Sensory modalities are not separate modalities: plasticity and interactions

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Historically, perception has been viewed as a modular function, with the different sensory modalities operating independently of each other. Recent behavioral and brain imaging studies challenge this view, by suggesting that cross-modal interactions are the rule and not the exception in perception, and that the cortical pathways previously thought to be sensory-specific are modulated by signals from other modalities.

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Abbreviations

fMRI functional magnetic resonance imaging PET positron emission tomography

Introduction

Cross-modal integration is performed on a vast level in the brain and contributes significantly to adaptive behavior in our daily life. Very little is known about how integration is achieved or its underlying neural mechanisms, however, because the overwhelming majority of studies on perception have focused on one sensory modality. Studying perception in an isolated single modality would be justifiable if different modalities processed sensory inputs independently of each other, as separate 'modules'. But are sensory modalities really separate modules? A variety of evidence seems to counter this notion of modularity. In this review, we summarize the evidence for vigorous interaction among sensory modalities.

Plasticity across sensory modalities

Both animal and human studies suggest that sensory modalities in early stages of development are not as inherently distinct and independent as was previously once thought. For example, in a study of cross-modal plasticity Sur *et al.* [1] removed the superior colliculus of both the ferret and the hamster on the day of birth by direct ablation. They also deprived the medial geniculate nucleus or the ventrobasal nucleus from their normal sensory input by sectioning the major input pathways. The retina then invaded these thalamic nuclei, which under ordinary circumstances relay auditory and somatosensory signals to the cortices, respectively. They found that visual responses (i.e. responses triggered by light stimulation on the retina) were elicited from neurons in the auditory or the somatosensory cortex.

More recently, Sur and co-workers $[2^{\bullet\bullet}]$ reported that these animals interpreted the activation of the rewired auditory cortex as visual sensation, indicating a functional relevance of the rewiring. There is also evidence for a cross-modal postnatal perceptual enhancement by prenatal sensory stimulation. When bobwhite quail chicks were prenatally exposed to an auditory, visual, tactile or vestibular stimuli, their postnatal auditory and visual responsiveness was enhanced, irrespective of the modality of prenatal stimulation [3–6].

Cross-modal plasticity has also been reported in humans that have had sensory deprivation in early life [7–13]. The typical finding in these studies has been that sensory deprivation in one modality starting from an early period of life causes the cortical area that is normally devoted to that modality to be used by some other modality. Early studies reported that visual event-related potentials tend to be enhanced in early onset deaf individuals [7,8].

Auditory-evoked potentials have also been recorded from posterior (occipital) regions in early and late blind subjects [9]. Although suggestive, however, these studies were neither able to identify exactly which part of the brain is responsible for the enhanced activity, nor able to examine functional relevance of the activity.

Studies that use a perceptual task have been more informative in this regard. For example, Uhl et al. [10] have provided evidence for posterior DC potentials in blind subjects engaging in a tactile reading task. Similarly, a positron emission tomography study (PET) by Sadato et al. [11] indicated the activation of primary and secondary visual cortical areas induced by Braille reading in early blind subjects, but the deactivation of the same areas relative to the rest condition induced by Braille reading in sighted subjects [11]. Simple tactile stimuli that did not require discrimination produced no activation of visual areas in either group.

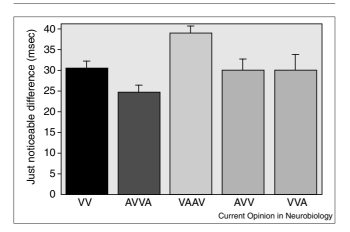
Furthermore, the same researchers applied a train of pulses of transcranial magnetic stimulation to the occipital region of blind subjects engaged in the Braille reading task, and showed that indeed it degraded performance but selective to this task only [12].

Taken together, these animal and human studies indicate a surprising degree of neural plasticity in early stages of life, and clearly argue against the most stringent version of brain segregation and modularity of sensory modalities.

