

Contents lists available at ScienceDirect

Journal of Experimental Child Psychology



journal homepage: www.elsevier.com/locate/jecp

Children's perception of visual and auditory ambiguity and its link to executive functions and creativity



Mihaela Taranu^a, Marina C. Wimmer^{a,*}, Josephine Ross^b, Dávid Farkas^c, Raymond van Ee^{d,e,f}, István Winkler^g, Susan L. Denham^a

^a Cognition Institute and School of Psychology, University of Plymouth, Plymouth PL4 8AA, UK

^b Psychology, School of Social Sciences, University of Dundee, Nethergate, Dundee DD1 4HN, Scotland, UK

^c Institute of Cognitive Neuroscience and Psychology, Research Centre of Natural Sciences, Hungarian Academy of Sciences, H-1117 Budapest, Hungary

^d Biophysics, Donders Institute for Brain, Cognition and Behavior, Radboud University, 6500 GL Nijmegen, The Netherlands ^e Department of Brain and Cognition, Leuven University, 3000 BE Leuven, Belgium

^f Department of Brain, Behavior and Cognition, Philips Research, High Tech Campus, 5656 AE Eindhoven, The Netherlands ^g Department of Cognitive Science, Faculty of Natural Sciences, Budapest University of Technology and Economics, H-1111 Budapest, Hungary

ARTICLE INFO

Article history: Received 3 October 2018 Revised 25 March 2019

Keywords: Perceptual bistability Visual bistability Auditory bistability Perceptual switching Executive functions Creativity

ABSTRACT

The phenomenon of perceptual bistability provides insights into aspects of perceptual processing not normally accessible to everyday experience. However, most experiments have been conducted in adults, and it is not clear to what extent key aspects of perceptual switching change through development. The current research examined the ability of 6-, 8-, and 10-year-old children (N = 66) to switch between competing percepts of ambiguous visual and auditory stimuli and links between switching rate, executive functions, and creativity. The numbers of switches participants reported in two visual tasks (ambiguous figure and ambiguous structure from motion) and two auditory tasks (verbal transformation and auditory streaming) were measured in three 60-s blocks. In addition, inhibitory control was measured with a Stroop task, set shifting was measured with a verbal fluency task, and creativity was measured with a divergent thinking task. The numbers of perceptual switches increased in all four tasks from 6 to 10 years of age but differed across tasks in that they were higher in the verbal transformation and ambigous structure-from-motion tasks than in the

* Corresponding author.

E-mail address: marina.wimmer@plymouth.ac.uk (M.C. Wimmer).

https://doi.org/10.1016/j.jecp.2019.03.010

0022-0965/© 2019 Elsevier Inc. All rights reserved.

ambigous figure and auditory streaming tasks for all age groups. Although perceptual switching rates differed across tasks, there were predictive relationships between switching rates in some tasks. However, little evidence for the influence of central processes on perceptual switching was found. Overall, the results support the notion that perceptual switching is largely modality and task specific and that this property is already evident when perceptual switching emerges.

© 2019 Elsevier Inc. All rights reserved.

Introduction

The complex and dynamic natural world that we inhabit presents our sensory systems with a challenging problem, namely the decomposition of continuous streams of sensory input into stable and veridical representations of objects in the environment around us. One paradigm that has been used to study the perceptual strategies employed by our sensory systems is perceptual bistability, the phenomenon in which perception switches between alternative interpretations of an unchanging ambiguous stimulus. There are many qualitatively different ways to elicit perceptual bistability, and it has been reported in vision, audition, and even olfaction (Kleinschmidt, Sterzer, & Rees, 2012). The ambiguities inherent in perception have recently been brought to public attention through social media spreading of examples of perceptual ambiguity in the visual ("blue/black versus white/gold" dress) and auditory ("Yanny/Laurel") domains (Lafer-Sousa, Hermann, & Conway, 2015; Watson, 2018). These viral phenomena have highlighted gaps in our knowledge concerning why people perceive stimuli in different ways and how individual differences in the ability to perceive different interpretations of a stimulus emerge (Lafer-Sousa et al., 2015). The abilities to explore multiple ways of parsing the sensory scene and to flexibly switch between alternatives are thought to be prerequisites of cognitive flexibility, creativity, imagination, and perspective taking (Bialystok & Shapero, 2005; Wiseman, Watt, Gilhooly, & Georgiou, 2011).

Perceptual switching is typically experienced through prolonged exposure to a stimulus that itself is unchanging yet has more than one viable interpretation. The "correct" interpretation of the stimulus, therefore, is ambiguous. The ability to perceptually switch develops from 4 years of age onward in both vision and audition (Sussman, Wong, Horvath, Winkler, & Wang, 2007; Wimmer & Doherty, 2011). For example, children begin to perceive both interpretations of the duck/rabbit ambiguous figure at 4 or 5 years of age provided that children are first informed about the two interpretations (Doherty & Wimmer, 2005; Gopnik & Rosati, 2001; Mitroff, Sobel, & Gopnik, 2006; Rock, Gopnik, & Hall, 1994; Wimmer & Doherty, 2011; Wimmer & Marx, 2014) or between 5 and 9 years of age when they are uninformed (Mitroff et al., 2006). By 6 years of age, children spontaneously perceive multiple interpretations in verbal transformation tasks (e.g., repeated presentation of the nonword "flime" results in perception typically switching back and forth between "flime" and "clime") (Warren & Warren, 1966). Regarding the auditory streaming paradigm, the ability to switch depends on the pitch difference between low and high tones, with 5- to 11-year-olds requiring larger differences than adults to perceive the two dominant alternative interpretations (Sussman et al., 2007). Thus, the ability to switch interpretations develops between 4 and 6 years of age, but the specific developmental trajectory may be task and modality specific. This possibility has not yet been explored given that to date no comparison across tasks and modalities has been conducted within the same developmental sample. An important theoretical question is whether the processes underlying perceptual switching in the visual and auditory domains are domain specific or domain general (Denham et al., 2018; Pressnitzer & Hupé, 2006). In the current research, we adopted a developmental approach including 6- to 10-year-old children to allow examining the question of domain generality or specificity in visual and auditory switching.

