memory" neuron receives strong excitatory input from the "brightness" neuron and feeds back to itself; thus its response will be close to maximum when the bee encounters a change in light intensity. The "counting working memory" neuron also feeds back to itself, but it is only weakly stimulated by the "brightness" neuron; thus its response will be proportional to the number of times the bee has moved between dark and bright areas. Note that numbers are not registered as integers, but accumulated as magnitudes (as in the approximate number system described in humans for estimating numerosities higher than 4; Feigenson et al., 2004). Finally, the "evaluation" neuron is excited by the "brightness working memory" neuron and inhibited by the "counting working memory" neuron. The "evaluation" neuron thus accumulates information about the stimulus while the bee is inspecting it, and so this neuron provides a continuously updating evaluation of the numerosity of the stimulus (Figure 1).

The flight path of the bee, and the resulting visual input sequence, is crucial for generating a correct evaluation. A recent experiment (Skorupski et al., 2018) analyzed the bees' flight trajectories when choosing



Figure 2. Estimating Numerosity in a Counting Task

Using realistic visual input and following simple rules for interpreting the evaluation provided by the neural network, the model can provide enough information to reproduce the decisions the bee made in the counting task from Skorupski et al. (2018).

(A) The flight path of a bee trained to choose two items and not four items. The bee inspects each item one by one, flying over them at a distance of 1–2 cm. During training, the bee was rewarded with sugar solution hidden in a hole in the middle of the correct stimulus (indicated by a small circle). When a stimulus is chosen, the bee hovers in front of the hole, trying to feed from it; when rejected, she leaves the stimulus without trying to feed. Reproduced from Skorupski et al. (2018).

(B) Example set of visual input to the model during scanning. From a short distance, with limited field of view, the visual input is akin to moving a spotlight across the image.

(C) The responses of the "brightness" neuron (gray line), "brightness working memory" neuron (dotted line), the "counting working memory" neuron (dashed line), and the "evaluation" neuron (black line) for the flight path shown in (A). Decisions to land on a stimulus (light gray arrow) are made when the scan is finished (there are no more items in sight) and the response of the "evaluation" neuron is high (above approximately 0.8 here). Decisions to leave the stimulus (black arrows) are triggered when the "evaluation" falls below a threshold (approximately 0.8 here). When the bee decides to leave a stimulus, the network is reset, and it is reactivated once the bee has left the stimulus.

between patterns with different number of elements and concluded that bees inspect pattern elements sequentially, flying over each item once. Moreover, bees keep very close to the pattern during scanning (Guiraud et al., 2018; Ings et al., 2012). We assumed a 1–2 cm viewing distance; from this distance, our wide-field neuron's receptive field of 60° only covers 1.2–2.3 cm in diameter of the stimulus. Thus its input is akin to a moving spotlight across the pattern (Figure 2).

Our network outputs an estimate of the numerosity that can be used to solve a number of different types of quantification tasks (counting, equals to, less than, or more than) depending on how the "evaluation" neuron's response is interpreted in decision making. For the "choose two not four" task from Skorupski et al. (2018), we employed the rule that the bee is prompted to leave the stimulus if the "evaluation" neuron's response falls below a critical level (approximately 0.8 here). This can happen if the bee has passed over more items than she is looking for (>3 items here) and when she has been searching for the sugar reward without success long enough for the working memory to degrade. We also assume that the bee would land on the stimulus should she complete a scan without being



Figure 3. Estimating Numerosity in a Numerical Ordering Task

The model reproduces the choices of the bee in a complex numerical ordering task from Howard et al. (2018), including the preference for the empty stimulus ("zero") and an increasing success of discrimination with increasing numerical distance

(A) The evaluation given by the model for the stimuli used in Howard et al. (2018) shows a decreasing response with increasing numerosity. Each stimulus contains a varying number (0–6) of differently shaped (circle, triangle, or square) items. 1, 3, 15, or 21 individual patterns per number were used. The dots represent the "evaluation" responses to each pattern, the black lines indicate their means, and the gray areas depict the standard deviations. We assumed the bee scans each pattern by flying over each item once, as described in Skorupski et al. (2018).

(B) If the bee chooses patterns to scan randomly and lands with a likelihood directly proportional to the state of the "evaluation neuron" at the end of the scan, the distribution of landings on zero versus other numerosities follows Weber's law of number discriminability (as found in Howard et al., 2018). Note that this decision rule does not involve comparing two stimuli.

