A learning-based approach to artificial sensory feedback leads to optimal integration

Maria C Dadarlat1–4, Joseph E O’Doherty1–3 & Philip N Sabes1–4

Proprioception—the sense of the body’s position in space—is important to natural movement planning and execution and will likewise be necessary for successful motor prostheses and brain–machine interfaces (BMIs). Here we demonstrate that monkeys were able to learn to use an initially unfamiliar multichannel intracortical microstimulation signal, which provided continuous information about hand position relative to an unseen target, to complete accurate reaches. Furthermore, monkeys combined this artificial signal with vision to form an optimal, minimum-variance estimate of relative hand position. These results demonstrate that a learning-based approach can be used to provide a rich artificial sensory feedback signal, suggesting a new strategy for restoring proprioception to patients using BMIs, as well as a powerful new tool for studying the adaptive mechanisms of sensory integration.

Humans plan and execute movements under the guidance of both vision and proprioception1–2. In particular, maximally precise movements are achieved by combining estimates of limb or target position from multiple sensory modalities, weighting each by its relative reliability3–6. Furthermore, in the absence of proprioception, even simple multi-joint movements become uncoordinated7,8. Therefore, we should not expect continuous process of adaptive recalibration even in adult humans. Therefore, we should not expect multi-sensory integration, whereby multiple sensory signals are combined, to improve the precision of sensory estimates, is learned from cross-modal experience during development15,16 and relies on a continuous process of adaptive recalibration even in adult humans and monkeys17–19. Recent theoretical work from our laboratory suggests that multisensory integration can be learned with experience through a simple Hebbian-like learning rule20. In this model, successful integration of two sensory signals depends not so much on choosing the right patterns of neural activity to encode spatial information but rather on the presence of spatiotemporal correlations between input signals, which allow downstream neurons to learn the common underlying cause—for example, hand position.

As a demonstration of the potential value of somatosensory feedback, it has been shown that including natural kinesthetic feedback improves BMI control in intact monkeys to near natural levels9. The ideal artificial proprioceptive signal would be able to fulfill the same functions that proprioception has in natural motor control: providing sufficient information to allow competent performance in the absence of other sensory inputs and permitting multisensory integration with vision to reduce movement variability when both signals are available. Here we present proof of concept that both of these goals can be achieved using multichannel intracortical microstimulation (ICMS).

Most efforts to develop artificial sensory signals have taken a biomimetic approach: trying to recreate the patterns of neural activity that underlie natural somatosensation10–14. We propose a complementary approach that focuses not on reproducing natural patterns of activity but instead on taking advantage of the natural mechanisms of sensorimotor learning and plasticity. In particular, the process of multisensory integration, whereby multiple sensory signals are combined, to improve the precision of sensory estimates, is learned from cross-modal experience during development15,16 and relies on a continuous process of adaptive recalibration even in adult humans and monkeys17–19. Recent theoretical work from our laboratory suggests that multisensory integration can be learned with experience through a simple Hebbian-like learning rule20. In this model, successful integration of two sensory signals depends not so much on choosing the right patterns of neural activity to encode spatial information but rather on the presence of spatiotemporal correlations between input signals, which allow downstream neurons to learn the common underlying cause—for example, hand position.

As a demonstration of the potential value of somatosensory feedback, it has been shown that including natural kinesthetic feedback improves BMI control in intact monkeys to near natural levels9. The ideal artificial proprioceptive signal would be able to fulfill the same functions that proprioception has in natural motor control: providing sufficient information to allow competent performance in the absence of other sensory inputs and permitting multisensory integration with vision to reduce movement variability when both signals are available. Here we present proof of concept that both of these goals can be achieved using multichannel intracortical microstimulation (ICMS).

Most efforts to develop artificial sensory signals have taken a biomimetic approach: trying to recreate the patterns of neural activity that underlie natural somatosensation10–14. We propose a complementary approach that focuses not on reproducing natural patterns of activity but instead on taking advantage of the natural mechanisms of sensorimotor learning and plasticity. In particular, the process of multisensory integration, whereby multiple sensory signals are combined, to improve the precision of sensory estimates, is learned from cross-modal experience during development15,16 and relies on a continuous process of adaptive recalibration even in adult humans and monkeys17–19. Recent theoretical work from our laboratory suggests that multisensory integration can be learned with experience through a simple Hebbian-like learning rule20. In this model, successful integration of two sensory signals depends not so much on choosing the right patterns of neural activity to encode spatial information but rather on the presence of spatiotemporal correlations between input signals, which allow downstream neurons to learn the common underlying cause—for example, hand position.

RESULTS

Behavioral task and feedback signals

Two rhesus macaques were trained to make instructed-delay center-out reaches to invisible targets (Fig. 1a) in a virtual reality environment (Supplementary Fig. 1) guided by feedback that represented the vector (distance and direction) from the middle fingertip to the reach target (Fig. 1b). This ‘movement vector’ was not explicitly shown; instead, it was encoded by one of three feedback types: a visual signal (VIS), a signal delivered through patterned multichannel ICMS pulse trains (ICMS) or a combination of these two signals (VIS+ICMS).