Here we distinguished between the ability to perceive more than one interpretation (ability to switch) and the rate of ongoing perceptual switching between the alternatives given that the basic ability to switch exists. Because typical phase durations are rather long, instead of analyzing switching rate (switches per second—a very small quantity), we considered the mean number of switches within a given duration (here 60 s) as a proxy for switching rate. Preliminary evidence from the verbal transformation task suggests that even after the ability to switch has developed, there are ongoing developmental increases in switching rate between 6 and 10 years of age and between 10 years and adulthood (Warren & Warren, 1966). In addition, for ambiguous structure-from-motion tasks, the rate of switching increases between 10 years and young adulthood (Ehlers, Struber, & Basar-Froglu, 2016).

of switching increases between 10 years and young adulthood (Ehlers, Struber, & Basar-Eroglu, 2016), suggesting that the mechanisms underlying perceptual switching undergo further maturation after the ability to switch has developed. There are several reasons to expect changes in switching rate between 6 and 10 years of age. In the visual domain, even though important perceptual functions emerge during the first years of life, there

visual domain, even though important perceptual functions emerge during the first years of life, there are ongoing developments such as shifts from feature processing styles predominant until 4 years of age to increasingly global processing by 7 years of age (Nayar, Franchak, Adolph, & Kiorpes, 2015). Susceptibility to visual illusions, such as the Ebbinghaus illusion, increases with age, particularly between 4 and 10 years (Bremner et al., 2016; Doherty, Campbell, Tsuji, & Phillips, 2010), suggesting a default local processing style in younger children changing to a more global processing style in older children. Feature versus global processing in adults has been shown to affect ambiguous perception and switching rate (Gale & Findlay, 1983; Long & Toppino, 2004). Thus, switching rate might change after the ability to switch has developed due to changes in perceptual processing (Doherty et al., 2010). However, it remains unclear what specific mechanisms determine the rate of perceptual switching in children. Although inhibitory control is associated with the ability to switch (Wimmer & Doherty, 2011; Wimmer & Marx, 2014), its influence on the rate at which perception switches has not been explored in children. For instance, in adults inhibitory control measured with a Stroop task was found to be negatively related to switching rates in the auditory streaming task (Farkas, Denham, Bendixen, Tóth et al., 2016), whereas no such relation was found in the Necker cube task (Díaz-Santos et al., 2017). Thus, in adults, the relation between inhibitory control and switching rate is inconsistent. In children, the relation has not yet been investigated.

Another executive function of potential relevance is set shifting, the ability to switch between tasks or mental sets (Miyake et al., 2000), something that has been shown to relate to 6-year-olds' ability to experience alternative interpretations of ambiguous figures when prompts and biased interpretations are provided (Bialystok & Shapero, 2005). However, the evidence is again inconclusive. Set shifting does not relate to 4- and 5-year-olds' ability to perceptually switch per se (Wimmer & Doherty, 2011), nor does it relate to perceptual switching rates in ambiguous structure from motion (Chamberlain, Swinnen, Heeren, & Wagemans, 2017) or auditory streaming (Farkas, Denham, Bendixen, Tóth et al., 2016) tasks. Measuring both inhibitory control and set shifting in this study together with a range of ambiguous tasks could allow us to address some of the inconsistencies in the current literature.

Executive functions have also been found to be associated with creativity (Benedek, Jauk, Sommer, Arendasy, & Neubauer, 2014; Gonen-Yaacovi et al., 2013; Sharma & Babu, 2017). Creativity, as measured with the pattern meanings task (Wallach & Kogan, 1965), is positively correlated with the perceptual switching in the vase-face and Necker cube tasks in 16- to 18-year-olds (Doherty & Mair, 2012). A positive association is also evident in adults between divergent thinking and perceptual switching in the duck/rabbit task (Wiseman et al., 2011). These results suggest that subjective experience of perceptual switching may be related to discovering new solutions in creativity tasks (Schooler & Melcher, 1995). In contrast, Farkas, Denham, Bendixen, Tóth et al. (2016) found no correlation between divergent thinking measures of creativity and perceptual switching in the auditory streaming task in adults. Differences in perceptual tasks and measures of creativity make it difficult to draw any conclusions regarding the relation between creativity and perceptual switching, and no previous study has explored the link between creativity and perceptual switching rates in children.

In sum, we offer the first systematic investigation of perceptual switching behavior in response to ambiguous stimuli in 6-, 8-, and 10-year olds across different tasks and modalities (two visual and two auditory tasks). We anticipated that the number of switches (switching rate) would increase with age

on the verbal transformation task based on preliminary evidence (Warren & Warren, 1966), but it was unclear whether this would generalize to the other tasks tested here. If perceptual switching depends on general maturational processes and general perceptual processing styles as previously found in perceptual illusions (Bremner et al., 2016; Doherty et al., 2010; Nayar et al., 2015), then we would expect to find the same age effects across all ambiguous perception tasks. If perceptual switching is consistently related to executive functions or creativity, then this would suggest a domain-general aspect to the development of a flexible perceptual system. If there is no systematic relation, then this would speak for separate, yet generic, task-specific systems underlying perceptual switching (Denham et al., 2018).

Method

Participants

Overall, 66 children (28 girls and 38 boys) from a local state-funded primary school took part in the study. Children were predominantly Caucasian native English speakers, coming from a mix of socioe-conomic backgrounds. There were 22 6-year-olds (M = 72 months, SD = 4), 23 8-year-olds (M = 96 months, SD = 4), and 21 10-year-olds (M = 123 months, SD = 3). Participants had parental consent and gave their assent on the day of testing.

Design

Each child performed nine tasks: four perceptually ambiguous tasks (ambiguous figure, ambiguous motion, verbal transformation, and auditory streaming), a visual control task, an auditory control task, a day–night Stroop task (inhibition), a pattern meanings task (creativity), and a verbal fluency task (set shifting).

Materials and procedure

Children were seen individually in two 30-min sessions in a quiet room at their school. They sat approximately 60 cm from a Dell Latitude E6520 computer (15-inch monitor with 1600×900 resolution). Sounds were presented binaurally through headphones (Sony, MDR-NC7 B) adjusted to a comfortable volume. The experimenter sat next to the children and reminded them to focus on their task. Unlike in adult research where head movement is constrained, the children were simply asked not to move their heads and to look at the screen. Task order was counterbalanced within and between sessions with the constraint that perceptual tasks from the same modality (e.g., visual) never followed each other.

Perceptually ambiguous tasks

The *ambiguous figure task* (Jastrow, 1899) (Fig. 1, upper left panel) depicted an ambiguous line drawing $(7.7 \times 5 \text{ cm})$ subtending 7.2×4.81 (horizontal × vertical) degrees of visual angle. Participants were asked to report whether they perceived a duck or a rabbit. The body of the duck or rabbit was added to the ambiguous head during the disambiguation (training) phase.

The *ambiguous structure-from-motion task* (Wallach & O'Connell, 1953) (Fig. 1, lower left panel) consisted of a revolving cylinder with two transparent layers of 200 randomly positioned white dots over a black background moving in opposite directions with a sinusoidal speed profile (Klink et al., 2008). The cylinder subtended 5.11×6.15 (v × h) degrees of visual angle. A single dot was 12.29×12.35 arcmin (v × h) and moved with a peak angular speed of 6.96 degrees per second. Participants were asked to report whether they perceived the front face of the cylinder moving leftward or rightward. Disambiguated versions were created by dimming the luminance of either the leftward or rightward moving dots.