See also Figure S1 and Table S1.

prompted to abandon it. In either event, the decision to leave or find the sugar reward will reset the network by inhibiting both the "brightness working memory" and the "counting working memory" neurons. Using these assumptions, we find that the model reliably predicts the rejections and landings shown in Skorupski et al. (2018) based on the visual input during its flight path (Figure 2). The same model outputs can be used for selecting "higher number" or "greater" by inverting the decision rule, i.e., the bee should leave the stimulus after finishing the scan but lands on it once the evaluation falls below the threshold. Moreover, the same method can be used to address landmark counting (Chittka and Geiger, 1995; Dacke and Srinivasan, 2008). Here, the bee is viewing large landmarks from a distance, instead of small items from close up; the behavioral rule is to interrupt flight and land when the "evaluation" neuron's response falls below a threshold.

We then turned our attention to a recent study in which bees succeeded in numerical ordering tasks (including an adequate response to zero), which was interpreted by the authors as indicating that bees form concepts of "less than," "greater than," and "zero" (Howard et al., 2018). The flight paths were not recorded in this article, compelling us instead to assume an idealized flight path based on the scanning rules described in Skorupski et al. (2018), one that passes over each item once, at a hovering distance of 2 cm. The decision to land on the feeder is determined by the value of the "evaluation" neuron after completing the scan. For the "less than" task, the decision to land would be guided by a high value, whereas the "more than" task would be guided by a lower value. Note that the network does not compare two sets of stimuli; instead, a decision is made concerning each stimulus independently based on its overall score. We simulated the input to the neural network that could result from flying a scanning path and calculated the output from the "evaluation" neuron to predict the probability of choosing a pattern. For the stimuli used in Howard et al. (2018), the model's numerosity estimations match the performance of the bees (Figure 3 and Table S1--"less than" task; for the "more than" task, see Table S1). The model's accuracy when choosing from two stimuli follows Weber's law, which states that accuracy is expected to improve with numerical distance. Finally, the model reliably rates "zero" (an empty sheet) over other numbers in the "less than" task. Our counting network

actually outperformed the behavior of real bees; they failed at choosing zero over two in the original experiment.

DISCUSSION

The model rests on the assumption that during counting bees scan pattern elements sequentially and visit each element once (proposed in Skorupski et al., 2018). There is evidence to show that this scanning behavior might be required to extract visual pattern information. The time bees require to solve a visual discrimination task increases with task complexity (Ings et al., 2012; Nityananda et al., 2014), indicating that they use some sort of sequential scanning behavior. As bees observe pattern elements from a distance of only a few centimeters (Guiraud et al., 2018; Ings et al., 2012), the larger part of the pattern will be out of the bees' field of view at any given time, suggesting a one-by-one inspection strategy. How bees know which pattern element they have already visited is not yet fully understood; however, as they can clearly avoid revisits to previously emptied feeders (Lihoreau et al., 2012) and previously emptied nectaries within one flower based on their position (Bar-Shai et al., 2011), we expect them to be able to use their working memory to avoid revisits to pattern elements as well. The visual inspection strategies that other non-human animals use during counting remain unexplored; there is, however, evidence for animals following a set visual path during object recognition tasks (in chickens; Dawkins and Woodington, 2000). Overall, the sequential scanning behavior described in Skorupski et al. (2018) might be a version of a more universal strategy, co-opted and tailored to the task of counting.

The model offers a non-countable magnitude estimation similar to the core system 1 humans employ for approximating large numbers, and not the core system 2 we use to discriminate a small number of items "at a glance" (subitizing) (Feigenson et al., 2004; but see Gallistel and Gelman, 2000 for the argument that both systems use magnitude estimation). As the success of discrimination is expected to be ratio dependent only in the first case, and this has been argued for bees in recent experiments (Howard et al., 2018), there is some support for this kind of implementation, but more experiments will be needed. Consider, however, that there are other known examples of approximate magnitude estimation circuits in insect brains. Ants, for example, measure distance by integrating step count (Wittlinger et al., 2006, 2007), whereas bees keep track of the total amount of image motion in the lateral regions of their visual field (Srinivasan et al., 1997; Stone et al., 2017). The "item counter" might use the same neural architecture as these distance meters, or even recruit the visual distance circuit directly.

