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# Action-based predictions affect visual perception, neural processing, and pupil size, regardless of temporal predictability



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#### ABSTRACT

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Sensory consequences of one's own action are often perceived as less intense, and lead to reduced neural responses, compared to externally generated stimuli. Presumably, such sensory attenuation is due to predictive mechanisms based on the motor command (efference copy). However, sensory attenuation has also been observed outside the context of voluntary action, namely when stimuli are temporally predictable. Here, we aimed at disentangling the effects of motor and temporal predictability-based mechanisms on the attenuation of sensory action consequences. During fMRI data acquisition, participants (N = 25) judged which of two visual stimuli was brighter. In predictable blocks, the stimuli appeared temporally aligned with their button press (active) or aligned with an automatically generated cue (passive). In unpredictable blocks, stimuli were presented with a variable delay after button press/cue, respectively. Eye tracking was performed to investigate pupil-size changes and to ensure proper fixation. Self-generated stimuli were perceived as darker and led to less neural activation in visual areas than their passive counterparts, indicating sensory attenuation for self-generated stimuli independent of temporal predictability. Pupil size was larger during self-generated stimuli, which correlated negatively with the blood oxygenation level dependent (BOLD) response: the larger the pupil, the smaller the BOLD amplitude in visual areas. Our results suggest that sensory attenuation in visual cortex is driven by action-based predictive mechanisms rather than by temporal predictability. This effect may be related to changes in pupil diameter. Altogether, these results emphasize the role of the efference copy in the processing of sensory action consequences.

#### 1. Introduction

The sensory consequences of one's own actions result in a less intense experience than identical but externally generated events. This phenomenon, called sensory attenuation (Brown et al., 2013; Hughes et al., 2013), is thought of as one of the mechanisms allowing an organism to distinguish between internal and external events (Frith et al., 2000; Haggard and Tsakiris, 2009). Attenuation of self-generated sensory events is a phenomenological experience (Blakemore et al., 1999a; Sato, 2008; Weiss et al., 2011) reflected in reduced neural processing (Schafer and Marcus, 1973; Blakemore et al., 1998, 1999b; Martikainen, 2004; Bäß et al., 2008; Aliu et al., 2009; Shergill et al., 2013; Straube et al., 2017; Arikan et al., 2019; Pazen et al., 2019;

Uhlmann et al., 2020; Schmitter et al., 2021). These findings have predominantly been discussed within the framework of internal forward models which suggests that the sensory consequences of one's actions are predicted based on efference copies generated during motor planning (Wolpert et al., 1995). Despite the efference-copy hypothesis being suggested as a general mechanism occurring in all modalities (Brown et al., 2013), sensory attenuation has mainly been investigated in somatosensation (Blakemore et al., 1998, 1999a, 1999b) and audition (Sato, 2008; Aliu et al., 2009; Bäß et al., 2009; Weiss et al., 2011; Sanmiguel et al., 2013; Mifsud et al., 2016a), with evidence in the visual domain remaining inconclusive. While some studies reported lower perceptual sensitivity (Cardoso-Leite et al., 2010; Dewey and Carr, 2013) and/or reduced neural responses (Leube, 2003; Straube et al., 2017;

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Arikan et al., 2019; Pazen et al., 2019; Uhlmann et al., 2020; Schmitter et al., 2021) for self-generated visual stimuli, others observed no (Schwarz et al., 2018) or ambiguous effects (Yon and Press, 2017).

The notion of motor action being crucial for sensory attenuation is challenged by the observation that mere temporal prediction of a stimulus can also attenuate perceptual and neural processing (Summerfield et al., 2008; Bendixen et al., 2009; Alink et al., 2010; Todorovic et al., 2011; Kok et al., 2012; John-Saaltink et al., 2015). It has been suggested that besides motor predictions, better temporal predictability of stimulus onset may contribute to sensory attenuation and neural suppression effects for actively generated stimuli, simply because of a heightened temporal control when the presentation of a stimulus is caused by one's own action (Hughes et al., 2013). Many paradigms indeed compare self-generated stimuli with stimuli that are externally generated at random time points, which creates a confound as the temporal predictability between actively and externally generated stimuli differs in such paradigms. As such, sensory attenuation for self-generated stimuli may not solely be due to efference-copy mechanisms but be explained at least partially by increased temporal predictability of these stimuli.

This study aimed to disentangle the roles of action-based and general temporal prediction mechanisms in sensory attenuation of visual action consequences. During functional magnetic resonance imaging (fMRI), participants engaged in a visual intensity judgment task, judging which of two subsequent stimuli was brighter. Stimuli were elicited either by an active button press or by the computer; stimulus onset was either temporally predictable or unpredictable. We manipulated luminance, the physical quantity related to the intensity of a visual stimulus, and considered brightness, representing the perceived intensity as dependent variable. This was motivated by the extensive work on sensory attenuation in the auditory (e.g., Sato, 2009; Weiss et al., 2011) and somatosensory (Blakemore et al., 1998, 1999a) senses. The results of these lines of research suggest that perceived stimulus intensity is attenuated for sensory events that result from one's own actions. Neurons in visual cortex have long been known to be sensitive to orientation (Graham et al., 1993; Ling et al., 2009) and contrast (Boynton et al., 1996; Avidan et al., 2002), and were thought of as less (if at all) responsive to uniform illumination (Hubel and Wiesel, 1968). However, more recent evidence suggests that primary visual areas in humans can respond to changes in luminance (Haynes et al., 2004; Vinke and Ling, 2020) (but also see: Cornelissen 2006). In the light of this, we expected that visual events would be perceived as darker and result in a suppressed BOLD response in primary visual cortex for active as compared to passive trials. Furthermore, we hypothesized that temporal predictability of stimulus onset and voluntary action would interact such that the attenuation (behavioral and neural) was strongest for trials being active and predictable.

Finally, we were interested in the role of pupil size in sensory attenuation effects. It is long established that pupil size is influenced by stimulus luminance (Loewenfeld, 1958; Larsen and Waters, 2018). However, pupil size is also affected by brightness such that stimuli perceived or expected as brighter give rise to stronger constriction, even if luminance remains physically unchanged (Laeng and Endestad, 2012; Binda et al., 2013; Naber and Nakayama, 2013). Thus, we expected that dilation was predictive of perceived stimulus intensity (i.e., lower perceived intensity resulting in bigger pupil size). In addition, we aimed to explore the relation between pupil size and the neural effects of sensory attenuation.

# 2. Materials and methods

## 2.1. Participants

25 subjects (15 female, 10 male, age = 23.8, SD = 2.2) participated in the experiment. All participants were right-handed (Edinburgh Handedness Inventory), had normal or corrected-to-normal vision and reported no history of neurological or psychiatric diseases. Three participants were excluded after data acquisition – two because of heavy head motion and one due to suspected neurological issues. Accordingly, the fi-

nal sample comprised 22 individuals (13 female, 9 male, age = 23.6, SD = 2.0). The study protocol was approved by the local ethics committee in accordance with the Declaration of Helsinki, and all participants provided written informed consent.

#### 2.2. Stimuli and procedure

Participants performed a visual intensity judgment task in the MRI scanner with constant, minimal lighting distributed equally across the scanner room. The stimuli were presented on a screen (refresh rate 60 Hz) located behind the scanner which participants saw via an MR-compatible eye-tracking mirror. Additionally, participants were equipped with two button pads, one for each hand, that were placed on the respective legs. The software used for stimulus presentation was PsychoPy (version 1.64; Peirce et al., 2019). Luminance measurements were performed using an i1Display Pro photometer (X-Rite Pantone, Grand Rapids, USA).