The sound used in the verbal transformation task (Warren & Gregory, 1958) (Fig. 1, upper right panel) was the word "life" spoken by a native female English speaker. Word duration was 0.39 s



Fig. 1. Stages for each perceptual task: (1) initial perception of an ambiguous stimulus; (2) disambiguation of two perceptual alternatives; (3) ongoing perception of the ambiguous stimulus during the test phase.

and mean pitch was 203 Hz. To create the ambiguous stimulus, the word was looped without pause 153 times. Participants were asked to report whether they heard the word "life" or "fly." For disambiguation, the words "life" and "fly" were presented once.

The auditory streaming task (van Noorden, 1975) (Fig. 1, lower right panel) was a sequence of repeating low-high-low pitched sounds (LHL_, where "L" and "H" were complex sounds and "_" indicated silence with the same duration as the sounds). The L sound was a recording of a water droplet hitting glass (a wine glass), and the H sound was a recording of a water droplet hitting ceramic (a china cup). The pitch difference between the two sounds was 28 Hz, with a mean pitch for L of 402 Hz (range = 397-408) and for H of 430 Hz (range = 428-431). Stimulus onset asynchrony (SOA: onset to onset time interval) was 150 ms. In general, a trade-off between the frequency difference (Δf) and SOA between the H and L tones determines whether people hear the sounds primarily as originating from a single source (or stream) or from two separate sources. When Δf is very small (<1 semitone [ST]), participants always perceive the sounds as coming from a single source. When Δf increases (>4 ST) and/or SOA decreases, the L and H tones are more likely to be heard as coming from two separate sources (van Noorden, 1975), and it has been found that over a very wide range of parameters both interpretations may be experienced (Denham, Gyimesi, Stefanics, & Winkler, 2013), making the stimulus ambiguous.

Participants were asked to report whether they perceived one dripping tap or two dripping taps. For disambiguation, participants were shown a picture and provided with sound of one tap and two taps. The one-tap sound contained three water droplets falling from the tap once, all of equal intensity (i.e., one LHL_ cycle). The two-tap sound contained a 19-dB intensity difference between the L and H sounds (L = 76 dB and H = 57 dB) to emphasize the perceptual popout of the L sound. Children needed to close their eyes and report what they heard (i.e., one or two taps).

All test and training stimuli can be found in the online supplementary material.

Training proceeded as follows. First, the ambiguous stimulus was presented for 15 s, after which children were asked what they saw or heard. Second, the stimulus was disambiguated until children understood the two possible categories. Third, the ambiguous stimulus was presented again (Fig. 1). Prior to the test phase, participants were instructed on which buttons they should use to indicate what they saw or heard. Testing proceeded once children understood the perceptual categories, the button assignments, and the task they were to perform.

During the test phase, children were instructed to keep the button pressed as long as they perceived one interpretation and to switch to the other button as soon as their perception changed. Button boxes were on the right and left sides of the table, with pictures of the disambiguated interpretations attached. Category position (left/right) assignment was randomized across participants except for the ambiguous motion task, in which case it was kept constant (left/right). The program recorded the button presses, from which the *number of switches* was calculated for each participant.

For each task, three 60-s blocks were separated by a pause. Participants performed the other nonperceptual tasks among the four perceptual tasks.

Perceptual control tasks

One visual control task and one auditory control task were administered to ensure that the children followed instructions.

In the visual control task, children looked at an unambiguous picture (drawing of a girl) for 60 s and a morphing animation (a horse morphed into a sheep) for 60 s. They were instructed to press a key on the keyboard whenever they saw the picture changing. Children were excluded from the analyses if they reported a change in the unambiguous picture or if they did not report any change in the morphing animation. Two participants were removed from the final analyses for failing this task (see Results).

In the *auditory control task*, children listened to two 30-s sound segments, both consisting of the two interleaved dripping taps but with one segment having the disambiguated form (intensity difference) and the other segment having the ambiguous form; segment order was randomized across participants. Participants were required to report the correct perceptual category, as instructed during training. One participant was excluded for failure in this control task (see Results).

Stroop task

Inhibition was measured using a day-night Stroop task (Simpson & Riggs, 2005). Two pictures with a day scenario and a night scenario were presented (Fig. 2); children were required to press a dark blue button in response to the day scenario and a yellow button in response to the night scenario. Participants had a pretest phase where they could practice the task. During the test phase, 16 pictures were presented in a pseudorandom order (DNNDNDDNDNDNDNDNDN, where "D" was a day picture and "N" was a night picture) and participants were asked to respond as accurately and quickly as possible. Each picture was presented for a maximum of 8 s with a 2-s intertrial interval. Accuracy and reaction times (RTs) were recorded.

To measure basic working memory components of the Stroop task (i.e., remembering what button to press for which picture), participants saw two abstract images (Fig. 2) and pressed the associated dark blue or yellow button. Otherwise, the procedure was the same as above. To isolate the inhibition component of the task, mean RT in the memory control task (memory RT) was subtracted from mean RT in the Stroop task (Stroop RT) to yield inhibition RT.

Pattern meanings task

Creativity was assessed using the pattern meanings task of Wallach and Kogan (1965), which is a divergent thinking test. Eight abstract patterns were presented individually on 20×14 -cm laminated cards (Fig. 3). Children were asked to describe "all the things you think it could be or that it reminds



Fig. 2. Images used in the Stroop task (two leftmost images) and the related memory control task (two rightmost images).



Fig. 3. Example of a pattern used in the pattern meanings task.

you of." The main variable of interest was the total number of responses generated by each participant (cTotal). Two independent raters coded 30% of the data to determine interrater reliability. Cohen's kappa test showed good agreement between the two raters, cTotal, κ = .854 (95% confidence interval [CI] = .704–1.00), *p* < .001. Because interrater agreement was very good, only one of the raters coded the remaining 70% of the data.