This task was chosen to test whether the ICMS signal could provide position information that could both be integrated with vision and could replace it. By using natural movement, we obtained the most direct and precise estimates of how well the ICMS signal encoded sensory information about the limb (for example, not...
confounded by additional performance noise due to imperfect BMI control). However, natural movement leaves natural proprioception intact, which would have made an ICMS signal encoding absolute limb position redundant. By encoding the relative positions of the limb and target, the VIS and ICMS signals provided a feedback variable that was both required to complete the task and not available from natural sensory signals.

The VIS signal was a random moving-dot flow field (‘dot-field’), wherein the direction and speed of the flow indicated the direction and distance to the target, respectively (Fig. 1b and Supplementary Video 1). The reliability, or precision, of the dot-field was manipulated by changing its coherence—the percentage of dots moving in the same direction—during reach planning and execution. Performance on VIS-only conditions. Behavioral performance measures, averaged across the last seven testing sessions for each monkey, are shown for each sensory feedback type and (for VIS and VIS+ICMS trials) as a function of visual coherence. Error bars denote bootstrapped s.e.m. The ICMS-only data points, which are independent of visual coherence, are extended across the plot to aid visual comparison. (a) Number of movement subsegments. (b) Movement path length, normalized by the initial distance to the reach target. (c) Movement time, normalized as in b. See Online Methods for a detailed description of task performance measures.
trials improved monotonically with increasing dot-field coherence for all behavioral metrics (Fig. 2), demonstrating that differences in performance reflect the precision of the sensory cue.

After the monkeys could perform the reaching task using visual feedback, we tested our hypothesis that spatiotemporal correlations between vision and ICMS could drive integration of the new sensory modality. We did so by exposing the monkeys to paired, correlated VIS+ICMS feedback signals both during the instructed delay period (as static information) and throughout the reach (dynamically updated feedback). The visual signal was first set to 100% coherence but was gradually reduced across training blocks to increase the relative value of the ICMS signal (ultimately settling at 25% for monkey D and 20% for monkey F). Under this training regime, the animals learned to integrate the two sensory signals; that is, the addition of ICMS improved behavioral performance (see below). Animals needed more explicit instruction to learn to initiate movement on ICMS-only trials (see Online Methods). Once that was accomplished, the training regime changed to include ICMS-only trials (33%), a pragmatic choice intended to speed learning. A summary of the behavioral training regime can be found in Supplementary Tables 1 and 2. We periodically assessed learning (approximately every 500–1,000 training trials) by including testing blocks: trials of VIS-only, VIS+ICMS and ICMS-only trials wherein the visual dot-field coherence could take a range of values ([0,15,25,50,100] for monkey D; [0,10,15,25,50] for monkey F).

Figure 3 Evolution of performance over training (monkey F). Behavioral performance measures are shown as a function of the cumulative number of VIS+ICMS trials performed (training and testing). The data, collected during testing sessions, were smoothed for clarity (Gaussian window with s.d. of 2,800 training trials for monkey F). The visual coherence on training trials was decreased across training sessions (indicated by gray bars at the bottom of the figure and vertical gray lines at the transitions). The left, thin green line denotes the onset of ICMS-only trials, where target sizes were temporarily larger than in the other trial conditions; the right, thick green line denotes the beginning of ICMS trials with targets of standard size. (a) Percentage correct trials. (b) Number of movement segments, including the initial movement and corrective submovements. (c) Movement time for the trial, normalized by the initial distance to the reach target. (d) Path length, normalized as in c. See Supplementary Table 2 for details of the training and testing schedule.

Figure 4 Monkeys estimate both target distance and direction from sensory feedback. Data points represent individual trials. Black solid lines are unity and the thick colored lines represent linear fits between the movement and target variable. Fits were performed separately for the distal (target angle [−π:0]) and proximal (target angle [0:π]) halves of the workspace. (a,b) Initial movement angle versus target angle for monkeys D and F, respectively. (c,d) Initial movement distance versus target distance for monkeys D and F, respectively. Region within the dashed black lines falls within the diameter of the target.
Substitution of vision by ICMS

We analyzed the data from testing blocks to determine how well the animals could interpret the ICMS signal, using it in place of vision to perform accurate reaches. Once the monkeys began making reaches on ICMS-only trials, they became increasingly proficient across training sessions (Fig. 3 and Supplementary Fig. 3), and performance on ICMS trials was ultimately comparable to performance with low to intermediate visual coherences (15–25% for monkey D, 15% for monkey F; Fig. 2 and Supplementary Tables 3 and 4). A more qualitative impression of the performance comparisons can be obtained from sample movement paths for various feedback conditions (Supplementary Fig. 4).