The task consisted of judging which one of two gray discs was brighter. Both stimuli were in the color of the gray of the monitor (R = G = B). The first stimulus was always presented at a luminance of 29.5 cd/m<sup>2</sup>, whereas the second could either have a luminance of 26.6, 28.0, 29.5, 31.0, or 32.7 cd/m<sup>2</sup>. Brightness is a non-linear function of luminance, that is, equal increments in luminance do not correspond to equal increments in brightness (Stevens, 1957, 1966; Poynton, 1993). We chose precisely the aforementioned luminance values since they could be expected to be approximately perceptually equidistant according to the CIE 1976 L\* function, given that the white of monitor  $(R = G = B = max; 68.8 \text{ cd/m}^2)$  is used as reference white for L\*. All stimuli were presented against a background which also had the color of the monitor gray and always had a luminance of 16.1 cd/m<sup>2</sup>. We added a white frame (68.8 cd/m<sup>2</sup>) to this background to control for any unsought anchoring effects (Gilchrist and Bonato, 1995; Gilchrist et al., 1999). To diminish possible crispening effects (i.e., facilitated brightness discrimination for stimuli similar in intensity to the background; Takasaki, 1966), a black outline (2 pixel wide) was added to the stimulus (Whittle, 1992).

Trials started with a black fixation dot (0.5° visual angle), followed by an actively elicited or automatically triggered cue (enlarged dot, 0.9° visual angle, 300 ms) which indicated to the participant that the first stimulus would be launched shortly. The strength of the effect of sensory attenuation decreases within a few hundred milliseconds after action execution (Bays et al., 2005; Aliu et al., 2009), thus, we expected sensory attenuation to affect the first of the two stimuli more strongly than the second, as it was closer in time to the button press. Consequently, all imaging and pupillometry analyses will refer to the first stimulus, which was physically identical throughout the experiment and will be referred to as "stimulus of interest" from here on. Subsequently, the stimulus of interest and the comparison stimulus were presented (both 1000 ms and 2° visual angle), separated by a variable inter-stimulus interval (ISI; 1000, 1350, 1650, 2000 ms). Upon offset of the comparison stimulus, participants were prompted with the question "Which stimulus was brighter?" and answered by button press within a time window of 2500 ms. The response was followed by an inter-trial interval (ITI) which defaulted to a duration of 500 ms, with added jitter based on the remaining time from the active button press phase and the response phase (see Fig. 1).

The experiment comprised four conditions each of which differed slightly in the details of the trial structure. Trials were manipulated with regard to (1) voluntary generation and (2) the temporal predictability of the stimulus of interest. For the main effect of the efference copy, half of the trials involved a motor action (active) which elicited the stimuli, while no movement was performed in the other half (passive) and stimuli were presented automatically. In active trials, the appearance of the fixation dot indicated the start of a four second time window in which a voluntary button press (right index finger) had to be performed. Participants were instructed to withhold their action for a few milliseconds

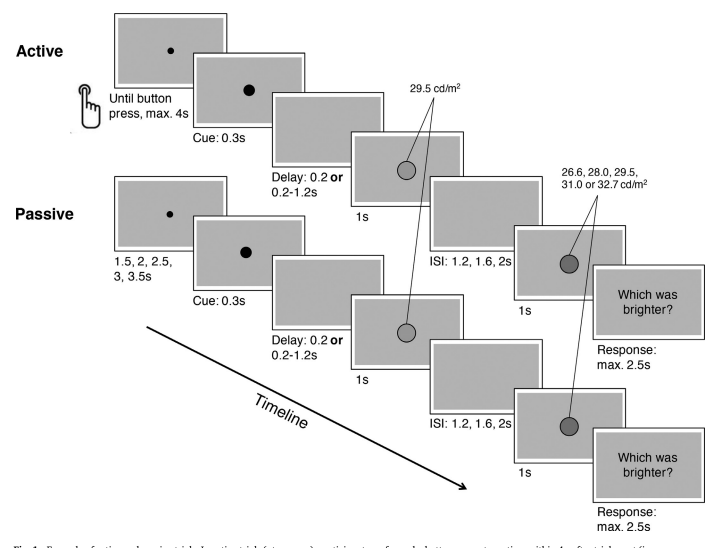


Fig. 1. Example of active and passive trials. In active trials (top row), participants performed a button press at any time within 4 s after trial onset (i.e., appearance of fixation dot). The button press triggered a cue which was followed by the presentation of the visual stimulus. Stimuli were presented either after a variable delay (unpredictable, 200, 450, 700, 950, 1200 ms) or after a constant period (predictable, 200 ms). Following the stimulus of interest, after a random ISI (1.2–2 s) a comparison stimulus was presented. Participants' task was to report which stimulus they perceived as brighter. In passive trials (bottom row), the trial structure was identical except that participants did not perform button presses, instead cue and stimuli appeared automatically.

so that the button press could be consciously prepared and occur in a willed manner, rather than being an automatic mechanism in response to the fixation dot (Rohde and Ernst, 2013). As a direct consequence of the button press, the fixation dot would enlarge and serve as cue signaling the subsequent presentation of the stimuli. In passive trials, no button press was executed. The cue forecasting the upcoming stimulus was identical to the active conditions (enlarged dot) but was generated automatically by the computer. The cue could appear at variable times after trial onset (1500, 2000, 2500, 3000, 3500 ms) to mimic the temporal differences that might occur in the active condition. Notably, the cue indicated a button press (active: the actual button press; passive: a simulated button press performed by the computer) in all conditions and always informed the participant that the action consequence, i.e., the stimulus of interest, would occur after a predictable or unpredictable amount of time in the predictable and unpredictable conditions, respectively. Furthermore, predictability of stimulus onset was manipulated by altering the duration of the time interval between cue offset and onset of the stimulus of interest. In temporally predictable trials, the stimulus of interest was always presented 200 ms after cue offset (thus 500 ms after cue onset). The brief delay was introduced so that participants had the chance to also build up a temporal expectation about stimulus onset in passive trials. Temporal cueing experiments evidenced

that stimulus onset asynchronies (cue onset to stimulus onset) as short as 250 or 500 ms are sufficient to lead to temporal predictability effects in detection and discrimination tasks, respectively (Griffin et al., 2001; Nobre and van Ede, 2018). Moreover, short cue-intervals of 250 ms have also been shown to be sufficient to lead to neuronal sensory attenuation in visual cortex (Fischer et al., 2013). Lastly, the brief cue-stimulus interval was chosen as a trade-off between being long enough to prepare for the onset of the stimulus in the passive conditions but short enough to account for the effect of sensory attenuation diminishing over time (Bays et al., 2005; Pinheiro et al., 2019). In contrast, in unpredictable trials, the onset of the stimulus of interest occurred after a randomly chosen interval (either 200, 450, 700, 950, or 1200 ms, picked with equal probability), relative to cue offset.