Verbal fluency task

Set shifting was measured using a verbal fluency task (Troyer, Moscovitch, & Winocur, 1997). Children had 60 s to name as many animals as they could think of ("Please tell me as many animals as you can") (semantic fluency) or to name words that start with the letters F. A. and S (phonetic fluency). The order of the four verbal fluency subparts (F, A, S, animals) was randomized across participants. Responses were recorded using a digital voice recorder. The responses for each of the four test subparts were transcribed, and words belonging to the same category were determined. For example, if in response to the animal naming task participants generated the words "cat," "dog," "octopus," and "fish," the first two words were considered part of one category (i.e., domestic animals) and the last two words were considered part of a second category (i.e., aquatic animals). Similarly, if in response to the letter F naming task participants responded with "fish," "finish," and "focus," the first two words were considered part of one category (i.e., words that start with the letters "fi-") and the last word was considered part of the beginning of another category (i.e., words that start with the letters "fo-"). The total number of switches between categories (VF switch) was the dependent measure for set shifting. Two independent raters coded 30% of the data to determine interrater reliability: Cohen's kappa test, VF switch, $\kappa = .847$ (95% CI = .691–1.00), p < .001. Because the agreement between the two raters was very good, only one person coded the remaining 70% of the data.

Data analysis

To examine both the ability to switch and the number of switches in the perceptual bistability tasks, two 3×4 (Age Group [6-year-olds vs. 8-year-olds vs. 10-year-olds] \times Task [ambiguous figure vs. ambiguous motion vs. verbal transformation vs. auditory streaming]) repeated-measures analyses of variance (ANOVAs) were computed with task as a within-participant variable and age group as a between-participants variable. Performance on the Stroop, pattern meanings, and verbal fluency tasks was examined using several univariate ANOVAs with age group as a between-participants variable. Bonferroni post hoc tests were used throughout. For simplicity, after reporting the results from ANOVA, whenever we used multiple pairwise *t*-test comparisions we report only the lowest *t* value for all significant effects, with plural indicated by appending an "s" (e.g., $ts \ge 6.71$, ps < .001). We used linear regression to examine factors predicting the number of perceptual switches in each task.

Results

Of the original sample, 3 participants (2 6-year-olds and 1 8-year-old) were excluded from the final analyses for failure in the auditory and visual control tasks.

Perceptually ambiguous tasks: Ability to switch

For comparison with the developmental literature in ambiguous perception, we first examined whether children aged 6 to 10 years would be able to switch at all (yes = 1, no = 0; see Fig. 4). The ability to switch was statistically different from zero across all four tasks for all age groups ($ts \ge 6.71$, ps < .001).

There was no effect of age on pass/fail switching scores, F(2, 60) = 2.181, p = .12, $\eta_p^2 = .07$; the 6-year-olds (M = .88) were equally able to switch as the 8-year-olds (M = .86, p = 1.00), who in turn were equally able to switch as the 10-year-olds (M = .95, p = .17). There was a difference in the ability to switch across tasks, F(3, 180) = 8.74, p < .001, $\eta_p^2 = .13$.

Post hoc comparisons showed that the mean proportion of participants able to perceptually switch varied for the different tasks as follows: ambiguous motion (M = .99), ambiguous figure (M = .86), auditory streaming (M = .76), and verbal transformations (M = .98), with significant differences between the ambiguous figure and ambiguous motion tasks (p = .022), between the ambiguous motion and auditory streaming tasks (p = .002), and between the verbal transformation and auditory streaming tasks (p = .002). No other differences were significant (p > .05).

Perceptually ambiguous tasks: Number of switches

The mean number of switches across participants is summarized in Fig. 5 for each of the three blocks separately for each age group.

The number of switches increased with age, F(2, 60) = 4.06, p = .022, $\eta_p^2 = .12$, where the 6-year-olds (M = 16.68) reported fewer switches than the 10-year-olds (M = 24.81, p = .025). The 8-year-olds (M = 18.72) did not differ in their number of switches from either age group (ps > .12). The number of switches also differed across tasks, F(3, 180) = 98.43, p < .001, $\eta_p^2 = .62$, where more switches occurred in the verbal transformation task (M = 47.40) than in all other tasks (ps < .001). There were also more switches in the ambiguous motion task (M = 17.10) than in both (p < .001) the ambiguous figure task (M = 8.15) and auditory streaming task (M = 7.61), which did not differ (p = 1.00). There was no Age Group × Task interaction, F(6, 180) = 1.95, p = .08, $\eta_p^2 = .06$.



Fig. 4. Proportions of children able to switch. Mean pass/fail scores for each perceptual task for the different age groups are shown. Error bars show standard errors of means. AF, ambiguous figure; AM, ambiguous motion; VT, verbal transformation; AS, auditory streaming.



Fig. 5. Numbers of switches. Mean numbers of switches for each task for different age groups are shown. Error bars show standard errors of means. AF, ambiguous figure; AM, ambiguous structure from motion; VT, verbal transformation; AS, auditory streaming.

Inhibition, creativity, and set shifting

A summary of performance on the Stroop, pattern meanings, and verbal fluency tasks is presented in Table 1.

Stroop task

Accuracy on the Stroop task was at ceiling across all age groups (6-year-olds: M = .90; 8-year-olds: M = .93; 10-year-olds: M = .95); therefore, no further statistical analyses were conducted on accuracy.

Stroop response time decreased with increasing age, F(2, 62) = 14.23, p < .001, $\eta_p^2 = .32$, particularly between adjacent ages of 6 and 8 years (p = .003) but not between 8 and 10 years (p = .17). Memory control response time also decreased with increasing age, F(2, 62) = 14.40, p < .001, $\eta_p^2 = .32$, particularly between adjacent ages of 8 and 10 years (p = .006) but not between 6 and 8 years (p = .09). There was no effect of age on inhibition (Stroop RT – memory control RT), F(2, 62) = 1.66, p = .20, $\eta_p^2 = .05$.

Pattern meanings task

The total number of responses, cTotal, did not differ with age, F(2, 62) = 2.40, p = .10, $\eta_p^2 = .07$.

Verbal fluency

Table 1

The number of category switches, VF switches, increased with increasing age, F(2, 62) = 7.95, p < .001, $\eta_p^2 = .21$, especially between 6 and 8 years (p = .008) but not between 8 and 10 years (p = 1.00).

Mean task performance for each age group.

| Task | Measure | 6-year-olds (<i>n</i> = 20) | 8-year-olds (<i>n</i> = 22) | 10-year-olds (<i>n</i> = 21) |
|------------------|-------------------|------------------------------|------------------------------|-------------------------------|
| Stroop | Stroop RT | 1444 (415) | 1086 (312) | 886 (280) |
| | Memory control RT | 1141 (292) | 968 (241) | 722 (220) |
| | Inhibition RT | 303 (347) | 117 (445) | 164 (162) |
| Pattern meanings | cTotal | 19.25 (5.24) | 24.41 (8.98) | 23.24 (8.85) |
| Verbal fluency | VF switches | 5.85 (3.56) | 9.05 (2.70) | 9.67 (3.54) |

Note. Standard deviations are in parentheses. Reaction times (RTs) are displayed in milliseconds. Inhibition RT = Stroop RT – Stroop memory control RT.