We next quantified how well the monkeys made use of the distance and direction information encoded in the ICMS signal. To do this, we analyzed the distance and direction of the initial movement segment of the reach, which reflected the animals’ estimates of the required movement vectors derived from sensory feedback during the instructed delay period (Fig. 4). Although demonstrating some idiosyncratic biases (see below), monkey D was also able to derive good estimates of target direction from ICMS (ICMS, $R^2 = 0.900$; Fig. 4a; 100% VIS, $R^2 = 0.957$; Fig. 4a). Monkey F was highly adept at estimating target angle (ICMS, regression $R^2 = 0.948$; Fig. 4b), performing as well with ICMS as with the highest visual coherences (50% VIS, $R^2 = 0.945$; Fig. 4b). Both monkeys were somewhat worse at estimating distance than direction from ICMS. Owing to differences in performance across the workspace, we analyze distance estimation separately for the two half-planes: the more proximal workspace, with target angles [$−\pi$, 0], and the more distal workspace, with target angles [0, $\pi$]. Monkey D could accurately estimate the distal half of the workspace ([0, $\pi$], $R^2 = 0.494$; Fig. 4c) but was less able to do so in the proximal half ([−$\pi$, 0], $R^2 = 0.108$; Fig. 4c). Still, these values are comparable to those the animal achieved with the highest coherence VIS feedback ([0, $\pi$], $R^2 = 0.365$; [−$\pi$, 0], $R^2 = 0.176$; Fig. 4c). For monkey F, distance estimates were equally good across the workspace ([−$\pi$, 0], $R^2 = 0.432$; [0, $\pi$], $R^2 = 0.473$; Fig. 4d) and largely fell within one target radius of the correct distance (but not necessarily in the target on each trial, because of directional error), although these values are lower than those he achieved with high-coherence VIS feedback ([−$\pi$, 0], $R^2 = 0.716$; [0, $\pi$], $R^2 = 0.751$; Fig. 4d). In summary, the task performance observed in the ICMS-only condition was driven by the animals’ ability to decode both distance and direction information from the ICMS signal.

Finally, we analyzed corrective submovements to determine whether the monkeys used ICMS as a continuous feedback signal during movement. As an alternative, it is possible that the animals only used the ICMS signal to estimate the target location during the instructed delay period, with subsequent corrective submovements guided either by the remembered target location or simply by a random search. However, we found that submovement direction correlated well with the direction of the online movement vector (target – hand position) at the end of the previous submovement (Fig. 5). In fact, the precision of the corrective movements in the ICMS-only condition was comparable to that seen with high visual coherences and was considerably lower than that observed with 0% VIS, where no directional information is available, or that which would be expected by chance. These results suggest that the VIS and ICMS cues were indeed being used as online feedback signals. Furthermore, if the animals had simply memorized the location of the target during the instructed delay period, we would expect a decline in precision across sequential corrective movements. Instead, the correlation between cued and executed submovements was largely consistent across corrective submovement number, with no clear increase in error variance for later submovements. These results strongly indicate that the animals were using the online feedback to execute corrective submovements.

Minimum-variance integration of vision and ICMS

In addition to serving as the training condition, VIS+ICMS trials provided a test of the animals’ ability to improve performance by combining information from the two sensory cues. This ability emerged during training, with performance on VIS+ICMS trials becoming progressively better than for VIS trials, even before the animals could complete reaches with ICMS alone (Fig. 3 and Supplementary Fig. 3), supporting the idea that multisensory integration drives ICMS learning. Moreover, the hallmarks of multisensory integration were observed in the asymptotic performance on VIS+ICMS trials, after learning was complete (Fig. 2). At intermediate dot-field coherences (10–25%), where performance between the two unimodal cues was similar, VIS+ICMS reaches were significantly better; that is, faster and straighter (Supplementary Table 3). In contrast, at high (50–100%) and low (0%) dot-field coherences, behavior in the bimodal condition approximated that observed with the more reliable of the unimodal cues (Supplementary Table 3).
We next asked whether the visual and ICMS cues were integrated in an optimal—that is, minimum variance—manner, as is the case for natural visual and somatosensory signals. The answer came from an analysis of the statistics of the initial reach directions, which gave the most direct readout of the animals’ estimate of target direction following the instructed delay period. The minimum variance model makes specific predictions about both the variance and bias of this estimate, and we consider each in turn.

We first considered how the variance of the initial reach angle depends on feedback condition (Fig. 6a). For VIS trials, the initial angle variance increased markedly with decreases in coherence, as expected if the variance reflects the residual uncertainty about cue direction after the instructed delay. Variability in the ICMS trial was comparable to that in VIS trials at 15–25% visual coherence, consistent with the other movement metrics above. From these unimodal variances, we determined what the initial angle variance should be for the VIS+ICMS condition, under the model of minimum variance integration. We computed this prediction under two limiting conditions: assuming that the initial angle variance arises only from variability in the sensory estimates of target direction (Fig. 6a) or assuming that the measurements also include the maximal consistent level of downstream (for example, motor) variability (see Online Methods; Fig. 6a). The empirical variances observed in the VIS+ICMS condition followed the predicted trend closely, lying between the two limiting predictions in the region where the animal received most of the multisensory training (20–25% coherence; see Online Methods). This comparison suggested that, after training, the animals optimally combined the ICMS signal with vision. We next tested this conclusion further by analyzing the pattern of mean initial angles.