Thus, the four conditions were: active predictable (AP), active unpredictable (AU), passive predictable (PP), and passive unpredictable (PU). Each condition (AP, AU, PP, PU) included 60 trials that were evenly distributed across three experimental runs, yielding 240 trials in total (note, that for one participant only 232 trials were collected as data collection was aborted 8 trials prior to the end). Within a run, all conditions were presented and trials were grouped by condition into mini-blocks of 10 trials (2 per run). We pseudo-randomized the order of conditions within a run - so that active and passive conditions alternated - as well

as the ISIs and the delay between cue and the stimulus of interest (the latter just for the passive conditions). Prior to each mini-block, participants were visually instructed about the following condition. To assure an identical number of scanned volumes per trial, the differences in trial duration were compensated for in the ITI. The default ITI (500 ms) was extended by the difference between maximum jitter time and presented jitter time in the current trial for (1) all predefined jittered elements (passive button press, unpredictable stimulus appearance, ISI) and (2) all varying elements controlled by the participant (reaction time for active button press and judgment). Consequently, the length of each trial amounted to a total of 12.5 s.

#### 2.3. Data acquisition

#### 2.3.1. Functional MRI

MRI Data collection was conducted using a 3 Tesla MR scanner (Siemens Magnetom TIM Trio, Erlangen, Germany) at the Department of Psychiatry and Psychotherapy, Philipps-University Marburg, using a 12-channel head-coil. Participant's heads were stabilized in order to reduce head motion artifacts. Time courses of functional activation were obtained using a T2\*-weighted gradient-echo planar imaging sequence (EPI) sensitive for the blood oxygenation level dependent (BOLD) contrast. Settings were adjusted as follows: echo-planar images, 64 × 64 matrix; 34 slices descending; field of view [FoV] = 192 mm; repetition time [TR] = 1650 ms; echo time [TE] = 25 ms; flip angle =  $70^{\circ}$ ; slice thickness = 4.0 mm, gap size = 15%, and voxel resolution =  $3 \times 3 \times 4.6 \text{ mm}$ . Slices were acquired parallel to the intercommissural line (anterior commissure-posterior commissure). During each run of the experimental paradigm, 626 transversal functional whole brain images (including cerebellum) were recorded in descending order. Additionally, a higher resolution T1-weighted volume covering the whole brain was obtained using a magnetization-prepared rapid gradient-echo sequence in sagittal plane (176 slices, TR = 1900 ms, TE = 2.26 ms, FoV =  $256 \times 256$  mm<sup>2</sup>, flip angle 9°, matrix size =  $256 \times 256$  voxels, voxel size =  $1.0 \times 1.0 \times 1.0$  $1.0 \text{ mm}^3$ ).

# 2.3.2. Eye tracking

Simultaneously to MR acquisition, we recorded the eye movements and pupil diameter of our participants' the right eye. Data was collected at a sampling rate of 250 Hz, using an MR-compatible EyeLink 1000 eyetracker system (SR Research, Osgoode, ON, Canada) that was placed behind the MR scanner, such that participants' eye movements were recorded via the mirror mounted at the head coil. Prior to each of the three experimental runs, the eye gaze position on the monitor was calibrated using an automated nine-point calibration procedure. The calibration was accepted when the mean error was less than 0.75° of visual angle according to the corresponding validation procedure.

#### 2.4. Data analysis

#### 2.4.1. Behavioral data

To analyze participants' perception in the intensity judgment task, we calculated the proportion of trials in which the stimulus of interest was perceived as darker. Crucially, the stimulus of interest remained physically constant throughout the experiment; luminance values were manipulated only for the comparison stimulus. Thus, if brightness perception was skewed into one direction, this effect would be purely due to perceptual differences. Trials in which participants (1) failed to perform the active button press (in active trials) and/or (2) failed to report their judgment were excluded (1.95% of trials). Subsequently, the responses of "second stimulus brighter" were calculated for each participant, condition and luminance level of the second (comparison) stimulus. Logistic psychometric functions were fitted using Psignifit 4 (Schütt et al., 2016) for MATLAB (R2014a Mathworks, Sherborn, Massachusetts), implementing a maximum-likelihood estimation. Based on

the function, the thresholds and the slopes were derived for each participant and each condition. Thresholds reflect the intensity value at which participants perceive the comparison stimulus as brighter in 50% of the trials, whereas the slopes refer to the participants' ability to discriminate between the stimuli of different intensities. Here, the measure of main interest were the perceptual thresholds, because they reflect differences in the perceived brightness of the stimuli as a function of the four experimental conditions (Weiss et al., 2011). When comparing the thresholds for conditions (i.e., active and passive), a shift towards the lower intensities (to the left) indicates that, on average, the comparison stimulus was judged as brighter more often than the stimulus of interest, which always had the same luminance. Thus, a leftward shift is associated with sensory attenuation of the first visual event (stimulus of interest), since the second (comparison) stimulus was perceived as brighter in comparison. To statistically analyze the perceptual differences between the four conditions for the estimated thresholds and slopes, 2-by-2 repeatedmeasures analyses of variance (rmANOVA) were conducted separately for both measures (IBM SPSS Statistics 21). Within-subject factors were suppression (active vs. passive) and predictability (predictable vs. unpredictable) and the significance level was set at p < .05.

#### 2.4.2. Eye tracking

Eye tracking data were used to assure stimuli were properly fixated and to correlate pupil dilation and brain activation. For fixation control, we analyzed the gaze coordinates pertaining to the time when the stimulus of interest was presented. A region of interest in which participants had to fixate was defined by a circle around the center of the screen with a radius of 1.75°, thus covering the whole stimulus area while also taking into account the accuracy of the eye-tracker. After performing a drift-correction for each trial (using the gaze position recorded during the 0.3 s central fixation cue), the percentage of samples in which participants' gaze was inside of our fixation region of interest (ROI) was determined, where missing data (e.g., due to blinks) were classified as being outside the ROI. An rmANOVA with the factors action (active vs. passive) and predictability (predictive vs. unpredictable) was conducted to test for differences in gaze behavior. For the pupil size analysis, we first normalised the pupil size data. To this end, we used all the pupil traces from 500 ms prior to the onset of the stimulus of interest to its offset. We then computed mean and standard deviation across all of these data within each participant (i.e., 1 value per participant for each measure) and subtracted this mean from all data and divided the result by this standard deviation (i.e., normalized pupil data to z-scores). Note that thanks to the within-subject design, this normalization does not affect any statistics on the pupillometry data. As main trial-specific measure we used the mean of this normalized pupil data during the presentation of the stimulus of interest (1000 ms) for each trial. These trial-specific estimates of pupil size were used as a covariate in subsequent fMRI analyses to investigate the relation between pupil size and the hemodynamic response. Blinking has been shown to co-occur with button presses (van Dam and van Ee, 2005) and thus systematic differences between active and passive conditions may impact pupil size estimates. We tested for differences in blinking behavior between conditions and found no main effect of action (F(1,21) = 2.230, p = .150,  $\eta_p^2 = .096$ ) and no action\*predictability interaction (F(1,21) = 0.001, p = .988,  $\eta^2_{p < .001}$ ). There was a main effect of predictability (F(1,21) = 7.853, p = .011, $\eta^2_p = .272$ ).

#### 2.4.3. fMRI data

Preprocessing. The analysis of MRI data was performed using Statistical Parametric Mapping (SPM12, https://www.fil.ion.ucl.ac.uk/) in Matlab (R2014a Mathworks, Sherborn, Massachusetts). EPI images were realigned to the mean image to correct for head movements. Each individual's structural scan (T1 weighted) was co-registered to the functional data, segmented and normalized to the standardized Montreal Neurological Institute (MNI) template ICBM152. Using the standardized structural, all EPI images were warped to MNI space (resampled

to a voxel size  $2 \times 2 \times 2$  mm) and smoothed with a full-width at half maximum kernel ( $8 \times 8 \times 8$  mm).