Interaction across modalities

In the phenomenological and psychophysical literature, a wealth of literature on mature perceptual systems indicates

Figure 1



Visual temporal resolution modulated by sounds. Temporal resolution is often expressed in terms of just noticeable differences (JNDs), the smallest temporal gap between two signals at which the two signals are not perceptually fused. JNDs are plotted for conditions of a vision only baseline (V-V), a preceding sound and a following sound (A-V-V-A), two sounds inserted between the two visual signals (V-A-A-V), one sound preceding the two visual signals (A-V-V), and one sound following the two visual signals (V-V-A). During the task, two LEDs were switched on with asynchrony varying from -60 to +60 ms, and the subject had to judge which LED was first. The sounds were always presented through a single speaker located between the two LEDs. The JND is smaller (thus, the visual temporal resolution is better) in the A-V-V-A condition, and larger (worse) in the V-A-A-V condition. The results of the last two conditions, particularly that of the A-V-V condition, indicate that the modulation effect is not due to general warning or arousal effects [22].

that there is vigorous interaction and integration among the sensory modalities — a far greater interaction than what is expected from a 'naive modularity' point of view. One of the most intriguing examples of cross-modal interaction is synesthesia, in which an involuntary conscious sensation (such as color) is induced by a stimulus in another modality (such as sound). Synesthesia can occur in normal, healthy populations, in brain-damaged or sensorydeafferented patients, or in people who are addicted to hallucinogenic drugs [14••].

Vision alters other modalities

In normal people, the effects of vigorous cross-modal integration are made apparent under cleverly designed artificial conditions. The McGurk effect exemplifies such a condition [15]. The McGurk effect is a perceptual phenomenon in which vision alters speech perception (e.g., the sound 'ba' tends to be perceived as 'da' when it is coupled with a visual lip movement associated with 'ga' [15]).

The spatial location of a sound source can also be drastically influenced by visual stimulation. This effect is known as the 'ventriloquist effect' [16], and is experienced frequently in daily life when watching television or movies in which the voices are perceived to originate from the actors on the screen, despite a potentially large spatial discrepancy between the two.

It has been shown that tactile location, such as location of a finger pointing, can also be 'captured' by visual location [17]. All these effects emphasize the strong effect of visual signals on the other modalities, consistent with the commonsense notion that human is primarily a visiondominated animal.

Sound alters the temporal aspects of vision

Although the best-known cross-modal effects are those of vision influencing other modalities, visual perception can also be altered by other modalities. All the earlier reports of alteration of visual perception by other modalities have been in the temporal domain. For example, the perceived duration [18] or rate [19–21] of a visual stimulus has been shown to be influenced by accompanying sound signals. More recently, Scheier et al. [22] have shown that visual temporal resolution can be either improved or degraded by sounds, depending on the temporal relationship. They showed that when two lights are turned on with a small temporal delay (in the -60 to +60 ms range), the accuracy of temporal order judgment is better when a sound precedes and another follows the visual stimuli (the audiovisual-visual-audio [A-V-V-A] time order). In contrast, the subjects' performance becomes worse when two sounds are inserted between the two visual stimuli (V-A-A-V time order), as illustrated in Figure 1.

Sound alters other aspects of vision

The great body of behavioral findings on cross-modal interactions has been accounted for in a 'modality appropriateness' hypothesis [23]. This hypothesis postulates that the modality that is most appropriate or reliable with respect to a given task is the modality that dominates the perception in the context of that task. Vision has a higher spatial resolution, hence its dominance in spatial tasks (e.g. the ventriloquist effect and visual capture), whereas audition has a higher temporal resolution, hence its dominance in temporal tasks.

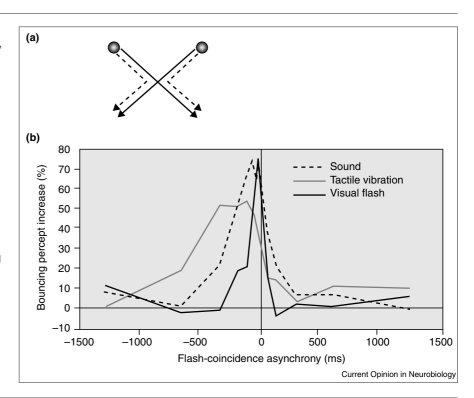
Alteration of vision by sound, however, turns out to be not limited to temporal aspects. The perceived intensity of a visual stimulus has been shown recently to be enhanced in presence of sound [24]. Moreover, the quality or structure of visual perception itself can be altered by auditory stimuli, according to the latest psychophysical studies.