The animals exhibited idiosyncratic patterns of mean initial reach angle as a function of target angle. For monkey D, these patterns were clearly distinct between the VIS and ICMS trials (Fig. 6b; also see Supplementary Fig. 5 for monkey F, whose patterns were less well defined). Since the required movements were the same across cue conditions, these patterns likely arise from biased estimation of the target direction. Therefore, they offered another opportunity to test whether the VIS and ICMS signals were combined optimally. Minimum variance integration predicts that in the VIS+ICMS condition, as the visual coherence increases from 0% to 100%, the animals should transition from relying primarily on the ICMS cue to primarily on the visual cue. This trend could be seen qualitatively in the pattern of mean initial angles for monkey D (Fig. 6b): at 15% coherence, the VIS+ICMS mean was close to that observed with ICMS alone; at 100% it was close to that observed with VIS. The relative weighting of the two modalities were estimated quantitatively by modeling the VIS+ICMS mean as an affine combination of the unimodal biases (Online Methods, equation (3)). As expected, the weighting of the visual cue smoothly transitioned from zero to unity as the visual coherence increased (Fig. 6c). Under the model of minimum variance integration, each sensory cue should be weighted inversely proportional to its variance (Online Methods, equation (6)). Using the unimodal variance data from Figure 6a, we obtained quantitative predictions for the cue weighting in the VIS+ICMS trials (Fig. 6c; Online Methods, equations (7) and (8)); these were in good agreement with the empirical data.

**DISCUSSION**

We have shown that multichannel patterned ICMS of primary somatosensory cortex can be used to provide monkeys with continuous information about hand position that enables goal-directed reaching. In particular, the monkeys were able to use ICMS to estimate the distance and direction between their hand location and the reach target. Furthermore, when both visual and ICMS feedback was available, the...
monkeys combined these signals to achieve better task performance, and they did so at or near theoretical optimal levels, as is observed for natural sensory signals.4,5

A key result of the paper is that, with only eight electrodes, we were able to deliver continuous spatial information with a reliability comparable to that achieved with the visual cue. Initial angle estimation with the ICMS signal had the same variance as that observed with 15–20% visual coherence and was only about three times more variable than that observed with the highest coherence for both animals. Furthermore, when the signals were used online, ICMS performance was even closer to that achieved with visual feedback (Fig. 5b). The better performance (relative to vision) of ICMS during corrective movements could be due to either the inherent delays in visual feedback, the relative importance of somatosensory feedback for online movement controls, the shorter integration times available for online corrections, or a greater contribution of motor noise.

Learning to integrate a novel sensory signal
The ability to use the ICMS signal was necessarily learned during the training process, since, by design, the patterns of ICMS did not mimic naturally occurring signals in the brain. This learning could have been driven by several possible mechanisms of learning. Consider that, in this experiment, the visual and ICMS signals changed in a correlated fashion. Previous modeling work from our lab showed that in a network with unsupervised, Hebbian-like learning, such correlations are sufficient to learn optimal integration20; however, other learning mechanisms may also have contributed, including error-corrective or reinforcement learning24,25 of a sensory-to-motor mapping from ICMS to the appropriate movement. Our experimental design cannot definitively distinguish between these possibilities, but the emergence of multisensory integration before the animals could perform with ICMS alone (Fig. 3 and Supplementary Fig. 3) suggests that unsupervised, multisensory learning played a large, if not dominant, role.

While there is evidence of multisensory integration at all visual coherences (Fig. 2), optimal performance in the VIS+ICMS condition was only observed at intermediate visual coherences; performance at the lowest and highest coherences approached optimality, but did not achieve it (Fig. 6). A likely explanation is that monkeys learned to integrate vision and ICMS optimally when vision matched or was close to the training coherence (25% for monkey D; 20% for monkey F). In fact, a similar effect was observed in the network model of unsupervised sensory integration20: small but apparent departures from optimal integration were seen when the unimodal variances deviated too far from the regime in which the network was trained. Those observations are qualitatively consistent with the experimental results we presented here.