1st level analyses. On the single subject level, a general linear model (GLM) was designed. The design matrix contained 11 regressors: one regressor for each first (stimulus of interest) and second (comparison) stimulus of all experimental conditions (AP\_1, AU\_1, PP\_1, PU\_1, AP\_2, AU\_2, PP\_2, PU\_2), modeling the duration of stimulus presentation (1 s). Furthermore, the cue, separated for active and passive trials, and participants' judgment, indicated by a button press, were each included as a single stick function regressor. In addition, the 6 realignment parameters entered the GLM as regressors of no-interest to control for motion-induced artifacts and a high-pass filter was set to a cut-off period of 128 s to remove slow frequencies. BOLD responses were modelled by convolving all regressors of interest with the canonical hemodynamic response function (HRF). Based on the parameter estimates, T-contrasts of the first four stimulus regressors (AP\_1, AU\_1, PP\_1, PU\_1) against implicit baseline were fed into a flexible-factorial design for the group-level analysis with the factors action (active vs. passive) and predictability (predictable vs. unpredictable). Thus, the grouplevel analysis solely focusses on the timepoint around the stimulus of interest.

2nd level analysis. As our main interest was sensory attenuation in the visual system, the occipital cortex was selected as region of interest (ROI) using a mask of the occipital lobe in the Wake Forest University (WFU) Pickatlas (Maldjian et al., 2003) based on the Automated Anatomical Labelling (AAL) atlas (Tzourio-Mazoyer et al., 2002). Subsequently to the ROI analysis, a whole brain analysis was conducted to investigate which areas outside of visual cortex might have been involved in attenuating mechanisms in the context of predictability and voluntary action.

Contrasts of interest. To investigate the main effects of action and predictability, we contrasted active and passive conditions [(PP + PU) - (AP + AU)] and predictable and unpredictable conditions [(AU + PU) - (AP + PP)], respectively. Finally, we examined the suppression\*predictability interaction ([(AU - AP) - (PU - PP)], [(AP - AU) - (PP - PU)]). For all results reported, the voxel-wise significance level was set at p < .001 (uncorrected). For the ROI analysis, small volume correction was applied. For the whole-brain analysis, a correction for errors of multiple comparisons was applied at the cluster level (FWE cluster level correction implemented in SPM12). Thus, we only report clusters which remained significant after the control for cluster-wise multiple comparisons (pFWEc < 0.05), unless specified otherwise.

Correlation with pupil data. For a second, exploratory analysis, a new GLM was set up to account for changes in participant's pupil size and its association with neural processing. In addition to the regressors from the first GLM, participant's trial-specific pupil size during the presentation of the stimulus of interest was included as a parametric regressor. Thus, each first stimulus regressor (AP\_1, AU\_1, PP\_1, PU\_1) was weighted parametrically by the individual pupil size. T-contrasts of the parametric modulators against implicit baseline were fed into a 2nd level group analysis (flexible factorial design). Here, we tested both positive and negative correlations between pupil size and BOLD response. For this analysis, we employed family wise error correction (FWE) at a significance level of pFWEc < 0.05.

### 3. Results

#### 3.1. Behavioral results

To determine the effect of action and predictability on brightness perception, we determined the luminance at which the second stimulus was judged equally bright as the stimulus of interest (point of subjective equality – PSE). To this end, we fitted psychometric functions to the fraction of judgements "second stimulus brighter" as function of log luminance of the second stimulus per individual (Fig. 2A). There was a main effect of action (F(1,21) = 4.54, p = .045,  $\eta^2_p = .178$ ) on the PSE, while

we observed no main effect of predictability (F(1,21) = 1.56, p = .226,  $\eta^2_p$  = .069) nor an action\*predictability interaction (F(1,21) = 0.266, p = .611,  $\eta^2_p$  = .013). PSEs were lower in active (M: 29.35 cd/m², SD: 1.02 cd/m²) than in passive (M: 29.46 cd/m², SD: 1.03 cd/m²; Fig. 2B).² Hence, despite substantial between-subject variance, the within-design reveals a subtle but significant tendency to perceive the stimulus of interest as darker in the active than in the passive condition. The ability to distinguish luminance levels between the two stimuli is reflected in the slope of the psychometric functions. We find no main effect of either factor (action: F(1,21) = 0.865, p = .363; predictability: F(1,21) = 0.002, p = .965); although there was an interaction (F(1,21) = 4.48, p = .047; Fig. 2C), follow-up tests did not show a significant effect for action, neither for predictable (t(21) = 1.32, p = .202) nor for unpredictable stimuli (t(21) = 1.33, p = .197).

#### 3.2. fMRI results

*ROI analysis.* For passive as compared to active conditions (passive > active), we observed bilateral activation in visual cortex, more specifically in calcarine and lingual gyri (x, y, z=10, -86, 10; T=4.82, kE = 1004; see Fig. 3A, Table 1) indicating BOLD suppression for active conditions.

To investigate the influence of predictability on sensory attenuation, conditions of predictable events were subtracted from unpredictable ones (unpredictable > predictable). No clusters of activation were found in visual cortex for this contrast. Similarly, for the action\*predictability interaction, reflecting modulatory influences of action generation and temporal predictability on brightness perception, no significant clusters of activation were observed.

As predictability was expected to have an effect, we performed an additional post-hoc analysis to examine whether our manipulation of inserting jittered delays per se worked. To this end, we split our unpredictable trials according to the delay inserted between action and the stimulus of interest (200, 450, 700, 950, 1200 ms) and formed regressors for each delay. Contrasts against implicit baseline were fed into our second level analysis, where we examined the effect of delay. We observed a main effect of delay in the visual cortex (x, y, z = 22, -80, 10; T = 4.04, E = 164, E = .017, uncorrected, with activity linearly increasing as a function of delay (Fig. 4B). This effect was similar in both active and passive conditions; even though activity was overall lower during active trials, both conditions show a linear effect of delay with the lowest activity closest to the action/cue.

Whole-brain analysis. In agreement with the ROI analysis, the whole-brain analysis revealed a suppression effect (passive > active) in visual cortex (x, y, z=10, -82, 10; T=5.19, kE=1179; Fig. 3B, Table 1). Additionally, we observed suppression in motor-related areas (primary somatosensory: x, y, z=-36, -22, 54, T=11.86, kE=2249; supplementary motor area (SMA): x, y, z=-4, -12, 52, T=8.95, kE=1186; cerebellum lobule VI: x, y, z=22, -52, -22, T=8.3, kE=1147) and in the thalamus (x, y, z=-12, -18, 2, T=7.96, kE=615). Although our main hypothesis was focused on suppression in active conditions, we also explored the opposite contrast (active > passive). Most importantly, we did not observe any activity in visual areas during the active conditions (for full results see Supplement). It should be noted that activity in visual cortex was however observed in active conditions prior to stimulus presentation, namely during the cue (see Supplement).

Also consistent with the ROI analysis, no main effect of predictability (unpredictable > predictable) was observed in the whole-brain analysis, even at a lower threshold of p < .001 uncorrected. The exploration of the contrast predictable  $\rangle$  unpredictable (Fig. 3C, Table 2) revealed a widely spread activation pattern in the cerebellum involving lobules

<sup>&</sup>lt;sup>2</sup> Note that the psychometric functions were fit to log luminance and arithmetic mean and standard deviation were also determined in log luminance and mapped back to the original values for reporting; that is, in linear units the reported mean is the geometric mean (which is the arithmetic mean in log space).