Correlations among inhbition, creativity, set shifting, and age

Correlations were calculated among the Stroop inhibitory control measure, pattern meanings task overall performance, age, and number of switches in the verbal fluency task (Table 2). Age correlated only with the number of switches in the verbal fluency task. After partialing out age, performance in the pattern meanings task correlated with the number of switches in the verbal fluency task.

Predictors of the number of perceptual switches in the ambiguous tasks

Four linear regression analyses were conducted to investigate whether the number of switches in each perceptual task was predicted by the number of switches in the other perceptual tasks, age group, inhibition (Stroop inhibitory control), set shifting (verbal fluency), and creativity (pattern meanings task). All variables were introduced in one step. There was no multicollinearity in any of the models, and the variance of the inflation factor was less than 2.0.

The regression models were significant for the number of perceptual switches in the ambiguous figure task, $R^2 = 35.1\%$, F(7, 55) = 4.25, p = .001, verbal transformation task, $R^2 = 36.4\%$, F(7, 55) = 4.49, p = .001, and auditory streaming task, $R^2 = 46.9\%$, F(7, 55) = 6.93, p < .001. The model was not significant for the number of switches in the ambiguous motion task, $R^2 = 6.8\%$, F(7, 55) = 0.575, p = .77 (Tables 3–6).

The number of switches in the ambiguous figure task was predicted by the number of switches in auditory streaming (Table 3). The number of switches in the verbal transformation task was predicted by the number of switches in auditory streaming and pattern meanings performance (Table 5). The number of switches in auditory streaming was predicted by the number of switches in the ambiguous figure and verbal transformation tasks (Table 6). There were no further significant predictors.

Discussion

This is the first investigation of perceptual switching across four different tasks (two auditory and two visual) within the same developmental sample. In line with separate strands of evidence from visual (duck/rabbit ambiguous figure task) and auditory (verbal transformations and auditory streaming) tasks (Sussman et al., 2007; Warren & Warren, 1966; Wimmer & Doherty, 2011), our current findings indicate that the ability to switch per se is present from 6 years of age across all tasks tested, thereby applying to multiple tasks and domains. However, this ability varies across individuals and tasks. In addition, the rate of perceptual switching increases between 6 and 10 years of age, and this was the case in all four tasks. Because the increase in perceptual switching rate with age is not task or domain specific, this suggests a role for generic maturational processes as well.

A long-standing question in perceptual bistability has been whether perceptual switching is caused by domain-specific or domain-general mechanisms. Our results in this developmental study are consistent with recent findings in adults (Denham et al., 2018), which showed that although many properties of perceptual switching are very similar across tasks and modalities, spontaneous perceptual switching is not centrally controlled.

 Table 2

 Correlations among age, executive functions, and creativity.

| | Inhibition | Creativity total | Verbal fluency switch |
|-----------------------|------------|------------------|-----------------------|
| Age | 21 | .20 | .42*** |
| Inhibition | _ | .08 | 01 |
| Creativity total | .13 | - | .48 |
| Verbal fluency switch | .08 | .45*** | _ |

Note. Correlations above the diagonal and partial correlations (partialing out age) below the diagonal among the Stroop, pattern meanings, and verbal fluency tasks are shown.

* p < .001.

| Deamagaiam | a maleraia | forthe | meren la au | ~£ | arrital as | : | 41-0 | a mala i arrea era | 6 | 40.01. |
|------------|------------|---------|-------------|-----|------------|---|------|--------------------|-------|--------|
| Regression | anaivsis | for the | number | OI. | switches | m | the | ampiguous | ngure | LASK. |
| | | | | | | | | | | |

| Independent variable | b | b SE | β | t |
|--------------------------|-------|------|-----|-------|
| Age group | 0.55 | 0.57 | .12 | 0.97 |
| Number of switches in AM | 0.08 | 0.09 | .10 | 0.88 |
| Number of switches in VT | -0.03 | 0.03 | 14 | -1.03 |
| Number of switches in AS | 0.44 | 0.10 | .58 | 4.53 |
| Inhibition | 0.00 | 0.00 | .08 | 0.75 |
| Creativity | 0.08 | 0.12 | .09 | 0.67 |
| Set shifting | 0.19 | 0.27 | .09 | 0.69 |
| | | | | |

Note. AM, ambiguous structure from motion; VT, verbal transformation; AS, auditory streaming. *y* < .001.

Table 4

Regression analysis for the number of switches in the ambiguous motion task.

| Independent variable | b | b SE | β | t |
|--------------------------|-------|------|------|-------|
| Age group | 0.09 | 0.86 | .02 | 0.10 |
| Number of switches in AF | 0.18 | 0.20 | .154 | 0.88 |
| Number of switches in VT | 0.02 | 0.05 | .07 | 0.45 |
| Number of switches in AS | -0.16 | 0.17 | 17 | -0.95 |
| Inhibition | 0.00 | 0.00 | 04 | -0.26 |
| Creativity | 0.10 | 0.18 | .09 | 0.57 |
| Set shifting | 0.29 | 0.41 | .11 | 0.70 |

Note. AF, ambiguous figure; VT, verbal transformation; AS, auditory streaming.

Table 5

Regression analysis for the number of switches in the verbal transformation task.

| Independent variable | b | b SE | β | t |
|--------------------------|-------|------|-----|-------|
| Age group | 3.33 | 2.22 | .19 | 1.50 |
| Number of switches in AF | -0.55 | 0.53 | 14 | -1.03 |
| Number of switches in AM | 0.16 | 0.35 | .05 | 0.46 |
| Number of switches in AS | 1.55 | 0.40 | .50 | 3.81 |
| Inhibition | -0.00 | 0.01 | 02 | -0.18 |
| Creativity | 0.94 | 0.45 | .26 | 2.06* |
| Set shifting | 0.30 | 1.14 | .04 | 0.26 |
| | | | | |

Note. AF, ambiguous figure; AM, ambiguous structure from motion; AS, auditory streaming.

, an , p < .05.

^{**} p < .001.

Table 6

Regression analysis for the number of switches in the auditory streaming task.

| Independent variable | b | b SE | β | t |
|--------------------------|-------|------|-----|-------|
| Age group | -0.10 | 0.68 | 01 | -0.14 |
| Number of switches in AF | 0.62 | 0.14 | .47 | 4.50 |
| Number of switches in AM | -0.10 | 0.11 | 10 | -0.95 |
| Number of switches in VT | 0.14 | 0.04 | .42 | 3.81 |
| Inhibition | 0.00 | 0.00 | 11 | -1.23 |
| Creativity | -0.03 | 0.14 | 03 | -0.21 |
| Set shifting | -0.25 | 0.33 | 10 | -0.77 |

Note. AF, ambiguous figure; AM, ambiguous structure from motion; VT, verbal transformation. **** p* < .001.