Sekuler et al. [25] have shown that the presence of a sound can also alter the perceptual interpretation of an ambiguous visual motion event. Two identical visual targets moving across each other can be perceived either to bounce off or to stream through each other, as their trajectories are nearly identical (Figure 2a). Nonetheless, most observers report a perception of streaming, not bouncing motion.

If a brief sound is added at the moment that the targets coincide visually, however, visual perception is strongly biased in favor of bouncing motion [25]. The sound has to have a sharp onset to induce this effect. The ecological origins of

Figure 2

Cross-modal modulation of visual motion perception. (a) The ambiguous motion display, in which two objects move across each other. Observers typically perceive an object as streaming with an X-shaped trajectory when there is no accompanying sound or an accompanying sound that is not synchronous with the visual crossing. When there is a sound (or a flash or a tactile vibration) synchronized with visual coincidence. however, the perceptual dominance reverses and most observers now see the objects as bouncing against each other. (b) Temporal tuning curves of the bounce-inducing effects of a synchronous transient signal in auditory, visual or tactile modality. The percentage increase in the perception of bounce is plotted against asynchrony between the transient signal in each modality and the visual coincidence. Thus, qualitatively the same effect can be obtained as long as there is a transient sensory stimulus that is approximately synchronized with the critical visual event, regardless of the modality in which the transient stimulus is given.



this phenomenon are intuitively obvious: it should be related to the fact that the majority of collision events in the natural environment yield synchronized cross-modal signals.

Other studies on this phenomenon have revealed some unexpected properties. First, a transient sensory stimulus biases visual perception toward bouncing irrespective of its modality; for example, a brief visual flash or a brief tactile vibration on an observer's finger also induces bouncing perception [26,27]. The stimulus, however, has to be synchronized with the visual coincidence of two objects (the effective time window was found to be in the range of -300 to +200 ms, -600 to +100 ms and -100 to +100 ms for auditory, tactile and vision stimuli, respectively; see Figure 2b).

Thus, the same effect can be obtained qualitatively as long as there is a transient sensory stimulus that is approximately synchronized with the critical visual event, regardless of the modality in which the transient stimulus occurs. This type of perceptual modulation may serve as a collision detector at a functional level, yet there is a series of evidence indicating that it is implemented through a more generic and efficient mechanism of spatial attention [26–28]. Developmentally, at five months of age, human infants show behavioral evidence for the adult-like perceptual change caused by a synchronous sound [29–30]; this age is roughly the maturation onset of spatial attention mechanisms.

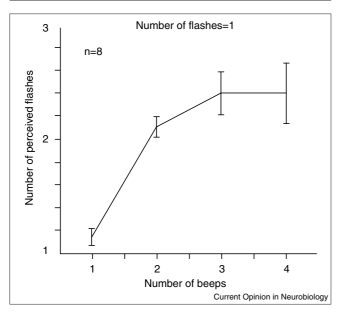
A recent study has shown that the alteration of vision by sound is not limited to the situations of ambiguity in the visual stimulus. The 'illusory flash effect' reports that an illusion in which

a radical change in the phenomenological quality of perceiving a non-ambiguous visual stimulus is induced by sound [31°]. When a single brief visual flash is accompanied by multiple auditory beeps, the single flash is perceived as multiple flashes (see Figure 3). Control conditions, catch trials and many other observations indicate that the illusory flashing phenomenon is indeed a perceptual illusion, and is not due to the difficulty of the task or some cognitive bias (caused by sound).