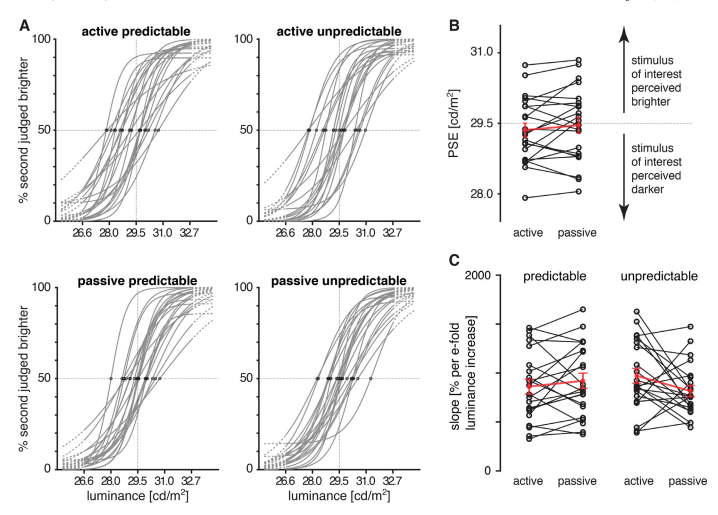


Fig. 2. Behavioral data. A. Fitted psychometric function per individual (N = 22) for the four conditions. Points of subjective equality marked with dots, extrapolation with dotted lines; x-axis logarithmically scaled. B. Point of subjective equality averaged over predictabilities, black lines: individual participants, red lines: mean and SEM. y-axis is logarithmically spaced, dotted line denotes veridical PSE. C) Slope of psychometric function for the four conditions,% 'second judged brighter' by e-fold increase of luminance; black and red lines as in panel C.

VIIa (x, y, z = 32, -82, -24, T = 5.78, kE = 3008). Furthermore, this contrast revealed clusters in the superior frontal gyrus (x, y, z = -26, 64, 10, T = 4.66, kE = 210), paracentral gyrus (x, y, z = -8, -24, 78, T = 5.02, kE = 471) and putamen (x, y, z = -16, 12, -12, T = 4.58, kE = 224). The action\*predictability interaction revealed one cluster in middle frontal gyrus (x, y, z = 42, 38, 30, T = 5.14, kE = 234), indicating lowest activation for the active unpredictable condition.

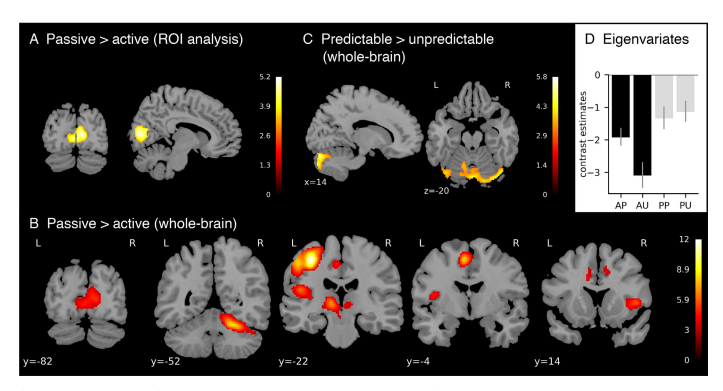
Correlation with pupil size. The analysis of pupil size revealed a main effect of action (F(1,21) = 86.01, p < .001,  $\eta_p^2 = .804$ ), such that pupils dilated more strongly in both active (AP: M = 0.24; AU: M = 0.22) as compared to passive (PP: M = -0.20; PU: M = -0.35) conditions (Fig. 5A). In addition, we observed a main effect of predictability (F(1,21) = 4.40, p = .048,  $\eta_p^2 = .173$ ) indicating larger pupil size for predictable relative to unpredictable trials. The interaction of action\*predictability showed no significant effect but a trend  $(F(1,21) = 3.91, p = .061, \eta^2_p = .157)$  thus we tested which condition was driving the difference by comparing pupil diameter between predictable and unpredictable trials in active and passive conditions. We observed that the effect was driven by the passive conditions, as in the active conditions, diameter did not differ statistically between predictable and unpredictable stimuli (t(21) = 0.24, p = .73), whereas pupils were more constricted for passive unpredictable as compared to passive predictable trials (t(21) = 2.80, p = .01).

In the parametric fMRI analysis with pupil size as a parametric modulator, a negative effect of condition was found: pupil dilation correlated

negatively with the hemodynamic response across all conditions. Thus, the smaller the pupil on a given trial, the stronger the BOLD response (Fig. 5B). This association was observed for a widespread cluster in occipital lobe (x, y, z = 3, -84, 15, T = 8.4, kE = 8135), and clusters in pre- and postcentral gyrus (x, y, z = 52, -12, 48, T = 7.5, kE = 3049; x, y, z = -38, -16, 46, T = 8.3, kE = 958) and rolandic operculum (x, y, z = 40, -16, 24, T = 5.7, kE = 43). The results showed no main effect of action, main effect of predictability or action\*predictability interaction.

#### 3.3. Eye-tracking results

Fixation analysis. If the ROI is defined by a radius of  $1.75^{\circ}$  around the center of the screen, then on average 95.6% of all samples recorded during the presentation of the stimulus of interest indicated that gaze was inside the ROI (second stimulus: 95.1%), where missing data (e.g., due to blinks) were classified as being outside the region of interest. An rmANOVA on the gaze coordinates during the stimulus of interest showed no significant differences between conditions: neither the main effect of action (F(1,21) = 0.130, p = .72), predictability (F(1,21) = 0.018, p = .77), nor the action\*predictability interaction (F(1,21) = 0.798, p = .38) reached significance. It is thus very unlikely that our behavioral and fMRI results can be explained by differences in fixation.



**Fig. 3.** Region of interest and whole-brain analyses. A. fMRI results showing BOLD suppression during active conditions (passive > active) in a cluster in visual cortex as assessed by a ROI analysis. B. fMRI results for the BOLD suppression in a network of clusters including visual cortex, somatosensory cortex and the cerebellum shown in the whole-brain analysis. C. As no main effect of predictability (unpredictable) predictable) was observed, the reverse contrast (predictable > unpredictable) was explored and revealed a cluster in the cerebellum. D. Eigenvariates, i.e., the first principal component of the time series, of the cluster with peak activity in Calcarine gyrus extracted from the ROI analysis cluster. Error bars show the standard error of the mean (SEM).

Table 1
BOLD suppression for action consequences as compared to identical but externally produced stimuli [(PP + PU) – (AP + AU)]. Coordinates are listed in MNI space. Significance threshold: p < .001 uncorrected, pFWEc = 0.05. V1, primary visual cortex; S1, primary somatosensory cortex; S2, secondary somatosensory cortex; IFG, inferior frontal gyrus; PMF, Posterior medial frontal; R, right; L, left.