Overall, our findings indicate task-specific rather than domain-specific (i.e., vision and audition) differences. Perceptual switching rates in the verbal transformation task were far higher than in any of the other tasks, and this was the only task that used concrete objects that children would have encountered prior to the experiment. Language-specific processing evident early in development leads to strong representations for words. For example, from 6 months of age onward, infants can already extract word forms from natural speech (Saffran, Aslin, & Newport, 1996), and these strong representations have been associated with faster retrieval and processing for words than, for example, simple tones (Rayner & Clifton, 2009). Modeling studies have shown that switching rate increases with the strength of the competing interpretations (Mill, Bőhm, Bendixen, Winkler, & Denham, 2013). Therefore, because children in our current age range would have already encountered the words "life" and "fly" frequently (Stuart, Masterson, Dixon, & Quinlan, 1993, 1996), we suggest that prior experience, and hence stronger representations, may explain the far greater number of switches reported in the verbal transformation task than any of the other tasks. In contrast, it is highly unlikely that the children would have heard the temporal patterns caused by the dripping taps in the auditory streaming task, or would have seen randomly positioned moving dots forming a rotating cylinder, before the experiment. Therefore, the explanation also accounts for the far smaller number of perceptual switches reported in these tasks. The intermediate number of switches reported in the duck/rabbit task may similarly be accommodated. In this case, although the children were undoubtedly familiar with ducks and rabbits, they might not have seen the line-drawing versions of these animals before. Thus, although the concepts were clear, the stimulus might not have mapped very well onto the children's internal representations of ducks and rabbits, hence reducing the number of perceptual switches. Our data are consistent with previous findings that prior knowledge and familiarity with the stimuli can exert top-down influences on perceptual switching (Long & Toppino, 2004; Rock et al., 1994). Future research may want to control for familiarity (e.g., using an equally unfamiliar verbal transformation stimulus such as a nonword) and examine the effect of familiarity on switching rates.

The regression analyses showed that although there are clearly task-specific processes at work, there is also some commonality in perceptual switching rates across tasks. In the current experiment, participants' switching rate in auditory streaming mutually predicted switching rates in the ambiguous figure and verbal transformation tasks. Therefore, it could be argued that some process related to the auditory streaming task is common to the ambiguous figure and verbal transformation tasks. However, the negative relationship between ambiguous figure and verbal transformation switching rates argues against the common process explanation. Based on findings in adults of strong correlations between perceptual switching rates in different tasks in the presence of evidence that different processes underlie switching in the different tasks, Denham et al. (2018) argued for common principles implemented by independent processes. The current findings are compatible with this view. Age was not a significant predictor for switching rate in the individual tasks, providing additional support for the notion that task-specific rather than generic developmental changes are involved in early perceptual switching. The influence of more central cognitive processes on perceptual switching rates in each of the tasks was generally negligible except for the prediction of switching rate in the verbal transformation task by creativity. This also is consistent with the patchy results from the adult literature. Based on these results and similar findings in adults, we suggest that the most likely explanation is that task-specific (distributed yet generic) properties of the neural circuitry give rise to the observed task-dependent dynamics of perceptual switching (Hupé, Joffo, & Pressnitzer, 2008; Pressnitzer & Hupé, 2006). These properties differ across individuals and mature through development.

One possible explanation for the developmental increase in switching rate is that the frontoparietal brain circuitry, putatively involved in the switching process (Brascamp, Kanai, Walsh, & van Ee, 2010; Brascamp, Sterzer, Blake, & Knapen, 2018; de Graaf, de Jong, Goebel, van Ee, & Sack, 2011), undergoes a maturational process (see also Dekker, Schwarzkopf, de Haas, Nardini, & Sereno, 2017, for higher-level processing developments in 6- to 12-year-olds; Ehlers, Struber, Stadler, & Basar-Eroglu, 2006). In a recent study of the neurobiological mechanisms underlying the effect of aging on perceptual rivalry, Arani, van Ee, and van Wezel (2018) argued that the age-dependent decrease in perceptual switching

rates in the elderly may be caused by changes in neural adaptation and neural noise. Consistent with these ideas, there is evidence of a developmental increase in neuronal noise in children, argued to be an indicator of an increase in the complexity of the neural circuitry (McIntosh et al., 2010). Most computational models of perceptual switching (e.g., visual: Brascamp, van Ee, Noest, Jacobs, & van den Berg, 2006; auditory: Mill et al., 2013) depend on noise, or more precisely a balance between noise and adaptation (Shpiro, Moreno-Bote, Rubin, & Rinzel, 2009), in order to simulate the dynamics of perceptual switching (e.g., see van Ee, 2009). Therefore, one possible explanation for the developmental increase in switching rate is an increase in the complexity of the neuronal circuitry and concomitant increase in neuronal noise.

The change in switching rate with age might also be linked to changes both in the ability to sustain attentional focus on the competing interpretations and in local and global aspects of perceptual processing strategies. Sustained attention matures during primary school (Betts, McKay, Maruff, & Anderson, 2006; Lin, Hsiao, & Chen, 1999). Attentional focus over the two competing alternating interpretations in perceptual bistability influences switching rate in vision (Denham et al., 2018; Intaitė, Koivisto, & Castelo-Branco, 2014), audition (Denham et al., 2018; Farkas, Denham, Bendixen, & Winkler, 2016), and multisensory perception (Alais, van Boxtel, Parker, & van Ee, 2010; van Ee, van Boxtel, Parker, & Alais, 2009). In other words, changes in attentional focus may provide a possible explanation for changes in switching rate with age.

In addition, perceptual switching may involve processes that undergo development until late adolescence and are associated with changes in perceptual processing style (Plude, Enns, & Brodeur, 1994; Porporino, Iarocci, Shore, & Burack, 2004). Perceptual processing may move gradually from featurebased processing to global processing (van Ee, 2011). The suggestion is that when two full (global) figures compete with one another, as opposed to small local competing features (which may drive patchwise alternation but not a switch between two full figures), perceptual switching is faster. There is evidence for this change, at least in the perception of visual illusions (Bremner et al., 2016; Káldy & Kovács, 2003; Nayar et al., 2015), and processing style has been previously associated with switching rate (Long & Toppino, 2004). However, we did not investigate children's processing style or attentional focus; establishing the validity of these explanations requires further investigation.