The crucial element in our neural architecture is the implementation of this "item counter" as a recurrent working memory circuit. Similar configurations have been found in insect brains (Douglass and Strausfeld, 2003; Eichler et al., 2017; Grünewald, 1999; Haag and Borst, 2001), and recurrent microcircuits have been suggested to serve the role of memory units encoding the distance traveled during path integration (Stone et al., 2017). In our model, recurrence creates an exponentially decaying memory trace that can be modified by further inputs. Behavioral experiments using a delayed-matching-to-sample paradigm have established that the bees' working memory (early short-term memory) indeed degrades in an exponential fashion and disappears in less than 10 s (Raine and Chittka, 2011; Zhang et al., 2005). In our model, we used one neuron (as theorized in Loewenstein and Sompolinsky, 2003). However, working memory is more likely to be implemented by a group of recurrently wired neurons. The memory capacity of such recurrent neural networks, called echo-state networks, depends on their size and on the level of noise (Jaeger, 2002); a small network, receiving the inevitably noisy visual input, is in line with the limited counting abilities of bees and of many other species.

The location of visual working memory in the bee brain is unknown. For olfactory learning, the antennal lobes have been identified as most likely sites for working memory/early short-term memory (Erber et al., 1980; Menzel, 1999); the most likely analog sites in the visual pathway would be in the medulla or lobula. In the medulla, recurrently wired neurons might exist in the serpentine layer (Douglass and Strausfeld, 2003); in the lobula, in the inner layers (Haag and Borst, 2001). There is evidence that in flies some short-term memory traces are stored in the central complex (Liu et al., 2006); for bees, Stone et al. (2017) put forward an argument for a recurrent microcircuit serving as the memory unit in the path integrator in the central complex. The linear neurons we used here are capable of less computation than a more complex spiking neuron, and even if we need to scale up the number of working memory neurons an order of magnitude to counterbalance neural noise, the necessary tens of neurons would still fit easily in the optic lobes or the central complex. Two simple working memory circuits, each of which receives the same input only weighted differently, may be easily co-opted for counting by neurons in

the mushroom body (Eichler et al., 2017; Grünewald, 1999; represented in our model as one "evaluation" neuron). To summarize, neurons of the types that are required for our simple theoretical model do exist in the insect brain. Although it is likely that more than four neurons are involved in real counting tasks (perhaps with multiple similar circuits operating in parallel), such circuitry would still not be prohibitively expensive to be accommodated in an insect brain.

The advantage of explicitly outlining a possible neural architecture is that such a model yields testable predictions. Our model makes the following predictions (among others). (1) The counting circuit we proposed relies on inputs from the color blind motion pathway, and so expected to use only long-wavelength-sensitive receptor inputs and be color blind as well (Paulk et al., 2008, 2009b). (2) The signal of the working memory circuit that acts as an "item counter" degrades exponentially with time; thus introducing delays during a scan is expected to cause the bee to underestimate numerosity. (3) We propose that the bee tailors her scan to the stimulus, in a way that keeps input noise to the minimum; forcing the bee to an altered scan route (e.g., viewing the stimulus from a larger distance than ideal) should degrade the performance. (4) The model makes the decision to land on a stimulus based on its numerosity, without directly comparing two stimuli. With appropriate tuning, this decision rule is enough to reliably choose the smaller or the larger numerosity but cannot be used to choose the middle number from three numbers; thus bees should fail at his task.

Fifty years ago, the ability to learn concepts was considered uniquely human and a sign of the highest form of intelligence. We now know from behavioral experiments that bees can solve the delayed-match-to-sample (Gross et al., 2009), sameness/difference (Giurfa et al., 2001), and above/below tasks (Avarguès-Weber et al., 2011), and can count and extrapolate to zero (Howard et al., 2018), but the behavioral strategies by which they do so and whether these indeed require the formation of concepts is a separate question. Our artificial neural network of a few nodes, when given appropriately structured visual input, can solve a variety of tasks without requiring any form of concept or "understanding." The model demonstrates this principle applied to counting and numerical ordering; however, counting is not the only learning task whose computational complexity has been called into question recently. After viewing an image, bees are able to distinguish between the same and a novel image, and can be trained to choose either the same or the different one, a task that appears to require the concepts of sameness and difference (Giurfa et al., 2001); modeling work, however, suggested a simple but neurobiologically plausible circuit that matches the bees' performances, and which does not involve any top-down processing (Cope et al., 2018). In the "above-andbelow" spatial conceptual task, bees are required to decide if any object is above or below a referent (Avarguès-Weber et al., 2011). A recent behavioral study that examined the bees' flight paths during this task (Guiraud et al., 2018) proposed that bees could turn the spatial relation task into a simple discrimination task by only inspecting the bottom part of any pattern and making a decision based on whether it is the referent (above which the item in question is found, thus the answer is "above") or anything else (thus the answer is "below"). Honeybees have been shown to selectively opt out from making a choice when they are uncertain, an indication of metacognition; however, the authors argue that the same neural circuit that implements simple associative learning could govern the behavior (Perry and Barron, 2013). Although it is often argued that cognitive ability correlates with brain size (see, e.g., Kotrschal et al., 2013 for an example on numerical abilities in guppies), our results and these studies show that seemingly advanced cognitive performance can be achieved with extremely small circuits. If this is so, a careful re-examination of the potential evolutionary advantage of bigger brains is in order (Chittka and Niven, 2009).