Area	Cluster extent	Side	x	у	z	T	$\mathbf{k}_{\mathrm{E}}$	$P_{\mathrm{FWEc}}$
ROI Analysis								
V1	Calcarine Gyrus	R	10	-82	10	5.19	1004	< 0.001
		Lingual Gyrus	L	-2	-82	4		4.67
		Calcarine Gyrus	R	8	-74	14		4.49
Whole-brain Analysis								
S1	Postcentral Gyrus	L	-36	-22	54	11.86	2249	< 0.001
	Precentral Gyrus	L	-50	-24	48	8.41		
	Precentral Gyrus	L	-20	-18	68	4.23		
Cerebellum	Cerebellum VI	R	22	-52	-22	8.3	1147	< 0.001
	Cerebellum VI	R	20	-66	-20	4.67		
	Cerebellum Crus 1	R	46	-50	-32	4.15		
Thalamus	Thalamus	L	-12	-18	2	7.96	615	< 0.001
	Thalamus	R	8	-16	2	4.42		
	Thalamus	R	2	-26	-2	3.98		
IFG	Pars Opercularis	R	46	14	4	5.79	520	< 0.001
	Insula	R	32	24	2	5.6		
	Pars Orbitalis	R	30	24	-12	4.94		
S2	Rolandic operculum	L	-48	-24	20	6.3	1028	< 0.001
	Precentral Gyrus	L	-56	4	28	6.18		
	Precentral Gyrus	L	-38	-16	8	5.34		
PMF	SMA	L	-4	-12	52	8.95	1186	< 0.001
	MCC	L	-8	-24	48	4.34		
	SMA	L	8	6	50	4.31		
V1	Calcarine Gyrus	R	10	-82	10	5.19	1179	< 0.001
	Lingual Gyrus	L	-2	-82	10	4.67		
	Calcarine Gyrus	R	8	-74	14	4.49		

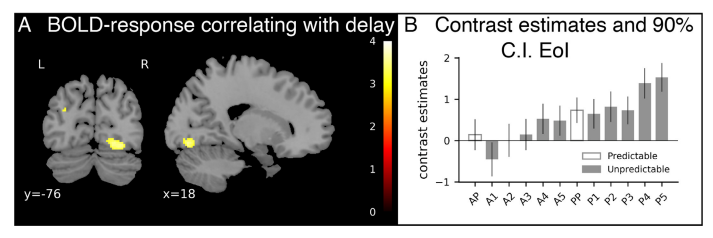
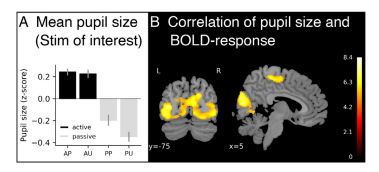


Fig. 4. Main effect of delay in a whole-brain analysis. A. BOLD activation in a cluster in visual cortex which correlated positively with delay in active and passive conditions. B. The bar plot illustrates the mean of extracted eigenvariates (i.e., the first principal component of the ROI's time series) for active and passive conditions when predictable and unpredictable (separately for all five delay increments) for the cluster peaking in visual cortex [22,-80,-10]. The error bars represent the standard error of the mean (SEM). AP = active predictable, PP = passive predictable, A\* = active, P\* = passive. The numbers 1 to 5 refer to the delay increments (1 = 1500 ms, 2 = 2000 ms, 3 = 2500 ms, 4 = 3000 ms, 5 = 3500 ms).



**Fig. 5.** Correlation of pupil size and BOLD response. A. Bar plot illustrates z-transformed pupil size averaged across participants separately for each condition. B. Activation of brain areas showing a negative linear correlation with pupil size included a network of occipital cortex, pre- and postcentral lobe and rolandic operculum.

Table 2
Processing of predictable as compared to unpredictable stimuli [(AP + PP) - (AU - PU)]. Coordinates are listed in MNI space. Significance threshold: p < .004 uncorrected, pFWEc = 0.05 R, right; L, left.

Area	Cluster extent	Side	x	y	z	T	$k_{\rm E}$	$P_{FWEc}$
Cerebellum	Crus 1, lobule VIIa	R	32	-82	-24	5.78	3008	< 0.001
	Crus 2, lobule VI	L	-2	-82	-28	5.72		
	Crus 1, lobule VIIa	R	10	-82	-26	5.2		
Superior	Superior Frontal Gyrus	L	-26	64	10	4.66	210	0.004
Frontal	Superior Medial Gyrus	L	-6	66	10	4.64		
Gyrus	Superior Medial Gyrus	L	-2	62	2	3.32		
Paracentral	Paracentral Gyrus	L	-8	-24	78	5.02	471	< 0.001
Gyrus	Paracentral Gyrus	R	10	-26	80	3.68		
	Postcentral Gyrus	L	-26	-32	72	3.58		
Dorsal	Putamen	L	-16	12	-12	4.58	224	0.003
Striatum	Olfactory Lobe	L	-20	6	-16	4.32		
	Caudate Nucleus	L	-10	16	-2	4.24		

#### 4. Discussion

Action-based sensory attenuation is a well-known effect in the auditory and tactile modalities, whereas heterogenous results have been reported in the visual domain. In this fMRI study, we examined whether action-based sensory attenuation can be observed in vision and which roles efference copy and temporal prediction mechanisms play in its generation. The results demonstrate that perceived intensity was lower and neural processing was suppressed in a network including visual, somatosensory and cerebellar brain areas in active as compared to passive conditions. There was no statistically significant effect of predictability, neither for the behavioral nor the neural data. Pupil size was larger in active as compared to passive trials and correlated negatively with BOLD response across all conditions. Overall, these data indicate that sensory attenuation and BOLD suppression are based on action-related

rather than temporal predictive mechanisms, possibly related to pupil

#### 4.1. Evidence for visual sensory attenuation: neural suppression

Our first main finding is that visual stimuli were perceived as darker and processed with less neural resources when they were actively generated, as compared to identical passively elicited stimuli. Thus, our data suggest that, beyond the auditory and somatosensory systems, sensory attenuation also occurs in the visual modality. Our ROI analysis showed reduced neural activity in visual cortex for actively generated visual stimuli. This finding is in line with previous work reporting BOLD suppression in visual cortex (Straube et al., 2017; Arikan et al., 2019; Pazen et al., 2019; Uhlmann et al., 2020, 2021; Schmitter et al., 2021) and reduced visual N1 components (Mifsud et al., 2018) for self-

generated compared to automatically presented visual stimuli. According to the notion of internal forward models (Wolpert et al., 1995; Miall and Wolpert, 1996), neural processing of action consequences is attenuated because of a cancelation of sensory information that matches the efference copy-based prediction (Blakemore et al., 1998; Shergill et al., 2013). Thus, the suppressed activity we observed in visual cortex for actively as compared to automatically generated stimuli may be a result of such action-based prediction.

An alternative framework that the BOLD suppression may be interpreted in is the "expectation by sharpening" hypothesis (Lee and Mumford, 2003) which is related to predictive coding theory (for a review see: Summerfield and de Lange 2014). While this account predicts that predictable stimuli result in reduced aggregate neural activity (Isaacson and Scanziani, 2011), it also assumes a sharper representation of predictable stimulus compared to unpredicted stimuli (Kok et al., 2012). BOLD suppression is thought to reflect a deactivation of neurons tuned away from the stimulus by which the activity of neurons coding the actual stimulus is masked. In line with this theory, sharpened sensory representations have also been reported in sensorimotor prediction for congruent action outcomes (Yon et al., 2018). Congruent action outcomes were decoded with higher accuracy than incongruent ones, while BOLD suppression was observed in voxels tuned away from the stimulus. The analysis presented in the present study cannot adjudicate between the two models. However, while the sharpening theory may explain the BOLD results, it predicts contrasting behavioral effects to what was observed here as the predictive coding framework generally assumes facilitated perception (Mumford, 1992; Friston, 2003).

