Cross-modal interactions are very common in perception. An important feature of many perceptual stimuli is their reward-predicting properties, the utilization of which is essential for adaptive behavior. What is unknown is whether reward associations in one sensory modality influence perception of stimuli in another modality. Here we show that auditory stimuli with high-reward associations increase the sensitivity of visual perception, even when sounds and reward associations are both irrelevant for the visual task. This increased sensitivity correlates with a change in stimulus representation in the visual cortex, indexed by increased multivariate decoding accuracy in simultaneously acquired functional MRI data. Univariate analysis showed that reward associations modulated responses in regions associated with multisensory processing in which the strength of modulation was a better predictor of the magnitude of the behavioral effect than the modulation in classical reward regions. Our findings demonstrate a value-driven cross-modal interaction that affects perception and stimulus encoding, with a resemblance to well-described modulatory effects of attention. We suggest that multisensory processing areas may mediate the transfer of value signals across senses.

reward value | sensory discrimination | audiovisual

The world is structured such that objects or events that cause sensations in one sensory modality influence those in another modality, a mechanism that underlies the near-ubiquitous phenomenon of multisensory interaction (1). This phenomenon has been the focus of a large and burgeoning theoretical and experimental literature (1–4). One feature of environmental stimuli important for adaptive behavior is their rewarding or reward-predicting properties. Surprisingly, we know little about how reward associations in one sensory modality influence processing in other modalities. In particular, whether an association with increased reward, known to increase the accuracy of perceptual processing within that sensory modality (5–10), can transfer between modalities for concurrently presented stimuli is unclear. This is important both because of what it reveals about the nature of cross-modal associations and because it might constitute an important perceptual mechanism in its own right. For example, one can easily imagine the importance of increased sensitivity to change in the visual scene to the concurrent sound of a tiger roaring or the mating call of a conspecific.

To study cross-modal transfer of value, we designed a visual orientation discrimination task in which visual stimuli were presented concurrently with one of two arbitrary pure tones previously associated with different levels of monetary reward. Critically, these tones were task-irrelevant and bore no relationship either to the orientation of the visual stimulus or to the outcome of the trial. (No feedback was presented about the accuracy of perceptual judgments, and performance on the orientation discrimination task was not related to the payment subjects received at the end of the trial.) We hypothesized that psychophysical measures of visual orientation judgment (d’ and performance accuracy) would show improvement in the presence of a sound with high (compared with low) reward association. Furthermore, simultaneous and spatially overlapping stimuli are more likely to have a common source and thus a shared reward association. Previous studies have shown that interactions between sensory modalities are strongest when stimuli are presented simultaneously and at contiguous spatial locations (1, 2, 11–13), especially at the earliest stages of processing (2, 11).

On this basis, we hypothesized that the effect of rewarded sounds on vision should be strongest when visual and auditory stimuli overlap in time and in space. Owing to the low temporal resolution of functional MRI (fMRI) signals, here we manipulated only the spatial overlap of stimuli while keeping their temporal alignment constant (with simultaneous presentation).

We had two specific questions regarding the neuronal underpinnings of any observed behavioral effects, which we addressed in simultaneously acquired fMRI data. First, we were interested in whether better behavioral performance owing to the presence of high-reward sounds was accompanied by a more differentiated stimulus representation in early visual areas, as assessed by classification accuracy in a multivariate pattern analysis (MVPA). Second, we were interested in whether this effect involved multimodal cortical areas known to play a role in the integration of audiovisual information, such as the superior temporal sulcus (STS) (14–16), or instead solely involved processing in classical reward-related areas, such as the ventromedial prefrontal cortex (vmPFC) (17, 18) or ventral striatum (19, 20).

Our first question draws on evidence demonstrating a nonuniform spatial distribution of reward effects across the visual cortex (21) and a sharpening of sensory representations through suppression of redundant signals elicited by reward/information predictive signals (10, 22). This evidence suggests that reward...
information might be reflected in the spatial pattern, rather than the absolute magnitude, of blood-oxygen-level–dependent (BOLD) responses. The second question is based on two possible mechanistic schemes of how reward impacts on perception, where reward could either directly impact neuronal (hence, visual) processing or via intermediate and presumably task-specific stages (here via regions involved in integrating auditory and visual information). Although both schemes predict a change in response magnitude or spatial pattern of BOLD responses in the visual cortex, only the latter predicts involvement of specific cross-modal areas. We provide evidence that supports task-specific influences of reward on perception.