In comparative cognition, there is little value in rating cognitive task difficulty based on how difficult it is using symbolic human-like thinking. What is worthy of our attention is the repertoire of innate and learnt behavioral routines that animals employ while completing the task, and the complexity of the task in terms of neurocomputation. Within this framework we have shown that counting and numerical ordering are computationally inexpensive, provided the animal employs an active, sequential scanning of pattern elements. Here we studied the scanning behavior of bees; similarly simple computational solutions may underpin numerical cognition in other animals that employ active scanning (e.g., Chittka and Skorupski, 2017; Dawkins and Woodington, 2000; Gegenfurtner, 2016). Furthermore, we have shown that counting does not need to rely on the internal representation of concepts. Sequential scanning drastically reduces the demand for the neural hardware required to solve the task. We conclude that active scanning behavior could play a major role in even the most complex cognitive tasks. Future studies in comparative cognition should benefit from shifting the focus from *what* an animal can do to *how* it does it and explore the intricacies of the sequential decision-making process (Chittka et al., 2012).

Limitations of the Study

(1) The model parameters were chosen to demonstrate that the circuit is able to estimate numerosity, but we do not investigate the learning or other processes that lead to the emergence of the synaptic weight parameters. (2) We do not test the robustness of the model to variations in input stimuli other than that inherent in the stimuli used in Skorupski et al. (2018) and Howard et al. (2018). We decided to omit these tests as a realistic estimate of model robustness would require more information on the scanning behavior (and thus the visual input) and the neural implementation than available.

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Transparent Methods, two figures, one table, and one data file and can be found with this article online at https://doi.org/10.1016/j.isci.2018.12.009.

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AUTHOR CONTRIBUTIONS

Conceptualization, V.V. and L.C.; Methodology, V.V.; Writing – Original Draft, V.V., Writing – Review & Editing, V.V. and L.C.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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REFERENCES

Agrillo, C., and Bisazza, A. (2018). Understanding the origin of number sense: a review of fish studies. Philos. Trans. R. Soc. Lond. B Biol. Sci. 373, 20160511.

Agrillo, C., and Bisazza, A. (2014). Spontaneous versus trained numerical abilities. A comparison between the two main tools to study numerical competence in non-human animals. J. Neurosci. Methods 234, 82–91.

Arenz, A., Drews, M.S., Richter, F.G., Ammer, G., and Borst, A. (2017). The temporal tuning of the *Drosophila* motion detectors is determined by the dynamics of their input elements. Curr. Biol. 27, 929–944.

Avarguès-Weber, A., Dyer, A.G., and Giurfa, M. (2011). Conceptualization of above and below relationships by an insect. Proc. Biol. Sci. 278, 898–905.

Bar-Shai, N., Keasar, T., and Shmida, A. (2011). The use of numerical information by bees in foraging tasks. Behav. Ecol. *22*, 317–325.

Burr, D., and Ross, J. (2008). A visual sense of number. Curr. Biol. 18, 425–428.

Chittka, L., and Geiger, K. (1995). Can honey bees count landmarks? Anim. Behav. 49, 159–164.

Chittka, L., and Niven, J. (2009). Are bigger brains better? Curr. Biol. *19*, R995–R1008.

Chittka, L., Rossiter, S.J., Skorupski, P., and Fernando, C. (2012). What is comparable in comparative cognition? Philos. Trans. R. Soc. Lond. B Biol. Sci. 367, 2677–2685.

Chittka, L., and Skorupski, P. (2017). Active vision: a broader comparative perspective is needed. Constr. Found 13, 128–129.

Cope, A.J., Vasilaki, E., Minors, D., Sabo, C., Marshall, J.A.R., and Barron, A.B. (2018). Abstract concept learning in a simple neural network inspired by the insect brain. PLoS Comput. Biol. 14, e1006435.

Dacke, M., and Srinivasan, M.V. (2008). Evidence for counting in insects. Anim. Cogn. 11, 683–689.

Dawkins, M.S., and Woodington, A. (2000). Pattern recognition and active vision in chickens. Nature 403, 652–655. Dehaene, S. (2001). Précis of the number sense. Mind Lang. 16, 16–36.