Prior evidence relating executive and higher-level functions with perceptual switching has been inconsistent. Although inhibitory control has been shown to underlie the ability to switch in 4- and 5-year-olds (Wimmer & Doherty, 2011; Wimmer & Marx, 2014), in the current study using slightly older children we found no evidence relating inhibitory control and switching rate, consistent with previous reports for adults (Denham et al., 2018; Díaz-Santos et al., 2017; Farkas, Denham, Bendixen, Tóth et al., 2016). Similarly, set shifting was not related to switching rate, in line with previous auditory streaming studies in adults (Farkas, Denham, Bendixen, Tóth et al., 2016; Farkas, Denham, & Winkler, 2018), ambiguous structure from motion (Chamberlain et al., 2017), and the Necker cube (Díaz-Santos et al., 2017). Overall, our study provides no evidence that either executive function (inhibition or set shifting) is related to switching rate during childhood. The only association between the perceptual tasks and central measures that remained significant after controlling for the effects of age was that between verbal transformations and creativity. This could be due to the nature of the tasks, with both relying on verbal processing. Similar patchy findings have been reported in the adult literature; whereas Doherty and Mair (2012) found a relation between pattern meanings performance and perceptual switching in both vase-faces and Necker cube tasks, Denham et al. (2018) found no correlation between self-reports of creativity and switching rate in either the ambiguous structure-from-motion task or auditory streaming task. Thus, our findings are in line with the contradictory evidence in adults, suggesting that the link between creativity and perceptual switching is somewhat dubious.

This study has a number of limitations. Questions such as whether perceptual switching is related to the ability to sustain attentional focus or to perceptual processing style were not explored, leaving uncertainty over their possible role in this process. Another limitation is that creativity, set shifting, and inhibitory control were assessed using only one measure for each. Although the measures used were selected based on previous findings in the literature, in the light of the task-dependent differences in perceptual switching shown in this study, it may be advisable in future research to implement a battery of tests to characterize each of these central factors to explore the specificity of the links between perceptual switching and different creativity and executive function measures.

In conclusion, the current study showed that the rate of perceptual switching increases between 6 and 10 years of age across a range of visual and auditory perceptual bistability tasks. Similar to adults, the results indicate that perceptual switching has task-specific characteristics and that these characteristics are already evident during childhood. Overall, these results show that perceptual flexibility is an inherent property of the perceptual system from very early on. Differences across tasks show that disambiguating ambiguity in our environment is a highly task-specific skill.

Acknowledgments

M.T. was supported by funding from the European Union's (EU) Marie Curie Initial Training Network, CogNovo (FP7-PEOPLE-2013-ITN-604764). R.v.E. was supported by the EU HealthPac grant (awarded to J. van Opstal), the Methusalem program of the Flemish Government (METH/14/02) awarded to J. Wagemans, and the Research Foundation Flanders. IW was supported by the Hungarian National Research, Development and Innovation Office (NKFIH K115385). We thank Martin Coath, Nicolas Pugeault, and Chris Klink for their help in designing the auditory streaming task animations and the ambiguous structure-from-motion cylinder. We also thank the technical support office at the University of Plymouth for programming the experiment and thank the children and teachers from Montpelier primary school who made this research possible.

Appendix A. Supplementary material

Supplementary data to this article can be found online at https://doi.org/10.1016/j.jecp.2019.03. 010.

References

- Alais, D. A., van Boxtel, J. J. A., Parker, A., & van Ee, R. (2010). Attending to auditory signals slows visual alternations in binocular rivalry. Vision Research, 50, 929–935.
- Arani, E., van Ee, R., & van Wezel, R. J. A. (2018). Changes in low-level neural properties underlie age-dependent visual decision making. *Scientific Reports*, *8*, 10789.
- Benedek, M., Jauk, E., Sommer, M., Arendasy, M., & Neubauer, A. C. (2014). Intelligence, creativity, and cognitive control: The common and differential involvement of executive functions in intelligence and creativity. *Intelligence, 46*, 73–83.
- Betts, J., McKay, J., Maruff, P., & Anderson, V. (2006). The development of sustained attention in children: The effect of age and task load. *Child Neuropsychology*, 12, 205–221.
- Bialystok, E., & Shapero, D. (2005). Ambiguous benefits: The effect of bilingualism on reversing ambiguous figures. Developmental Science, 8, 595–604.
- Brascamp, J., Kanai, R., Walsh, V., & van Ee, R. (2010). Human middle temporal cortex, perceptual bias, and perceptual memory for ambiguous three-dimensional motion. *Journal of Neuroscience*, 30, 760–766.
- Brascamp, J., Sterzer, P., Blake, R., & Knapen, T. (2018). Multistable perception and the role of frontoparietal cortex in perceptual inference. *Annual Review of Psychology*, 69, 77–103.
- Brascamp, J., van Ee, R., Noest, A. J., Jacobs, R. H., & van den Berg, A. V. (2006). The time course of binocular rivalry reveals a fundamental role of noise. *Journal of Vision*, 6, 1244–1256.
- Bremner, A. J., Doherty, M. J., Caparos, S., Fockert, J., Linnell, K. J., & Davidoff, J. (2016). Effects of culture and the urban environment on the development of the Ebbinghaus illusion. *Child Development*, 87, 962–981.
- Chamberlain, R., Swinnen, L., Heeren, S., & Wagemans, J. (2017). Perceptual flexibility is coupled with reduced executive inhibition in students of the visual arts. *British Journal of Psychology*, 109, 244–258.
- de Graaf, T. A., de Jong, M. C., Goebel, R., van Ee, R., & Sack, A. T. (2011). On the functional relevance of frontal cortex for passive and voluntarily controlled bistable vision. *Cerebral Cortex*, 21, 2322–2331.
- Dekker, T. M., Schwarzkopf, D. S., de Haas, B., Nardini, M., & Sereno, M. I. (2017). Population receptive field tuning properties of visual cortex during childhood. *Developmental Cognitive Neuroscience*. https://doi.org/10.1016/j.dcn.2019.01.001. Advance online publication.
- Denham, S. L., Farkas, D., van Ee, R., Taranu, M., Kocsis, Z., Wimmer, M., ... Winkler, I. (2018). Similar but separate systems underlie perceptual bistability in vision and audition. *Scientific Reports*, *8*, 7106.
- Denham, S. L., Gyimesi, K., Stefanics, G., & Winkler, I. (2013). Perceptual bistability in auditory streaming: How much do stimulus features matter? *Learning & Perception*, *5*, 73–100.
- Díaz-Santos, M., Mauro, S., Cao, B., Yazdanbakhsh, A., Neargarder, S., & Cronin-Golomb, A. (2017). Bistable perception in normal aging: Perceptual reversibility and its relation to cognition. *Aging, Neuropsychology, and Cognition, 24*, 115–134.
- Doherty, M. J., Campbell, N. M., Tsuji, H., & Phillips, W. A. (2010). The Ebbinghaus illusion deceives adults but not young children. Developmental Science, 13, 714–721.