Results

Behavioral Results. After subjects were familiarized with sounds and their respective rewards (Fig. 1A), they performed a visual orientation discrimination task in the presence of previously rewarded sounds (Fig. 1B and Materials and Methods). To test the effect of a sound’s reward association on visual orientation discrimination, we first examined the time course of the performance difference between Gabors + high-reward sounds and Gabors + low-reward sounds (Fig. 1C). High-rewarded sounds led to a better performance in orientation discrimination task, with the maximum effect seen at the beginning of the experiment and lower but steady levels in later trials. Average discrimination sensitivity, measured by $d'$ and accuracy, measured by percent correct rate, were higher for the Gabors accompanied by high-reward sounds (Fig. 1D). Repeated-measures ANOVA, with $d'$ as the dependent factor and reward and spatial congruency as the independent factor, revealed a significant main effect of reward ($F_{1,23} = 5.64, P = 0.02$). The interaction between reward and spatial congruence was not significant ($F_{1,23} = 1.15, P = 0.295$). Planned pairwise comparisons showed that both the high-reward–congruent (HC) and high-reward–incongruent (HIC) conditions had a significantly higher $d'$ value than the low-reward–congruent (LC) condition ($P = 0.01$). A comparison of HC vs. LC, $P = 0.044$ for comparison of HIC vs. LC; none of the other pairwise comparisons was significant, $P > 0.05$). We obtained similar results when percent correct rates were compared ($P = 0.03$ for comparison of HC vs. LC, paired $t$ test; all other pairwise comparisons were nonsignificant).

Fig. 1E shows the time course of the behavioral effect of sound and rewards inside the scanner. As time progressed, there was a marked decrease in the effect of reward, culminating in a reversal of the effect (i.e., lower performance for high-reward sounds) during the last few trials. This effect, in which the “extinction” of responses to a conditioned stimulus ultimately leads to a behavioral reversal after repeated exposure to nonreinforced conditioned stimulus, is well described in the conditioning literature (23). Extinction occurred only inside the scanner, most likely owing to either the differences between the scanning and behavioral testing environments or, more likely, the longer sessions for scanning.

Because we were interested in the initial (nonextinguished) effect of rewarded sounds on visual discrimination, we discarded these last trials (~3 minblocks of data, a total of 48 trials out of 288 trials, corresponding to 24 trials of each reward level; Fig. 1E). Fig. 1F shows $d'$ values and percentage correct of the remaining data. Repeated-measures ANOVA showed a significant main effect of reward on $d'$ ($F_{1,19} = 4.89, P = 0.03$), but the interaction between reward and spatial congruency was not significant ($F_{1,19} = 0.71, P = 0.41$). In a pairwise comparison of high rewards and low rewards, a significant effect of reward was present only when Gabor and sound were spatially congruent, for both $d'$ ($P = 0.03$ for HC vs. LC, paired $t$ test) and percent correct ($P = 0.044$ for HC vs. LC, paired $t$ test). All other pairwise comparisons of $d'$ and percent correct were nonsignificant. Herein we use $d'$ as a measure of performance. Before reward training, behavioral performance for the two sounds did not differ (Fig. S1 A and B). Cross-modal value had no effect on reactions times (Fig. S1 C and D).

The largest and most parsimonious effect of cross-modal reward was observed when stimuli in two modalities were spatially congruent (i.e., in HC vs. LC). However, even when the two stimuli were at different locations, performance was marginally better for high-reward stimuli, which explains why we did not find a significant interaction between reward and congruence. This finding indicates that our observed effects are not strictly spatially specific, and we believe it is likely that spatially nonspecific mechanisms, such as general arousal/alertness, play a role as well. Nevertheless, we note that spatially unspecific mechanisms (e.g., alertness/arousal) are not specific to cross-modal value transfer and could occur for any rewarding stimulus. These considerations aside, our main interest in the present study was related to cross-modal effects at early stages of cortical processing. For this reason, we use the contrast between HC and LC as a proxy for the cross-modal effect of value on behavioral orientation discrimination, and in our analyses this contrast will be the main contrast of interest.