Dehaene, S., and Changeux, J.-P. (1993). Development of elementary numerical abilities: a neuronal model. J. Cogn. Neurosci. 5, 390–407.

Dijkstra, K., van de Loosdrecht, J., Schomaker, L.R.B., and Wiering, M.A. (2018). CentroidNet: A deep neural network for joint object localization and counting. Conference: The European Conference on Machine Learning and Principles and Practice of Knowledge Discovery in Databases.

Douglass, J.K., and Strausfeld, N.J. (2003). Retinotopic pathways providing motion-selective information to the lobula from peripheral elementary motion-detecting circuits. J. Comp. Neurol. 457, 326–344.

Eichler, K., Li, F., Litwin-Kumar, A., Park, Y., Andrade, I., Schneider-Mizell, C.M., Saumweber, T., Huser, A., Eschbach, C., Gerber, B., et al. (2017). The complete connectome of a learning and memory centre in an insect brain. Nature 548, 175–182.

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Erber, J., Masuhr, T., and Menzel, R. (1980). Localization of short-term memory in the brain of the bee, *Apis mellifera*. Physiol. Entomol. *5*, 343–358.

Feigenson, L., Dehaene, S., and Spelke, E. (2004). Core systems of number. Trends Cogn. Sci. *8*, 307–314.

Gallistel, C.R., and Gelman, R. (2000). Non-verbal numerical cognition: from reals to integers. Trends Cogn. Sci. 4, 59–65.

Gegenfurtner, K.R. (2016). The interaction between vision and eye movements. Perception 45, 1333–1357.

Giurfa, M., Zhang, S., Jenett, A., Menzel, R., and Srinivasan, M.V. (2001). The concepts of "sameness" and "difference" in an insect. Nature 410, 930–933.

Gross, H.J., Pahl, M., Si, A., Zhu, H., Tautz, J., and Zhang, S. (2009). Number-based visual generalisation in the honeybee. PLoS One 4, e4263.

Grünewald, B. (1999). Morphology of feedback neurons in the mushroom body of the honeybee, *Apis mellifera*. J. Comp. Neurol. 404, 114–126.

Guiraud, M., Roper, M., and Chittka, L. (2018). High-speed videography reveals how honeybees can turn a spatial concept learning task into a simple discrimination task by stereotyped flight movements and sequential inspection of pattern elements. Front. Psychol. 9, 1347.

Haag, J., and Borst, A. (2001). Recurrent network interactions underlying flow-field selectivity of visual interneurons. J. Neurosci. 21, 5685–5692.

Hertel, H. (1980). Chromatic properties of identified interneurons in the optic lobes of the bee. J. Comp. Physiol. 137, 215–231.

Howard, S.R., Avarguès-Weber, A., Garcia, J.E., Greentree, A.D., and Dyer, A.G. (2018). Numerical ordering of zero in honey bees. Science 360, 1124–1126.

Ings, T.C., Wang, M.-Y., and Chittka, L. (2012). Colour-independent shape recognition of cryptic predators by bumblebees. Behav. Ecol. Sociobiol. *66*, 487–496.

Jaeger, H. (2002). Short Term Memory in Echo State Networks (GMD - German National Research Institute for Computer Science), GMD-Report 152.

Kotrschal, A., Rogell, B., Bundsen, A., Svensson, B., Zajitschek, S., Brännström, I., Immler, S., Maklakov, A.A., and Kolm, N. (2013). Artificial selection on relative brain size in the guppy reveals costs and benefits of evolving a larger brain. Curr. Biol. 23, 168–171.

Lempitsky, V. and Zisserman, A. (2010). Learning to count objects in images. NIPS10 Proc. 23rd Int. Conf. Neural Inf. Process. Syst. 1, 1324–1332.

Lihoreau, M., Raine, N.E., Reynolds, A.M., Stelzer, R.J., Lim, K.S., Smith, A.D., Osborne, J.L., and Chittka, L. (2012). Radar tracking and motionsensitive cameras on flowers reveal the development of pollinator multi-destination routes over large spatial scales. PLoS Biol. *10*, e1001392.

Liu, G., Seiler, H., Wen, A., Zars, T., Ito, K., Wolf, R., Heisenberg, M., and Liu, L. (2006). Distinct memory traces for two visual features in the *Drosophila* brain. Nature 439, 551–556.

Loewenstein, Y., and Sompolinsky, H. (2003). Temporal integration by calcium dynamics in a model neuron. Nat. Neurosci. *6*, 961–967.