ICMS feedback as a tool for studying multisensory circuits
The finding that animals can integrate and use an ICMS signal to direct movement offers a potentially powerful tool for studying information processing in sensorimotor circuits. For example, the posterior parietal cortex uses sensory feedback for a variety of multisensory computations, including estimation of the position of the limb and the location of targets.26–30 Computational models have been developed to demonstrate how neural circuits could perform these operations20,31,32, but testing these models has proven difficult. The challenge stems, in large part, from the fact that brain areas within the PPC exhibit complex, heterogeneous and partially redundant spatial representations33–35 and interact in a complex network22,36–38, often with overlapping function40,41. It may not be possible to discover how information is processed in this complex circuit only by manipulating the distal sensory inputs. ICMS feedback, by contrast, affords the experimenter proximal control of the afferent signal. This has two pertinent advantages. First, the anatomical origin of the signal can be controlled. In this particular study, we stimulated areas of somatosensory cortex that we know project directly to multisensory areas as such area 5 and ventral intraparietal cortex42,43; however, stimulation could be performed in other brain areas that participate more strongly in other cortical circuits, for example, in area 7 to study spatial representations in the circuits for saccadic eye movements. Second, because the ICMS signal bypasses peripheral receptors and subcortical processing, it gives the experimenter finer control over the statistics of the signal. Signal statistics figure prominently in current models of multisensory neuronal processing20,32, but the signal manipulations that would be most diagnostic for these models would be challenging or impossible to achieve with natural stimulation. With ICMS feedback, the manipulations become tractable: for example, changing the correlational structure between neurons in a given area or between neurons representing two sensory signals, and gaining full control over the timing and context of subjects’ exposure to the signal.

Furthermore, we think that the learning observed in the present study taps into the same mechanisms of plasticity that drives other forms of multisensory learning, such as intersensory calibration—the ability of two sensory modalities to come back into alignment following a perturbation17,19,39,44. ICMS feedback offers an ideal tool for studying these mechanisms, for example, for testing models of learning20 in real neural circuits, for the reasons described above—local access and control of statistics—and because we can observe changes in the behavioral and electrophysiological state of the animal from the very first exposure to the novel signal.

Neuroprosthetic applications
In this study, we show that ICMS can be used to deliver a task-relevant feedback signal that guides online, multidimensional movement control. We chose to encode the position of the hand relative to the reach target, as opposed to an absolute proprioceptive signal, because this allowed us to study artificial sensation in a simple, natural task without having to suppress natural proprioception. Because the reach target was never visible, estimation of the relative hand position was required in order to perform the task. To use this approach to provide proprioceptive feedback from a prosthetic device, the ICMS signal would instead encode the state of the device with respect to the body, for example joint or endpoint position or velocity. Because these variables are also available via visual feedback, the same learning mechanisms should apply.

We expect that ICMS feedback could play the same role for BMI control that proprioceptive feedback does for normal movement control. During normal movement, vision and proprioception make comparable contributions to limb state estimation1–3,45. While proprioceptive loss does not have a substantial effect on the simplest reaches when visual feedback is available46, it does impair performance of movements that require inter-joint coordination, which include most activities of daily life7. We have shown that the eight-channel ICMS signal used here can provide online feedback with reliability that is also comparable to that of vision. We expect that increasing the number of electrodes and optimizing the encoding scheme will further improve the quality and information capacity of this signal.

Our learning-based approach can be contrasted with a biomimetic approach, the attempt to reproduce natural patterns of sensory-evoked neural activity. In practice, a truly biomimetic stimulation scheme is not attainable, owing to a range of technical and scientific considerations, such as the lack of access to the full neuronal population, inability to translate electrical stimulation into naturalistic neural activity.
patterns, an incomplete understanding of neural encoding mechanisms and difficulty in characterizing those patterns in patients with sensory loss. Our approach circumvents these issues by taking advantage of the inherent plasticity of the brain. However, completely disregarding prior knowledge about natural neural signals, such as somatotopic organization, will impair initial performance and may limit the rate or extent of learning. In fact, learning-based and biomimetic approaches are complementary, and systems optimized for clinical applications would likely benefit from taking a combined approach.

METHODS

Methods and any associated references are available in the online version of the paper.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

ACKNOWLEDGMENTS

We thank M.R. Fellows for initial behavioral training; R.R. Torres for suggesting the 2AFc task for ICMS detection; A. Leggitt for help with data analysis; K.R. Andrews and K. MacLeod for animal-related support; and J.G. Makin, A. Yazdan-Shahmorad and T.L. Hanson for discussion and comments on the manuscript. This research was supported by the Defense Advanced Research Projects Agency (DARPA) Reorganization and Plasticity to Accelerate Injury Recovery (REPAIR, N66001-10-C-0510) and the US National Institutes of Health NEI (EY015679, EY007120).

AUTHOR CONTRIBUTIONS

M.C.D. and P.N.S. designed the experiments; M.C.D. and J.E.O. developed and tested multistimulus simulation capabilities, including behavioral validation; M.C.D. performed the experiments; M.C.D. and P.N.S. analyzed the data; M.C.D., P.N.S. and J.E.O. wrote the manuscript.