Modulation of Visual Cortex by Cross-Modal Reward Value. Fig. 2 shows the results of pattern classification analysis in individually defined visual areas (defined based on an orthogonal contrast; i.e., responses to the left visual hemifield > responses to the right hemifield and vice versa, for defining visual areas in the right and left hemispheres, respectively; Materials and Methods), as well as...
the average response magnitude in the same regions of interest (ROIs) (Fig. 2 A and B). Classification accuracies were highest for the HC condition [average accuracies: 58.6% for HC, 53.4% for HIC, 51% for LC, and 52.6% for the low-reward-incongruent (LIC) condition; \( P = 0.002, 0.01, 0.38, \) and 0.14, respectively, for comparison with chance, i.e., 50%, paired \( t \) test]. Repeated-measures ANOVA with accuracy as the dependent factor and reward and spatial congruence as independent factors revealed a significant main effect of reward (\( F_{1,16} = 4.77, P = 0.044 \)) and a significant interaction between reward and congruence (\( F_{1,16} = 4.55, P = 0.048 \)). Pairwise comparisons showed that the effect of reward was significant only when the sound and Gabor were spatially congruent (\( P = 0.007, \) HC vs. LC, paired \( t \) test). This effect, greater accuracy in HC compared with LC, was significantly correlated with the difference in behavioral \( d' \) between these two conditions (\( r = 0.61, P = 0.009 \)). Correlation between classification accuracy of the visual cortex and behavioral performance in all other pairwise conditions was nonsignificant (HC-LIC: \( r = -0.02, P = 0.91 \); HIC-LC: \( r = 0.11, P = 0.66 \); HIC-LIC: \( r = -0.07, P = 0.76 \)). The average response magnitude of the visual cortex was not affected by cross-modal value (\( P > 0.05 \) for all, for main effect or interaction with reward and pairwise comparisons).

We replicated these results when eye position offsets were included in our generalized linear models (GLMs) for 11 subjects for whom eye-tracking data were available (SI Materials and Methods and Fig. S2). These results show that the value associated with the sounds affects the accuracy of orientation coding in the visual cortex.

We conducted a number of additional tests to verify these results. First, we ensured that the differential effect of the two sounds on visual orientation coding was related to a difference in reward value as opposed to any difference in their physical attributes (frequency or perceived amplitude). To this end, we repeated our classification analysis for the data of the pretest block, in which subjects were not yet familiarized with the sound values. As shown in Fig. S2, in this pretest block, classification accuracies did not differ between the two sounds.

Second, we replicated our results using the same ROIs for all subjects (Fig. 2 C and D). These ROIs were defined based on the orthogonal contrast of responses to the left > responses to the right for the right visual cortex and vice versa for the left visual cortex at the group level, and were masked by anatomical ROIs of V1 and V2 (Materials and Methods). Classification accuracies were highest for the HC condition (average 59% for HC, 51.5% for HIC, 53% for LC, and 49% for LIC). Repeated-measures ANOVA revealed a significant main effect of reward (\( F_{1,16} = 7.05, P = 0.01 \), but a nonsignificant effect of congruence and the interaction term. In pairwise comparisons, accuracies were significantly different between the HC and LC conditions (\( P = 0.03 \) for comparison of HC and LC, paired \( t \) test). The greater accuracy in HC compared with LC was significantly correlated with the behavioral effect (\( r = 0.65, P = 0.008 \)).

By comparing our results were further correlated by a searchlight analysis in which accuracies were independently computed for each voxel (Fig. 2E and Materials and Methods). This analysis revealed two clusters, one in each hemisphere (demarcated by dashed circles in Fig. 3C), within the same mask used for the second analysis in which classification accuracies were significantly higher for contralateral HC compared with LC stimuli. The activation of the right cluster (peak \( x, y, z : 18, -73, -2 \) was significant when corrected for multiple comparisons (\( P < 0.05, \) small-volume familywise error-corrected). Thus, the same stimulus-specific regions of cortex demonstrated a reward-related change in stimulus representation when classification was independently assessed for each voxel. In addition to these clusters in early visual areas, congruent rewards increased the classification accuracies in a number of other regions, encompassing visual- as well as memory-, attention-, and reward-related areas (Table S2).