Matsuzawa, T. (2009). Symbolic representation of number in chimpanzees. Curr. Opin. Neurobiol. *19*, 92–98.

Menzel, R. (1999). Memory dynamics in the honeybee. J. Comp. Physiol. A 185, 323–340.

Menzel, R., Fuchs, J., Nadler, L., Weiss, B., Kumbischinski, N., Adebiyi, D., Hartfil, S., and Greggers, U. (2010). Dominance of the odometer over serial landmark learning in honeybee navigation. Naturwissenschaften *97*, 763–767.

Nieder, A. (2018). Evolution of cognitive and neural solutions enabling numerosity judgements: lessons from primates and corvids. Philos. Trans. R. Soc. Lond. B Biol. Sci. 373, 20160514.

Nityananda, V., Skorupski, P., and Chittka, L. (2014). Can bees see at a glance? J. Exp. Biol. *217* (Pt 11), 1933–1939.

Pahl, M., Si, A., and Zhang, S. (2013). Numerical cognition in bees and other insects. Front. Psychol. *4*, 162.

Paulk, A.C., Dacks, A.M., and Gronenberg, W. (2009a). Color processing in the medulla of the bumblebee (Apidae: *Bombus impatiens*). J. Comp. Neurol. *513*, 441–456.

Paulk, A.C., Dacks, A.M., Phillips-Portillo, J., Fellous, J.-M., and Gronenberg, W. (2009b). Visual processing in the central bee brain. J. Neurosci. *29*, 9987–9999.

Paulk, A.C., Phillips-Portillo, J., Dacks, A.M., Fellous, J.-M., and Gronenberg, W. (2008). The processing of color, motion, and stimulus timing are anatomically segregated in the bumblebee brain. J. Neurosci. 28, 6319–6332. Pepperberg, I.M. (2006). Grey parrot numerical competence: a review. Anim. Cogn. 9, 377–391.

Perry, C.J., and Barron, A.B. (2013). Honey bees selectively avoid difficult choices. Proc. Natl. Acad. Sci. U S A *110*, 19155–19159.

Rahnemoonfar, M., and Sheppard, C. (2017). Deep count: fruit counting based on deep simulated learning. Sensors 17, 905.

Raine, N.E., and Chittka, L. (2011). Flower constancy and memory dynamics in bumblebees (Hymenoptera: Apidae: *Bombus*). Entomol. Gen. *29*, 179–199.

Rose, G.J. (2018). The numerical abilities of anurans and their neural correlates: insights from neuroethological studies of acoustic communication. Philos. Trans. R. Soc. Lond. B Biol. Sci. 373, 20160512.

Rugani, R. (2018). Towards numerical cognition's origin: insights from day-old domestic chicks. Philos. Trans. R. Soc. Lond. B Biol. Sci. 373, 20160509.

Skorupski, P., MaBouDi, H., Galpayage Dona, H.S., and Chittka, L. (2018). Counting insects. Philos. Trans. R. Soc. Lond. B Biol. Sci. *373*, 20160513.

Srinivasan, M., Zhang, S., and Bidwell, N. (1997). Visually mediated odometry in honeybees. J. Exp. Biol. 200, 2513–2522.

Stoianov, I., and Zorzi, M. (2012). Emergence of a "visual number sense" in hierarchical generative models. Nat. Neurosci. *15*, 194–196.

Stone, T., Webb, B., Adden, A., Weddig, N.B., Honkanen, A., Templin, R., Wcislo, W., Scimeca, L., Warrant, E., and Heinze, S. (2017). An anatomically constrained model for path integration in the bee brain. Curr. Biol. 27, 3069– 3085.

Wittlinger, M., Wehner, R., and Wolf, H. (2007). The desert ant odometer: a stride integrator that accounts for stride length and walking speed. J. Exp. Biol. *210*, 198–207.

Wittlinger, M., Wehner, R., and Wolf, H. (2006). The ant odometer: stepping on stilts and stumps. Science 312, 1965–1967.

Yang, E.-C., and Maddess, T. (1997). Orientationsensitive neurons in the brain of the honey bee (*Apis mellifera*). J. Insect Physiol. 43, 329–336.

Zhang, S., Bock, F., Si, A., Tautz, J., and Srinivasan, M.V. (2005). Visual working memory in decision making by honey bees. Proc. Natl. Acad. Sci. U S A *102*, 5250–5255. ISCI, Volume 11

Supplemental Information

Insect-Inspired Sequential Inspection Strategy Enables an Artificial Network of Four Neurons to Estimate Numerosity Vera Vasas and Lars Chittka

Transparent Methods

CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Vera Vasas (v.vasas@qmul.ac.uk or vvasas@gmail.com).