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

Reprints and permissions information is available online at http://www.nature.com/reprints/index.html.
ONLINE METHODS

Subjects and implants. All animal procedures were performed in accordance with the US National Research Council’s Guide for the Care and Use of Laboratory Animals and were approved by the UCSF Institutional Animal Care and Use Committee. Two adult male rhesus macaque monkeys (Macaca mulatta) participated in this study. No statistical methods were used to predetermine this sample size. Rather, we chose to use two animals because the goal of this study was to demonstrate a novel form of learning, and we were able to show statistical significant effects of both learning and sensory integration independently in each animal. Clear demonstration of such learning effects in two macaque monkeys meets the generally employed standard in the field.

Each animal was chronically implanted with a 96-channel silicon microelectrode array coated with iridium oxide (Blackrock Microsystems, Salt Lake City, UT) over their left primary somatosensory cortices (Brodmann areas 1 and 2, S1). Most cells on monkey D’s array had receptive fields spanning the arm and shoulder (Supplementary Fig. 2), whereas those on monkey F’s array spanned the shoulder, back, side of the head, ear and occiput (Supplementary Fig. 2).

Behavioral task. The animals were trained to perform reaches in the horizontal plane to an unseen target in a two-dimensional virtual reality environment, where a mirror and an opaque barrier prevented direct vision of the arm (Supplementary Fig. 1). The mirror reflected visual input from a projector so that the visual cues appeared in the horizontal plane of the reaching hand. Fingertip position was monitored with an electromagnetic position sensor (Polhemus Liberty, Colchester, VT) at 240 Hz.

Each trial consisted of four epochs (Fig. 1a). (i) The monkeys moved the middle fingertip of their right hand to a fixed start position, located in the center of the screen and indicated by a circular visual target (10 mm radius). (ii) After a brief delay (0.25 and 0.5 s for monkeys D and F, respectively), the target cue was initiated, indicating the movement vector between the monkey’s current finger position and the center of the unseen reach target (12 mm radius). Targets were selected uniformly from an annulus centered on the start target with an inner radius of 40 mm and an outer radius of 115 mm (80 mm for monkey D). The movement vector cue was provided in the form of a dot-field (VIS, multichannel ICMS (ICMS) or both (VIS+ICMS). The monkeys were required to hold their position during this instructed-delay interval (0.2–0.7 s and 1–1.5 s for monkeys D and F, respectively). (iii) After a go cue (750 Hz tone, 0.5 s) the monkeys made a reach under the guidance of continuously updating VIS, ICMS or VIS+ICMS feedback. (iv) After acquiring the target and holding for 400 ms (monkey D) or 500 ms (monkey F), the monkeys received a liquid reward. Trials were terminated without reward if the monkeys moved too early during any of the delay intervals or if they failed to reach the target before a timeout (10 s). Different task parameters were selected for each animal to minimize the number of failed trials (for example, aborted hold at start or target) and therefore reflect the animals’ idiosyncratic behavioral tendencies.

Visual feedback. For vision, the movement vector was encoded using a random moving-dot flow field (‘dot-field’) consisting of approximately 600 dots over the visual display (roughly 53 cm × 33 cm, in the reaching plane). Each dot was initialized to a random location on screen and had a lifetime of 4 s (phases randomized), after which it reappeared at a new random location. Each dot in the dot-field moved at a speed proportional to the length of the movement vector, but could not exceed a maximum of 50 cm/s for monkey D and 40 cm/s for monkey F. A percentage of the dots moved coherently together in the direction of the continuously updating movement vector. The remaining dots moved in random directions, selected independently and uniformly from the circle. The percentage of dots moving coherently—the dot field coherence—was parametrically varied in order to manipulate the precision of the visual feedback.

ICMS. Intracortical microstimulation consisted of biphasic, charge-balanced pulse trains delivered asynchronously to each of eight electrodes in the array. The pulse trains were cathode-leading and symmetric, with 200 µs/phase and a 250-µs phase separation. The pulse amplitudes varied across electrodes, depending on perceptual threshold (see below) and ranged between 34 and 60 µA for monkey D and 30 and 80 µA for monkey F.

ICMS detection. A preliminary two-alternative forced choice task was used to determine the threshold pulse amplitudes at which the animals could detect ICMS on a given electrode. The monkeys first moved to a fixed start position near the midline (as in main task above) and maintained that position for 0.5 s. Next, there was a 0.5 s instructed delay period during which two reach targets were displayed, to the right and left of midline. The presence of an ICMS pulse train (100 Hz, 0.5 s) cued the animal to reach left; its absence cued the rightward reach. Animals were initially trained on this task using multielectrode stimulation, and the task was then used to identify electrodes on which ICMS was detectable. Eight such electrodes were identified for monkey F. For monkey D, seven such electrodes were identified; the final electrode could not be detected when stimulated alone with amplitudes of less than 60 µA.