**Fig. 2.** Effect of cross-modal value on visual areas. (A) Classification accuracies (bars) and average responses of individually defined visual ROIs (line). (B) Correlation between differences in classification accuracy and differences in \( d' \) values for HC vs. LC. Error bars represent SEM. (C) Same as A, computed in group-level ROIs. Identical ROIs were used for all subjects. (D) Correlation between difference in classification accuracy and difference in \( d' \) values for HC vs. LC in group-level, visual ROIs. (E) Searchlight results masked with group-level visual ROIs (as used for C and D), showing two clusters in right (red) and left (green) visual cortex. For demonstration purposes, data are thresholded at \( P = 0.05 \), uncorrected. The right cluster had significant activation when corrected for multiple comparison (\( P < 0.05, \) small-volume FWE corrected within ipsilateral visual ROI).

**Representation of Value Within Reward-Sensitive Regions and Cross-Modal Areas.** Based on our specific hypotheses as outlined earlier, we tested the effect of cross-modal value on accuracy in three anatomically defined ROIs corresponding to the ventral striatum, vmPFC, and cross-modal areas (Fig. 3). Note that our aim here was to compare two specific hypotheses, rather than to exhaustively test a large number of different brain regions, and as such, we present the present results as exploratory rather than definitive. All of these areas exhibited significantly greater activation for the Gabor + high-reward stimuli compared with the Gabor + low-reward stimuli (\( P < 0.05 \) for all, Wilcoxon signed-rank test), but only in cross-modal areas [i.e., STS/superior temporal gyrus (STG)] was the effect correlated with a behavioral change in \( d' \) (\( r = 0.51, P = 0.03 \)).

To rule out the possibility that this correlation is driven by potential outliers, we performed additional robust regression analyses (using the robustfit algorithm in MATLAB with the Huber weighting function). This analysis showed a significant correlation between the effect size in STS/STG and change in \( d' \) (\( r = 2.14, df = 15, P < 0.05 \)). Robust regression analysis also confirmed that effect size in vmPFC and striatum was not significantly correlated with behavioral change in \( d' \) (\( P > 0.05 \) for all). Formal model comparison using the Bayes information criterion (BIC) (SI Materials and Methods) strongly favored the model containing data from the STS (BIC: STS, \(-26.49 \); striatum, \(-31.58 \); vmPFC, \(-31.49 \)). The posterior probability of the STS model was 0.99. Within the context of our limited model space, this finding provides strong evidence in favor of the hypothesis that the behavioral effects that we observed are crucially mediated by...
multisensory processing areas. Importantly, before subjects learned the reward associations, STS/STG showed no difference in responses to high-reward stimuli and low-reward stimuli (Fig. S3; \( P > 0.05 \), Wilcoxon signed-rank test), indicating that a modulation by sound could not be related to any difference in the sounds’ inherent physical attributes (frequency or perceived amplitude).

We next tested whether the modulation of STS/STG after reward training was correlated with classification accuracies obtained from ipsilateral visual areas, an indication that feedback from cross-modal areas affects primary sensory areas (13). We found a significant correlation between STS/STG response modulation (i.e., high reward vs. low reward) and classification accuracies in visual areas \( (r = 0.50, P = 0.037) \). This correlation was still significant when a robust regression method was used \( (r = 2.13, df = 15, P < 0.05) \).

The results of our whole-brain analysis of the main contrast of high reward vs. low reward and high reward congruent vs. low reward congruent (HC-LC) indicate that in addition to reward and cross-modal regions, a number of areas classically involved in motor planning and executive functions (e.g., supplementary motor areas) were modulated by sound values (Table S1). Thus, our univariate analysis results show that during the orientation discrimination task, sound value is represented in reward processing and executive control networks, as well as in cross-modal areas. Activation within the STS/STG was correlated with the magnitude of the behavioral effect (SI Results), suggesting that this area may play a role in mediating the effect of cross-modal reward associations.