METHOD DETAILS

Generating the visual input

The relative amount of light (quantum catch) absorbed by the bee eye was set to 1 for white areas, 0 for black areas, and 0.5 for grey background areas. For the yellow and purple shown on Figure 2, the quantum catch for each receptor type can be calculated as (Backhaus and Menzel, 1987; Chittka and Kevan, 2005)

$$P = \int_{300}^{700} I_{s}(\lambda) S(\lambda) D(\lambda) d\lambda$$

where $I_s(\lambda)$ is the spectral reflectance function of the stimulus (Figure S2), $S(\lambda)$ is the spectral sensitivity function of the receptors from Peitsch et al. (1992), and $D(\lambda)$ is the spectrum of illumination, assumed to be unit across the visible spectrum but missing UV (matching experimental details for Skorupski et al., 2018). We used the response of the long-wavelength-sensitive receptors as the total quantum catch. (Colour and motion are processed separately in the bee brain (Paulk et al., 2008), and the latter only receives input from long-wavelength-sensitive receptors. Our model assumes input from small field on-off neurons of the motion pathway (as in e.g. Arenz et al., 2017). The quantum catch was scaled to yield 1 for white.

To calculate the visual input at each time step, we used a visual angle of 60 degrees and a viewing distance of 2 cm. We generated a partial image representing the field of view centred on a point on the flight path. The path used was the track specified on Figure 2 for the counting task (Skorupski et al., 2018) and idealized tracks (Data S1) for the numerical ordering task (Howard et al., 2018). Finally, the 'brightness' neuron's responses at each time step were calculated as taking the absolute differences between the quantum catches at the current and

the previous position pixel-by-pixel, and summing them, mimicking a summation of a collection of small field on-off visual neurons.

Neural model

In our minimal model, we used abstract linear transfer neurons, whose responses are bounded by 0 and 1. Thus, the output rates of neurons r_i at time *t* are given by the following piecewise linear transfer function:

if
$$x_i^t < 0$$
, $r_i^t = 0$;
if $0 \le x_i^t \le 1$, $r_i^t = x_i^t$;
if $1 < x_i^t r_i^t = 1$;

where x_i^t , the presynaptic input to neuron *i* at time *t* is calculated as the linear summation of input from presynaptic neurons *j*:

$$x_i^t = \sum_j w_{ji} r_j^{t-1}$$

where w_{ji} is the weight of the synapse from neuron *j* to *i*.

We used the following weight parameters for modelling the counting task from Skorupski et al. (2018) and the numerical ordering from Howard et al. (2018) (i – 'brightness' input neuron, b – 'brightness working memory' neuron, c – 'counting working memory' neuron, e – 'evaluation' neuron): $w_{ib} = 1.2$; $w_{ic} = 0.075$; $w_{bb} = 0.99$; $w_{cc} = 0.999$; $w_{be} = 1$; $w_{ce} = -1.1$.

DATA AND SOFTWARE AVAILABILITY

Description: Data S1

We implemented the calculations in MATLAB (Release 2018A, The MathWorks, Inc., Natick, Massachusetts, United States). Data S1 includes the scripts, the example stimuli and the flight paths from Howard et al. (2018) and Skorupski et al. (2018). Instructions for using the scripts are given as comments within each script.

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Software and Algorithms		
MATLAB	The MathWorks, Inc.	R2018A
Custom scripts for modelling the minimal neural	This paper	Data S1
network, and the associated input files: the stimuli		
and the flight paths from Howard et al. (2018) and		
Skorupski et al. (2018)		



Figure S1. The evaluation provided by the model follows the same pattern, that shown in Figure 3A, when the input is either a global brightness detector or an edge detector. Related to Figure 3. (A) We calculated the global brightness neurons' response using the total amount of quantum catch from its 60-degree visual field, subtracting the value at the previous time step from that at the given time step. The result was scaled log-linearly. The global brightness detector will thus respond to an increase in brightness with a phasic burst that scales log-linearly with the extent of the change (Hertel, 1980). We calculated the evaluation given by the model for the stimuli from Howard et al. (2018) using the following parameters: $w_{ib} = 0.8$; $w_{ic} = 0.09$; $w_{bb} = 0.99$; $w_{cc} = 0.999$; $w_{be} = 1$; $w_{ce} = -1.1$. (B) We defined the response for the edge detector neurons as the fraction of its 60-degree visual field that contains edges (Yang and Maddess, 1997). We calculated the evaluation given by the model for the stimuli from Skorupski et al. (2018) using the following parameters: $w_{ib} = 0.12$; $w_{bb} = 0.99$; $w_{cc} = 0.999$; $w_{cc} = -1.1$. The dots represent the response levels from the 'evaluation' neuron for each pattern; the black lines and the grey areas indicate the means and the standard deviations respectively.