In line with our hypothesis, we observed an attenuated BOLD response for actively elicited visual stimuli in primary visual cortex. Interestingly, this effect manifested itself in peripheral rather than foveal areas of V1 (Fig. 3A and B). At first glance, this finding is counterintuitive as an effect may be expected in foveal regions representing the stimulus. However, brightness judgments are strongly influenced by the luminance contrast between a stimulus and the background it is presented against, as has been evidenced by effects such as brightness induction (Heinemann, 1955) and anchoring (Gilchrist et al., 1999). Furthermore, luminance contrast has been shown to be represented not only in the lateral geniculate nucleus but also in V1 (Wiesel and Hubel, 1966; Johnson et al., 2001; Kinoshita and Komatsu, 2001; Vinke and Ling, 2020). In our experiment, luminance contrast was largest at the border between stimulus and background. Thus, we hypothesize that the cluster in primary visual cortex showing BOLD suppression may be a representation of the edge of the stimulus rather than the stimulus as a whole for which a more foveal effect would be expected. In that case, instead of using the luminance to compare the stimulus of interest and the comparison stimulus, participants might rather have based their intensity judgment on the luminance contrast between background and stimulus.

In addition to sensory attenuation in visual cortex, we observed a suppression effect in a network of brain areas involved in somatosensory and motor processing. Specifically, our data showed lower neural activity in postcentral gyrus during processing of actively as compared to passively elicited visual stimuli. This is in line with previous work (Straube et al., 2017; Arikan et al., 2019; Pazen et al., 2019; Uhlmann et al., 2020; Schmitter et al., 2021) and suggests that, in addition to processing of the sensory consequences of one's action in a given stimulus modality (here visual), also the somatosensory feedback of the action itself is attenuated. Furthermore, we observed suppressed activity during active relative to passive trials in several motor-related brain areas including SMA, cerebellum and precentral gyrus. Traditionally, these brain regions have been implicated in processes related to motor planning, execution and control (Ito, 1984; Doya, 2000; Picard and Strick, 2001). While one might expect stronger engagement (and increased activation) during active conditions, suppressed activity in these areas during active conditions has already been observed in previous work (Blakemore et al., 1998; Straube et al., 2017; Arikan et al., 2019; Pazen et al., 2019; Schmitter et al., 2021). Cerebellar areas and the SMA have also been linked to the sense of agency, more precisely to the loss of agency (David et al., 2007; Nahab et al., 2011; Sperduti et al., 2011). Given that in our experiment participants did not elicit the presentation of the stimulus in passive trials, it is possible that the stronger activation of SMA and cerebellum are a reflection of loss of agency over stimulus presentation. Alternatively, the cerebellum has been implicated in prediction error signaling for unpredictable stimuli (Schlerf et al., 2012; van Kemenade et al., 2019). It has been proposed that cerebellar activity during passive conditions (which are less predictable by nature) may indicate a prediction error resulting from the absence of efferent copy-based predictions (Pazen et al., 2019; Schmitter et al., 2021).

#### 4.2. Evidence for visual sensory attenuation: perceptual suppression

The behavioral results revealed lower perceptual thresholds in active compared to passive conditions, i.e., the stimulus of interest was perceived as darker when preceded by an active button press. This finding is in line with previous work (Cardoso-Leite et al., 2010; Vasser et al., 2019), parallels the BOLD suppression observed in the fMRI data and is well explained under the cancelation account. Presumably, due to reduced neural processing for stimuli generated by one's own action the stimulus is perceived as less intense. Yet the behavioral results presented here stand in contrast to a range of studies reporting no attenuation effect for self-initiated action consequences (Dewey and Carr, 2013; van Kemenade et al., 2016; Yon and Press, 2017; Schwarz et al., 2018) in similar experiments. It is possible that the heterogeneity of stimuli and experimental designs used in the literature, ranging from low level (Schafer and Marcus, 1973; Cardoso-Leite et al., 2010; Gentsch and Schütz-Bosbach, 2011; van Kemenade et al., 2016; Straube et al., 2017; Mifsud et al., 2018) to complex (Hughes and Waszak, 2014) and from discrete to continuously presented (Schmitter et al., 2021) stimuli, may have contributed to the ambiguous findings regarding the attenuation of visual stimuli. Furthermore, studies employed diverse measures such as speed judgments (Dewey and Carr, 2013), contrast discrimination (Roussel et al., 2013), stimulus detection (Cardoso-Leite et al., 2010; Schwarz et al., 2018), delay detections (van Kemenade et al., 2016; Straube et al., 2017; Arikan et al., 2019; Pazen et al., 2019; Uhlmann et al., 2020, 2021) and brightness judgments (Yon and Press, 2017). Further research is necessary to carefully establish the nature of stimuli and behavioral tasks leading to attenuated visual processing.

# ${\it 4.3. \ No \ attenuation \ of \ temporally \ predictable \ visual \ stimuli}$

Unexpectedly, the perceived brightness and neural processing of predictable and unpredictable stimuli did not differ significantly in our experiment. While null findings are no proof for the absence of an effect, they provide the possibility that the effect in question is unlikely to exist.

Our results suggest that the attenuation of behavior and neural processing is not confounded by temporal predictions but may rather be generated by motor-based predictions. This is in line with work from the auditory domain, showing that sensory attenuation prevails when temporal predictability (Klaffehn et al., 2019) or temporal predictability and temporal control (Harrison et al., 2021) are controlled for. Similarly, it has been reported that attenuation occurred only for touch generated by active movements and not by predictable passive movements (Kilteni et al., 2020). Combined with the present findings, this may suggest that the prediction of forward models can tolerate some degree of temporal uncertainty and cancelation of matching incoming signals is successful even at delay. However, another recent study evidenced that controlling for temporal predictability can abolish the attenuation of the auditory N1 for self-generated as compared to identical external tones (Kaiser and Schütz-Bosbach, 2018). These disparate findings may be related to differences regarding visual stimulation between

conditions (constant vs. variable) in the paradigms (Besle et al., 2004; Maddox et al., 2015).

Cortical processing of temporally predictable stimuli is often suppressed (Summerfield et al., 2008; Bendixen et al., 2009; Alink et al., 2010; Todorovic et al., 2011; Kok et al., 2012; John-Saaltink et al., 2015). Here, we manipulated the temporal predictability of visual events by varying cue-stimulus intervals, so that participants were unable to predict stimulus onset in -both actively and passively generatedunpredictable trials. However, participants were aware that a stimulus would certainly be presented on every trial. Thus, the absence of significant suppression effects for predictable as compared to unpredictable trials may be explained by the fact that the likelihood of occurrence of unpredictable stimuli was not unpredictable in our design, just the exact timing of stimulus presentation. Differences between studies might also be related to the possibility that the role of temporal predictability is not identical across modalities. It has been suggested that temporal predictability of auditory stimuli abolishes the attenuation effect, while visual components did not differ between temporally predictable and unpredictable passive stimuli (Mifsud et al., 2016b). Finally, alongside temporal predictability, further stimulus characteristics (i.e., temporal control, identity prediction) have been proposed to potentially account for the attenuation effects often observed for sensory action outcomes (Hughes et al., 2013). In the present paradigm, stimulus identity did not differ between conditions but, in contrast to active predictable conditions, participants did not have temporal control in passive conditions which may play a role in the results presented here.