**Discussion**

Reward associations influence perception (5–8, 10). Here we show that these influences extend across different sensory modalities. Sounds previously paired with monetary reward modulated visual perception, leading to better orientation discrimination when a task-irrelevant sound with high-reward associations was presented concurrently with the visual stimulus. There was also a suggestion that the effect of reward was strongest when the auditory and visual stimuli were spatially congruent, although the differences between congruent and incongruent conditions were not statistically significant. Stimulus-specific regions of early visual areas contained a more accurate representation of stimulus orientation when high-reward conditions (signaled by a task-irrelevant auditory stimuli), as opposed to low-reward or incongruent conditions, were presented. This effect was correlated with an increase in orientation sensitivity for high-reward stimuli. In the same regions of the visual cortex, the magnitude of responses did not differ between the two reward conditions, suggesting that reward modulates perception, at least in part, through an increase in the distinctness of neuronal representations rather than through a homogenous response enhancement. Responses within cross-modal areas (i.e., STS/STG) were significantly affected by the reward value of sounds, and this effect was highly correlated with both behavioral and BOLD correlates of visual orientation sensitivity, suggesting that these multimodal areas may underlie the effect of rewarded sound on vision.

The beneficial effect of reward information on sensory processing is in line with the results of a host of previous studies (5–8, 10, 21, 24, 25). These studies have demonstrated that in early sensory areas, the same neurons that process sensory information are modulated by reward (5, 10, 26) and thereby influence perception from the earliest stages of cortical processing. Our findings extend this previous work by showing that task-irrelevant cues presented in another sensory modality modulate perceptual processing according to their reward-predicting properties.

Simultaneously presented sounds can influence visual perception (1, 2) with visual detection or discrimination typically enhanced in the presence of sounds (27–29). Maximal enhancement occurs when visual stimuli are near the detection/discrimination threshold (28), the auditory and visual stimuli overlap in time and in space (13, 29, 30) and sounds carry biologically important information (27, 29, 31). We show that value can modulate audiovisual interactions (and, we hypothesize, multisensory interactions more generally), an effect congruent with ideas on the Bayesian brain (32–34), in which a core idea is that the brain embodies previous beliefs about the structure (i.e., statistics) of its environment. Specifically, here brief temporally (and perhaps spatially) contiguous stimuli are likely to be associated, and thus to share motivational properties. Therefore, visual stimuli have increased importance or salience (and hence presumably attention; ref. 35) conferred on them by concurrently presented high-reward sounds (34, 36). In this broader context, our results have implications for approaches in which basic perceptual and cognitive processes are reappraised in the light of the expectations of the brain about the causal structure of its environment, which here we extend to the domain of motivation.

We suggest that the cross-modal effects that we describe herein are triggered by the reward associations of a task-irrelevant stimulus in another sensory modality, and provide preliminary evidence that a sound’s reward information is transmitted to the visual cortex in the absence of visual stimuli, when attentional effects are minimal (Fig. S4). However, we note that it has proven difficult, if not impossible, to distinguish between reward and attention effects (37) in many studies, and in the present study as well. Attentional effects across sensory modalities are well described for both exogenous spatial attention (38) and object-based attention (39). The perceptual benefits of attention observed in these previous studies are similar to our effects but with several key differences. Exogenous attention [as in, e.g., Störmer et al. (38)] is mediated by the sudden appearance of a stimulus in a certain spatial location. This type of attention by itself cannot explain our effects, because exogenous...
cues will improve the processing of any stimulus that appears at the cued location. Nonetheless, we find that whereas discrimination sensitivity was increased for HC, it was strongly decreased for LC, which could not be predicted by exogenous attentional effects except when modulated by reward associations. In this case, reward could modulate the “salience” of an exogenous high-reward sound could be more effective at summoning attention to a specific spatial location. Endogenous attention, that is, the voluntary shift of attention to a certain location (40), might be triggered by rewarded sounds and could be then transferred to vision (39). This possibility is unlikely, however, given that classical endogenous attention effects need time to develop [a minimum of 300 ms (40)], whereas in the present study, sounds and Gabor stimuli were presented briefly and simultaneously. Future studies are needed to test whether the cross-modal value effect can occur when the influence of attention is minimized, as well as to map the temporal profile of these effects, which could point to the possible underlying mechanisms.