Figure S2. The spectral reflectance functions of the colours used in Skorupski et al. (2018). *Related to Figure 2 and Methods.*

vs.	0 items	1 item	2 items	3 items	4 items	5 items	6 items
0 item	0.5	0.53	0.58	0.63	0.68	0.74	0.85
1 item	0.47	0.5	0.55	0.6	0.65	0.72	0.84
2 items	0.42	0.45	0.5	0.55	0.6	0.67	0.81
3 items	0.37	0.4	0.45	0.5	0.56	0.63	0.77
4 items	0.32	0.35	0.4	0.44	0.5	0.58	0.73
5 items	0.26	0.28	0.33	0.37	0.42	0.5	0.67
6 items	0.15	0.16	0.19	0.23	0.27	0.33	0.5

B.

VS.	0 items	1 item	2 items	3 items	4 items	5 items	6 items
0 item	0.5	0.36	0.24	0.2	0.17	0.15	0.12
1 item	0.64	0.5	0.37	0.31	0.27	0.24	0.2
2 items	0.76	0.63	0.5	0.44	0.39	0.35	0.3
3 items	0.8	0.69	0.56	0.5	0.45	0.4	0.36
4 items	0.83	0.73	0.61	0.55	0.5	0.46	0.41
5 items	0.85	0.76	0.65	0.6	0.54	0.5	0.45
6 items	0.88	0.8	0.7	0.64	0.59	0.55	0.5

Table S1. The same model is capable of selecting 'less' or 'more' depending on the decision rule. Related to Figure 3. The numbers indicate the proportion of landings on the stimulus indicated in the rows vs. the stimulus in the columns. **Bold** letters highlight a preference $\geq 60\%$. (A) If the bee chooses patterns to scan randomly and lands with a likelihood directly proportional to the state of the 'evaluation neuron' at the end of the scan, then the bee will show a measureable preference for the stimulus containing fewer items. (B) If the bee chooses patterns to scan randomly, but lands with a likelihood inversely proportional to the state of the 'evaluation neuron' at the end of the scan, then the bee will show a measureable preference for the stimulus containing fewer items. (B) If the state of the 'evaluation neuron' at the end of the scan, then the bee will show a measureable preference for the stimulus containing fewer items.

A.

Supplemental References

- Arenz, A., Drews, M.S., Richter, F.G., Ammer, G., and Borst, A. (2017). The temporal tuning of the *Drosophila* motion detectors is determined by the dynamics of their input elements. Curr. Biol. 27, 929–944.
- Backhaus, W., and Menzel, R. (1987). Color distance derived from a receptor model of color vision in the honeybee. Biol. Cybern. 55, 321–331.
- Chittka, L., and Kevan, P.G. (2005). Flower colour as advertisement. In Practical Pollination Biology, Dafni, A., Kevan, P.G., Husband, B.C.,eds. (Enviroquest Ltd.), pp. 157-196.
- Hertel, H. (1980). Chromatic properties of identified interneurons in the optic lobes of the bee. J. Comp. Physiol. 137, 215–231.
- Howard, S.R., Avarguès-Weber, A., Garcia, J.E., Greentree, and A.D., Dyer, A.G. (2018). Numerical ordering of zero in honey bees. Science 360, 1124–1126.
- Paulk, A.C., Phillips-Portillo, J., Dacks, A.M., Fellous, J.-M., and Gronenberg, W. (2008). The processing of color, motion, and stimulus timing are anatomically segregated in the bumblebee brain. J. Neurosci. 28, 6319–6332
- Peitsch, D., Fietz, A., Hertel, H., de Souza, J., Ventura, D.F., and Menzel, R. (1992). The spectral input systems of hymenopteran insects and their receptor-based colour vision. J. Comp. Physiol. A 170, 23–40.
- Skorupski, P., MaBouDi, H., Galpayage Dona, H.S., and Chittka, L. (2018). Counting insects. Phil. Trans. R. Soc. B 373, 20160513.
- Yang, E.-C., and Maddess, T. (1997). Orientation-sensitive neurons in the brain of the honey bee (*Apis mellifera*). J. Insect Physiol. 43, 329–336.