# 4.4. Relationship between pupil size, visual cortex activity, and brightness perception

According to the pupillary light response effect, decreasing light intensity leads to stronger pupil dilation and increasing light intensity to stronger pupil constriction (Loewenfeld, 1958; Mathôt and Van der Stigchel, 2015). In our experiment, the stimulus of interest remained identical in luminance such that differences in pupil size could not have resulted from its physical properties. We observed that pupil size was larger in the active as compared to the passive conditions. Importantly, recent work suggested that beyond well-known influences of luminance, pupil size is also sensitive to changes in *perceived* luminance (i.e., brightness) (Laeng and Endestad, 2012). Therefore, larger pupil sizes in active trials might parallel the intensity judgment indicating that participants perceived the stimulus as darker as compared to the identical stimuli in passive trials. Thus, we propose that in addition to a behavioral intensity judgment, pupil size may be a marker providing information about perceived luminance.

An alternative explanation of the pupil size effect could be the difference in motor action and/or cognitive load between the active and passive conditions as both factors can result in pupil size changes, even in the absence of visual stimulation (Hess and Polt, 1964; Richer and Beatty, 1985; Hupe et al., 2009; Klingner et al., 2011). However, we also observed a negative correlation of pupil size with the BOLD-response in visual cortex. Thus, we find that with increasing pupil size, the activation of visual cortex decreases. This association between visual cortex and pupil size supports our hypothesis that the pupil size effect is related to visual processing of the stimulus, rather than to button press-related motion. Note, however, that the correlation was not specific to a certain visual brain area but rather generic for the whole visual cortex. One previous study, similarly, reported an inverse relationship between pupil size and the response of primary visual cortex (Bombeke et al., 2016) such that (apparently) brighter stimuli were associated with smaller pupil size and a more pronounced C1 component as compared to stimuli perceived as darker. Together, these and our findings suggest a link between pupil size and neural activity during brightness perception, however, a more mechanistic interpretation is not feasible at this point and further research is required to unravel the mechanisms underlying the link. As a side note, we find that pupil size is slightly larger when the

stimulus is predictable; this is in line with earlier findings that show larger pupil size for less uncertainty about an outcome (Preuschoff et al., 2011; Dragone et al., 2018).

#### 4.5. Effect of attention and task demand

Irrespective of (motor) prediction processes, stimulus appearance and its neural correlate can be altered by attention and task demand. The effective contrast of a visual stimulus (Carrasco et al., 2004) as well as the corresponding BOLD- (Kastner et al., 1999; Kanwisher and Wojciulik, 2000), and pupil responses (Hess and Polt, 1964; Klingner et al., 2011) can be enhanced when the stimulus is attended to or task demands are high. Furthermore, when attention is divided between two or multiple stimuli or tasks, the respective sensory brain areas are activated to a lesser degree (Loose et al., 2003). In the current experiment, active and passive conditions differed in that the active conditions comprised a motor and a subsequent visual task, whereas in the passive conditions participants focused on the visual task only. Therefore, it is possible that the enhanced brightness perception, the differences in pupil size, and the stronger BOLD-response observed in the passive conditions were a result of undivided attention towards the stimulus (brightness judgment), while participants were required to reallocate their attention in the active conditions (i.e., they had to switch from an active button press to a brightness judgment). The literature on task switching suggests that the reallocation of attention should be completed within 500 ms before the stimulus of interest is presented (Wasylyshyn et al., 2011; Worringer et al., 2019). However, in our experiment, the minimal time span between button press and onset of the stimulus of interest was 500 ms (maximum: 1500 ms). In line with this, we observed (1) no statistically significant mean difference in reaction times between active (M = 653 ms) and passive (M = 670 ms) conditions (non-parametric paired t-test: w = 87.0, p = .230, see Supplement) and (2) no correlation of pupil size and reaction times for active (Spearman correlation: rho = 0.167, p = .459) or passive (Spearman correlation: rho = 0.266, p = .232) conditions (see Supplement). Finally, two studies directly investigated the impact of attention on sensory attenuation by comparing the brain response (N1 component) to actively and passively generated stimuli (sounds) across conditions of varying allocation of attention (Timm et al., 2013; Harrison et al., 2021). The comparison of conditions with high and low attentional load (for active and passive conditions) did not reveal N1-amplitude differences, suggesting that selective attention did not diminish the attenuation effect. In contrast, the N2 component, indexing cognitive load rather than sensory processing (Folstein and Van Petten, 2007), was affected by the attention manipulation, such that the attentionally demanding conditions were associated with increased N2- amplitudes (Harrison et al., 2021). Thus, both our control analyses as well as previous work indicate that the effect of sensory attenuation unlikely results solely from differences in attentional load and task switching demands between the active and passive condi-

#### 4.6. Limitations

Motor actions, such as a button press, can introduce a variety of changes in neural processing. Given the absence of a motor baseline in the current study, we cannot completely rule out that the effect of sensory attenuation was mediated by processes unrelated to efference copy-based predictions. To preclude the possibility of spurious effects due to this disparity between active and passive conditions, a passive movement (i.e. an automatically moving button) or a motor baseline condition is needed. While the present study did not include either, sensory attenuation effects have repeatedly been shown to remain after factoring out the main effect of movement by means of a motor baseline condition both in electrophysiology (Martikainen, 2004; Baess et al., 2011; Gentsch and Schütz-Bosbach, 2011; Sanmiguel et al., 2013; Ford et al., 2014) and neuroimaging studies (Blakemore et al., 1998;

Kilteni and Ehrsson, 2020). Furthermore, several studies from our group observed sensory attenuation at the behavioral level and BOLD suppression in sensory cortices for actively elicited stimuli in experiments implementing a passive movement device (Arikan et al., 2019; Pazen et al., 2019; van Kemenade et al., 2019; Uhlmann et al., 2020). Such a passive movement device moves the participant's hand in the passive conditions and thereby imitates the hand movement performed in the active condition. Using this device, the active and passive conditions are equal in terms of the motion and differ only with regard to the presence of the efference copy as a result of the voluntary initiation of movement. Furthermore, it has been shown that even comparisons between active conditions provide some evidence for BOLD suppression specifically when the feedback of one's own hand has been provided compared to feedback from someone else's hand (Uhlmann et al., 2020). Thus, while it is possible that the attenuation effect observed here might be confounded by a lack of movement in the passive condition, we believe that it is unlikely given the experiments above which explicitly controlled for the main effect of movement.

#### 5. Conclusion

In this experiment, behavioral, neuroimaging as well as pupil size results substantiate the existence of sensory attenuation in the visual system. Stimuli elicited by a voluntary button press were perceived as darker, were associated with a suppressed BOLD response in visual cortex and led to larger pupil size. Interestingly, we demonstrate that pupil size was negatively correlated with the neural response in visual cortex. No significant effect of temporal predictability on the perception or processing of visual stimuli was observed. Our data suggest that sensory attenuation in vision likely relies more on mechanisms based on efference copies than on temporal predictions.

#### **Declarations of Competing Interest**

None.

#### Credit authorship contribution statement

Christina Lubinus: Conceptualization, Methodology, Formal analysis, Investigation, Writing – original draft, Visualization, Data curation. Wolfgang Einhäuser: Formal analysis, Writing – review & editing, Visualization, Funding acquisition. Florian Schiller: Methodology, Formal analysis, Writing – review & editing. Tilo Kircher: Writing – review & editing, Funding acquisition. Benjamin Straube: Conceptualization, Methodology, Writing – review & editing, Funding acquisition, Supervision. Bianca M. van Kemenade: Conceptualization, Methodology, Formal analysis, Investigation, Data curation, Writing – review & editing, Supervision.

#### Data and code availability statement

The data that support the findings of this study are openly available in Zenodo at: https://doi.org/10.5281/zenodo.7018607.

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#### Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.neuroimage.2022.119601.

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