Although mean changes in activity levels are often considered the signature of perceptual benefits of attention and reward, some studies have shown that reward- and information-predictive cues might suppress redundant visual responses and thus sharpen the representation of relevant visual information (10, 22). We did not find that incidental reward signals result in enhancement of visual response amplitudes; instead, our data support an emerging account in which reward/information cues increase the signal-to-noise ratio so as to improve perception (63). Thus, it is also possible that although responses to visual stimuli in the reward condition were larger in magnitude, they were also shorter in duration, and thus produced no clear differences in the BOLD contrast. Another possibility is that cross-modal value modulates subthreshold neuronal responses and thus does not produce a net response enhancement. This is an interesting topic for future study, and would be well suited to neuroimaging techniques with a high temporal resolution, such as EEG and magnetoencephalography.

We found reward associations in a number of regions classically linked into the expectations that the brain learns to associate with valuations. Among these, activity in the STS/STG, an area of known importance in cross-modal interactions (14–16), was correlated with the behavioral effect, and was a better predictor of behavior when a formal model comparison was performed. This suggests a model in which reward associations, either generated in regions typically associated with value or reflected in local processing in the sensory cortex (8, 26), influence sensory processing in other modalities via cross-modal association areas, such as the STS/STG. This suggestion is in line with a task-specific role of reward, in which the specific intermingling processing stages involved in a task, not the final effectors alone, are informed and influenced by value-related information. Whether activity in cross-modal areas plays a causal role in increased perceptual sensitivity remains to be established.

The present study has addressed the question of whether task-irrelevant reward associations in one sensory modality improve the processing of temporally congruent stimuli presented in another modality. We have shown that they do, and have presented evidence indicating that increases in perceptual accuracy are reflected in the discriminability of neuronal representations, very likely mediated by activity in cross-modal areas. This finding provides insight into the interactions that the brain has about the causal structure of its environment, and sheds light on a mechanism that in and of itself is likely important for adaptive behavior in ecological contexts.

Materials and Methods

Participants. Forty-four subjects participated in either a behavioral study outside the scanner (n = 24; 14 females; mean age, 25 y; range, 22–33 y) or a combined behavioral and fMRI experiment inside the scanner (n = 20; 11 females; mean age, 24 y; range, 23–30 y). The data for three subjects from the fMRI experiments were analyzed only in terms of behavioral effects, owing to technical problems with image acquisition. Each subject provided oral and written consent for his or her participation. The study was approved by the local Ethics Committee of Berlin Charité University Hospital.

Stimuli and Tasks. We used a Pavlovian conditioning paradigm to familiarize the subjects with sounds and their associated rewards (Fig. 1A). The pairing of sounds (high or low) with the visual Gabors was counterbalanced across subjects. The visual orientation discrimination task (Fig. 18) was adopted from one described in a previous study (29). In this task, after an initial fixation of variable duration, a Gabor stimulus was briefly (250 ms) presented at a parafocal location, and subjects reported its tilt orientation (i.e., two-alternative forced choice, clockwise or counterclockwise relative to the horizontal meridian). The tilt of the Gabor from horizontal was set to each subject’s discrimination threshold (75% performance). Subjects were instructed to maintain fixation throughout the trials (fixation circle diameter 1°, eye tracking described in SI Materials and Methods). At the same time as presentation of the visual stimulus, a tone was played either from the same side (spatially congruent) or opposite side (spatially incongruent) as the Gabor. The tone had either a low frequency (160 Hz) or high frequency (380 Hz) and was uninformative regarding the visual stimulus orientation. This was provided for a 2 × 2 factorial design with sound identity (high reward and low reward) and its spatial congruence (congruent and incongruent) as the two independent factors. No feedback regarding correct performance was provided to the subjects. We collected a total of 199–288 trials of the audiovisual task, corresponding to 72 trials for each of the four experimental conditions. To avoid extinction, short miniblocks of sound localization and orientation judgment tasks were interleaved (SI Materials and Methods).