Movement vector encoding. For ICMS, the movement vector was encoded in the spatial and temporal patterns of stimulation across the array (Fig. 1). Movement vector direction was encoded by the relative stimulation pulse rates across the electrodes. First, each of the eight electrodes was arbitrarily assigned one of eight preferred directions (PD), equally spaced around the circle. Then the stimulation pulse rate $f_i$ of electrode $i$ was calculated as a function of angle between the direction movement vector, $\theta$, and the electrode’s assigned PD:

$$f_i = f_0 \frac{1 + \cos(\theta - PD_i)}{2}, \quad i \in \{1:8\}$$

The frequency scaling factor, $f_0$, linearly encoded the movement vector distance, $d$, within the range of 100–300 Hz:

$$f_0 = \begin{cases} 300 & \text{if } d / d_{\text{max}} \geq 1 \\ 100 & \text{if } d / d_{\text{max}} < 1/3 \\ \frac{d}{d_{\text{max}}} & \text{otherwise} \end{cases}$$

where $d_{\text{max}}$ was 13.6 cm for monkey D and 10 cm for monkey F. The values of $d$ and $\theta$ were continuously updated during the reach to provide online feedback (Fig. 1d). The range of $f_0$ was restricted to the range [100, 300] to ensure that the monkeys could always detect the ICMS signal.

Experimental design. Behavioral sessions were divided into training blocks and testing blocks. The details of those blocks changed during the course of training, as described here and in Supplementary Tables 1 and 2.

Behavioral training. Monkeys first learned to perform the behavioral task using vision alone, with 100% coherence. Next, we began training with only VIS+ICMS trials. A dot-field coherence of 100% was used initially, and that value was slowly decreased across sessions to encourage the animals to use to the ICMS signal. Coherence values were lowered to 25% for monkey D and 20% for monkey F; these values depended on the animal’s performance in the VIS condition. This training regime was employed for approximately 20,000 training trials with monkey D and 40,000 training trials with monkey F, at which point the animals showed clear evidence of sensory integration of the VIS and ICMS signals—improved performance on VIS+ICMS trials compared to VIS trials, as evaluated on testing blocks (see below). We then altered the training regime to include 33% ICMS-only trials and 67% VIS+ICMS trials (see Supplementary Tables 1 and 2) once the animals were able to perform ICMS-only trials in the testing blocks (see below).

Behavioral testing. In between blocks of training, approximately every ~500–1,000 training trials, the animals performed a testing block to quantify performance across all feedback conditions. By the end of the experiment, a total of 11 feedback conditions were used: VIS, VIS+ICMS, and ICMS, with dot-field coherences of {0, 15, 25, 50, 100}% for monkey D and {0, 10, 15, 25, 50}% for monkey F, with the difference between animals reflecting individual performance in the VIS condition. At the beginning of the experiment, animals had not been exposed to lower visual coherences and could not perform the task at those coherences. Lower visual coherences for testing were introduced gradually during the course of the experiment as performance improved (see Fig. 3 and Supplementary Fig. 3). For all testing blocks, all conditions were randomized across trials.

As noted above, testing sessions revealed evidence of sensory integration of the VIS and ICMS signals before the animals began performing the ICMS-only task. At this stage, the animals still failed to initiate movement on ICMS-only blocks.
Data analysis. Behavioral performance measures. We quantified the animals’ ability to use the various sensory cues with the following performance measures (details of each are given below): (i) percentage correct trials, (ii) number of movement subsegments, (iii) normalized movement time, (iv) normalized path length and (v) mean and variance of initial angle of the movement. The first four metrics assessed animals’ use of the sensory cues throughout the trial, including movement planning during the instructed delay period and online movement control. The statistics of the initial angle assess only movement planning. The performance summaries in Figure 2 were computed from the last seven testing sessions available for each monkey.

(i) Percentage correct trials. This is the number of trials in which a monkey acquired a target and received a reward, compared to the total number trials in which a reach was initiated. We exclude in this analysis errors due to reaches beginning before the go cue, failure to initiate a trial, etc.

(ii) Number of movement subsegments. This metric quantifies the number of discrete submovements in a trial. Starting with the model assumption that submovements have bell-shaped velocity profiles47, we identified submovements by threshold crossings of the radial velocity plot of a trajectory, with a threshold of 20% of the maximum velocity on a given trial.

(iii) Normalized movement time. Since maximum movement velocity was largely independent of movement distance (data not shown), targets that were farther away took longer to reach. Therefore, we normalized the movement time by the distance from the starting point to the target.

(iv) Normalized path length. Similarly, we normalized the integrated path length by the distance from the starting point to the target.

(v) Mean and variance of the initial angle. For each monkey and feedback condition, we first computed a smoothed estimate of the mean initial angle as a function of target angle (robust locally weighted scatterplot smoothing, using the MATLAB “smooth” function, with a window of 40 data points). The initial angle variance was computed about this mean. Standard errors for the mean variance were estimated via bootstrapping.48

Quantifying use of direction and distance from the ICMS signal. A typical reach consisted of a long initial movement segment followed by one or more shorter, corrective submovements. The distance and direction of this initial reach can be taken to reflect the monkey’s estimate of target distance and direction, as decoded from the sensory information available during the instructed delay period.