The illusory double flash is perceptually very similar to the physical double flash. Furthermore, the illusion is very robust to the observer's knowledge about the physical stimulus, and to variations in stimuli parameters. The temporal tuning of this effect was also measured by varying the relative timing of visual and auditory stimuli. The illusory flashing effect decreased at separations greater than 70 ms; however, illusory flashing occurred as long as the beeps and flash were within about 100 ms - consistent with the integration time of polysensory neurons in the brain [32,33].

The alteration of vision by sound in this experiment was found to be asymmetrical: alteration occurred strongly only when a single flash was coupled with multiple beeps, and not when multiple flashes were paired with a single beep. In other words, strong alteration of vision by sound occurs only when sound is more discontinuous and structured than the visual stimulus. A similar phenomenon seems to be at work in a study that investigated the effect of vision on hearing [34]. Saldaña and Rosenblum's [34] data suggest that only the discontinuous visual stimulus has a strong effect on the perception of the sound.

Figure 3



Sound-induced illusory flashing. The number of flashes reported by the observers is plotted against the number of beeps that accompanied the visual stimulus, in trials where the visual stimulus comprises only one flash. Observers report seeing two or more flashes, when a single flash is accompanied by two or more beeps.

Cross-modal interactions depend on stimuli structure

The direction of cross-modal interactions has been thought to be determined by the relative appropriateness of the modalities involved in the task, as mentioned above. The evidence discussed above, however, seems to indicate that the direction of cross-modal interactions depends, at least in part, on the structure of the stimuli; that is, the modality that carries a signal which is more discontinuous (and hence more salient) becomes the influential or modulating modality. Such a hypothesis would also account for the findings of streaming/bouncing motion studies discussed above, in which the transient, hence more discontinuous and structured, stimulus alters the perception of a continuous visual stimuli regardless of its modality (auditory, tactile or visual) [25–30].

Neural mechanisms

At what point along the perceptual processing pathway do these cross-modal interactions take place? Recent data from brain imaging studies suggest that they occur at brain sites that used to be considered as modality-specific. For example, Calvert et al. [35] carried out a functional magnetic resonance imaging (fMRI) study that showed that the primary auditory cortex is activated when a talking face is viewed in the absence of sound. The activation was observed specifically in speech or pseudo-speech type of lip movements, but not in other type of lip movements.

The same research group has also reported that activity in visual (V5) and auditory (BA 41/42) cortices after exposure to bimodal (audio-visual) speech is enhanced relative to activity after a unimodal stimulus [36*]. The enhancement was found

to be contingent on congruency between the audio and visual signals. But as these studies deal only with speech perception, would their findings generalize to other types of stimuli?

Two very recent studies may have more general implications. An fMRI study has shown that tactile stimulation of a hand enhances activity in the visual cortex when the touched hand is on the same side as the visual stimulus [37••]. Likewise, an event-related potential (ERP) study also suggests that activity in the visual cortical areas is modulated by sound [38]. The modulation was found as early as 100 ms after the visual stimulus onset. This study used a brief flash and a brief beep as visual and auditory stimuli.

In contrast to speech stimuli, which inherently hinge on some higher level matching process (such as congruency detection), the stimuli used in the last two studies are very simple, and thus the results may have more general implications. In particular, the results of the ERP study, unlike any of the other findings discussed above, may have very general implications because they do not require congruency between the stimuli in the two modalities and do not seem to involve spatial attention mechanisms.

Together, the results of the studies summarized in this review challenge the common belief that 'modality-specific' cortices function in isolation from other modalities.

Conclusions

We have discussed a wide variety of evidence against the notion of strict modularity of sensory modalities. Both animal studies and human-deprivation cases provide evidence for a surprising degree of cross-modal plasticity in cortical processing. Psychophysical data indicate that interaction between modalities is the rule as opposed to the exception in brain function, and brain imaging and recording studies provide evidence against modularity and for interaction in areas traditionally thought to be unimodal.

Motivated by some of the recent psychophysical findings, we put forth a new hypothesis for multisensory interactions. We propose that the transient/discontinuous signals possess a special status in determining the direction of cross-modal interactions, with the transient stimulus strongly influencing the multimodal perception, regardless of the modality in which it occurs.

Acknowledgements

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