Univariate Analysis of the fMRI Data. Details of data acquisition and preprocessing are provided in SI Materials and Methods. To identify cortical regions modulated by sound rewards, we created a GLM with eight main regressors. These regressors included responses to HC, HIC, LC, and LIC conditions, modeled separately for left and right visual hemifields. Because reward conditioning and orientation discrimination miniblocks occurred alternately and in close temporal proximity to each other, four additional regressors were included to account for reward conditioning, so as to model two types of sound-reward pairs (high and low reward) on each side (left and right). Finally, two additional regressors were included to model the instruction display presented at the beginning of each miniblock (one for sound miniblocks and one for audiovisual miniblocks). Thus, the full GLM model included 12 regressors, which was modeled at each voxel time point at the level of each subject. This was done for a 2×2 factorial design with sound identity (high reward and low reward) and its spatial congruence (congruent and incongruent) as the two independent factors. No feedback regarding correct performance was provided to the subjects. We collected a total of 199–288 trials of the audiovisual task, corresponding to 72 trials for each of the four experimental conditions. To avoid extinction, short miniblocks of sound localization and orientation judgment tasks were interleaved (SI Materials and Methods).

Our behavioral analysis revealed that the effect of sounds on orientation discrimination performance decayed toward the end of the experimental session, with a reversal of the effect in the last three miniblocks (Fig. 1A). Thus, we included an additional regressor in our GLM to model the effect of the last reversal trials modeled with a boxcar function that covered all of the events after the third audiovisual miniblock of the last run (and thus undivided by different conditions). Six session-specific motion parameters were modeled as covariates of no interest. To test for regional-specific condition effects, we included linear contrasts for each subject and each condition (first-level analysis). These contrasts included the main effect of reward (HC + HIC) vs. (LC + LIC) pooled across congruency conditions and the contrast between HC and LC conditions. The resulting contrast images were entered into a second-level analysis, modeled by one-sample t tests (familywise correction). Multiple-brain results were thresholded at P < 0.001 (uncorrected, k = 10; Table S1).

To define the visually responsive ROIs, we identified areas activated by contralateral visual stimuli, showing higher activation in the contrast of left Gabor vs. right Gabor and vice versa as in our univariate, first-level GLMs. This contrast thus highlights right (higher activation for left Gabors vs. right Gabors) and left (higher activation for left Gabors vs. right Gabors) visual cortices for each individual subject. Note that this contrast (left vs. right Gabors) is orthogonal to both reward and spatial congruence (fully randomized across the two visual fields), and thus the activity of selected visual ROIs is not directionally biased to show a difference between any of the factors of interest.

We also reproduced our results with a second set of visual ROIs that were identical for all of the subjects. Here we used the second-level contrast of left Gabor vs. right Gabor to identify the right and left visual areas at the group level. We then masked these activations with anatomically defined ROIs of areas V1 and V2 constructed based on the anatomy toolbox of SPM (42). This enabled us to identify visually responsive regions that matched the topology probability maps of early visual areas known to be involved in low-level processing of stimulus orientation (43). Ideally, we also could have used detailed retinotopic mapping techniques to precisely map the topographic location of cortical areas and to identify our visual ROIs in relation to these topographic maps, but in the interest of time, we decided not to do so. For regions outside the visual cortex, ROIs were selected anatomically. The ventral striatum ROI consisted of the bilateral caudate, putamen, and globus pallidus, the vmPFC ROI comprised the bilateral gyrus rectus and medial
and the STS/STG ROI comprised bilateral superior and middle temporal gyrus as defined in the Anatomical Automatic Labeling atlas (44). The effect sizes of these ROIs (Fig. 3) were computed by averaging the contrast estimates across all voxels of an anatomical ROI.

MVPa of fMRI Data. We performed a MVPa to assess the effect of rewarded sounds on a trial-by-trial pattern classification accuracy of early visual areas (details provided in SI Materials and Methods). We used several approaches for the MVPa. In the first approach (Fig. 2 A and B), we performed a pattern classification analysis in individually defined visually responsive ROIs. The input of the support vector machine classifiers consisted of t-values of every trial for all voxels of a certain ROI. Eight classifiers were constructed, four for each hemisphere, to decode the tilt orientation (clockwise or counterclockwise) in the HC, HIC, LC, and LIC conditions of the contralateral visual field for a total of eight classifiers. The accuracies were then averaged across the two hemispheres (Fig. 2 A and B).

We also performed two supplementary analyses (Fig. 2 C and D). First, we used the same visual ROIs for all of the subjects, minimizing the possibility of subject-specific selection bias. Then we used a whole-brain searchlight method (SI Materials and Methods), which avoids ad hoc voxel selection (45). Note that here a whole-brain analysis was performed, and visual ROIs were subsequently used for small-volume correction.

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