(i) Direction estimation. We assessed the monkeys’ ability to estimate target direction from ICMS by regressing initial movement angle against target angle for ICMS-only trials. The initial movement angle was measured using the first movement subsegment, as described above. This assay ignores target-dependent biases in initial direction (see above) and is therefore a conservative estimate of the animal’s ability to decide target direction.

(ii) Distance estimation. We assessed the monkeys’ ability to estimate target distance from ICMS by regressing initial movement distance against target distance for ICMS-only trials. On a subset of trials, however, the initial movement deviated from the norm: animals sometimes made a small initial reach that was followed by several larger corrections. On these trials the distance of the initial reach segment was uncorrelated with movement distance. Therefore, for this analysis, we exclude trials for which the first movement segment was not the longest segment. This occurred in 30.2% of the trials for monkey D and 12.2% of the trials for monkey F.

Model prediction for initial angle variance. Under the model of minimum variance sensory integration8, we could predict the sensory variability in the bimodal condition from the variability for each unimodal condition. We focused on variability in the animals’ estimate of the target angle based on the sensory cues during the instructed delay period. Unimodal variances were computed from the variability in initial movement direction, \( \sigma^2_{\text{VIS}} \) and \( \sigma^2_{\text{ICMS}} \), calculated as described for performance measure (v) above. Next, for each coherence level, we predicted the bimodal variance under two limiting assumptions. First, we used the raw initial angle variance directly, which implicitly assumes that all movement variability derives from sensory variability:

\[
\sigma^2_{\text{VIS+ICMS}} = \frac{\sigma^2_{\text{VIS}} \sigma^2_{\text{ICMS}}}{\sigma^2_{\text{VIS}} + \sigma^2_{\text{ICMS}}} \tag{3}
\]

Second, we assumed that the movement variability was the largest possible value that would still be consistent with the data—this is the smallest initial angle variance across conditions, which we denote by \( \sigma^2_{\text{MIN}} \). Under this model, the bimodal initial angle variance is

\[
\sigma^2_{\text{VIS+ICMS}} = \frac{(\sigma^2_{\text{VIS}} - \sigma^2_{\text{MIN}})(\sigma^2_{\text{ICMS}} - \sigma^2_{\text{MIN}})}{\sigma^2_{\text{VIS}} + \sigma^2_{\text{ICMS}} - 2\sigma^2_{\text{MIN}}} + \sigma^2_{\text{MIN}} \tag{4}
\]

Model predictions for mean initial angle. The plots of mean initial angle for monkey D show a clear dependence on the feedback type (Fig. 6b). If we suppose that these differences reflect biases in the sensory estimates of target direction, then the minimum variance model can be used to predict, for each coherence value, the mean initial angle in the VIS+ICMS trials from those measured in the unimodal trials:

\[
\bar{\theta}_{\text{VIS+ICMS}} = \frac{\sigma^2_{\text{VIS}} \theta_{\text{VIS}} + \sigma^2_{\text{ICMS}} \theta_{\text{ICMS}}}{\sigma^2_{\text{VIS}} + \sigma^2_{\text{ICMS}}} \tag{5}
\]

Equation (5) can be summarized by the predicted visual cue weighting for each coherence,

\[
w_{\text{VIS}} = \frac{\sigma^2_{\text{VIS}}}{\sigma^2_{\text{VIS}} + \sigma^2_{\text{ICMS}}} \tag{6}
\]

which depends only on the unimodal initial angle variances.

We compared these model predictions to empirical values of the visual cue weighting, estimated from the mean initial angles. First, we divided the workspace into octants, and, for each octant and feedback condition, we computed the mean difference between the initial angle and the target angle, which we denote here as \( \bar{\theta}_{\text{octant},x} \) for octant \( x \) and condition \( x \). For each octant and coherence level, we then estimated the visual cue weighting as

\[
w_{\text{VIS}} = \frac{\delta_{\text{VIS+ICMS}} - \delta_{\text{ICMS}}}{\delta_{\text{VIS}} - \delta_{\text{ICMS}}} \tag{7}
\]

The standard error, \( s_{\text{VIS}} \), of each \( w_{\text{VIS}} \) was estimated from the standard errors of the component means, \( \delta_{\text{VIS}} \), by propagation of errors. Finally, for each coherence level we computed the mean visual cue weighting across octants, with each octant weighted by its standard error:

\[
w_{\text{VIS}} = \frac{\sum_{i=1}^{8} s_{\text{VIS}}^2 w_{\text{VIS}}}{\sum_{i=1}^{8} s_{\text{VIS}}^2} \tag{8}
\]

A Supplementary Methods checklist is available.

47. Novak, K.E., Miller, L.E. & Houk, J.C. The use of overlapping submovements in the control of rapid hand movements. *Exp. Brain Res.* 144, 351–364 (2002).

48. Efron, B. & Tibshirani, R.J. *An Introduction to the Bootstrap* (Chapman & Hall, 1993).