Doherty, M. J., & Mair, S. (2012). Creativity, ambiguous figures, and academic preference. Perception, 41, 1262–1266.

- Doherty, M. J., & Wimmer, M. C. (2005). Children's understanding of ambiguous figures: Which cognitive developments are necessary to experience reversal? *Cognitive Development*, 20, 407–421.
- Ehlers, J., Struber, D., & Basar-Eroglu, C. (2016). Multistable perception in children: Prefrontal delta oscillations in the developing brain. International Journal of Psychophysiology, 103, 129–134.
- Ehlers, J., Struber, D., Stadler, M., & Basar-Eroglu, C. (2006). Multistable perception in ten-year-old children. International Journal of Psychophysiology, 61, 377.
- Farkas, D., Denham, S. L., Bendixen, A., Tóth, D., Kondo, H. M., & Winkler, I. (2016). Auditory multi-stability: Idiosyncratic perceptual switching patterns, executive functions and personality traits. PLoS One, 11(5), e154810.
- Farkas, D., Denham, S. L., Bendixen, A., & Winkler, I. (2016). Assessing the validity of subjective reports in the auditory streaming paradigm. Journal of the Acoustical Society of America, 139, 1762–1772.
- Farkas, D., Denham, S. L., & Winkler, I. (2018). Functional brain networks underlying idiosyncratic switching patterns in multistable auditory perception. *Neuropsychologia*, 108, 82–91.
- Gale, A. G., & Findlay, J. M. (1983). Eye movement patterns in viewing ambiguous figures. In R. Groner, C. Menz, D. F. Fischer, & R. A. Monty (Eds.), Eye movements and psychological functions: International views (pp. 145–168). Hillsdale, NJ: Lawrence Erlbaum.
- Gonen-Yaacovi, G., De Souza, L. C., Levy, R., Urbanski, M., Josse, G., & Volle, E. (2013). Rostral and caudal prefrontal contribution to creativity: A meta-analysis of functional imaging data. *Frontiers in Human Neuroscience*, 7. https://doi.org/10.3389/ fnhum.2013.00465.
- Gopnik, A., & Rosati, A. (2001). Duck or rabbit? Reversing ambiguous figures and understanding ambiguous representations. *Developmental Science*, 4, 175–183.
- Hupé, J. M., Joffo, L. M., & Pressnitzer, D. (2008). Bistability for audiovisual stimuli: Perceptual decision is modality specific. *Journal of Vision*, 8(7). https://doi.org/10.1167/8.7.1.
- Intaitė, M., Koivisto, M., & Castelo-Branco, M. (2014). Event-related potential responses to perceptual reversals are modulated by working memory load. *Neuropsychologia*, 56, 428–438.
- Jastrow, J. (1899). The mind's eye. Popular Science Monthly, 54, 299-312.
- Káldy, Z., & Kovács, I. (2003). Visual context integration is not fully developed in 4-year-old children. *Perception*, 32, 657–666. Kleinschmidt, A., Sterzer, P., & Rees, G. (2012). Variability of perceptual multistability: From brain state to individual trait. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 367, 988–1000.
- Klink, P. C., van Ee, R., Nijs, M. M., Brouwer, G. J., Noest, A. J., & van Wezel, R. J. A. (2008). Early interactions between neuronal adaptation and voluntary control determine perceptual choices in bistable vision. *Journal of Vision*, 8(5). https://doi.org/ 10.1167/8.5.16.
- Lafer-Sousa, R., Hermann, K. L., & Conway, B. R. (2015). Striking individual differences in color perception uncovered by "the dress" photograph. Current Biology, 25, R545–R546.
- Lin, C. C., Hsiao, C. K., & Chen, W. J. (1999). Development of sustained attention assessed using the continuous performance test among children 6–15 years of age. Journal of Abnormal Child Psychology, 27, 403–412.
- Long, G. M., & Toppino, T. C. (2004). Enduring interest in perceptual ambiguity: Alternating views of reversible figures. Psychological Bulletin and Review, 130, 748–768.
- McIntosh, A. R., Kovacevic, N., Lippe, S., Garrett, D., Grady, C., & Jirsa, V. (2010). The development of a noisy brain. Archives italiennes de biologie, 148, 323-337.
- Mill, R. W., Böhm, T. M., Bendixen, A., Winkler, I., & Denham, S. L. (2013). Modelling the emergence and dynamics of perceptual organisation in auditory streaming. *PLoS Computational Biology*, 9, e1002925.
- Mitroff, S. R., Sobel, D. M., & Gopnik, A. (2006). Reversing how to think about ambiguous figure reversals: Spontaneous alternating by uninformed observers. *Perception*, *35*, 709–715.
- Miyake, A., Friedman, N. P., Emerson, M. J., Witzki, A. H., Howerter, A., & Wager, T. D. (2000). The unity and diversity of executive functions and their contributions to complex "frontal lobe" tasks: A latent variable analysis. *Cognitive Psychology*, 41, 49–100.
- Nayar, K., Franchak, J., Adolph, K., & Kiorpes, L. (2015). From local to global processing: The development of illusory contour perception. Journal of Experimental Child Psychology, 131, 38–55.
- Plude, D. J., Enns, J. T., & Brodeur, D. (1994). The development of selective attention: A life-span overview. Acta Psychologica, 86, 227–272.
- Porporino, M., Iarocci, G., Shore, D. I., & Burack, J. A. (2004). A developmental change in selective attention and global form perception. International Journal of Behavioral Development, 28, 358–364.
- Pressnitzer, D., & Hupé, J. M. (2006). Temporal dynamics of auditory and visual bistability reveal common principles of perceptual organization. *Current Biology*, 16, 1351–1357.
- Rayner, K., & Clifton, C. (2009). Language processing in reading and speech perception is fast and incremental: Implications for event-related potential research. *Biological Psychology*, 80, 4–9.
- Rock, I., Gopnik, A., & Hall, S. (1994). Do young children reverse ambiguous figures? *Perception*, 23, 635–644.
- Saffran, J. R., Aslin, R. N., & Newport, E. L. (1996). Statistical learning by 8-month-old infants. Science, 274, 1926–1928.
- Schooler, J. W., & Melcher, J. (1995). The ineffability of insight. In S. M. Smith, T. B. Ward, & R. A. Finke (Eds.), The creative cognition approach (pp. 97–133). Cambridge, MA: MIT Press.
- Sharma, S., & Babu, N. (2017). Interplay between creativity, executive function and working memory in middle-aged and older adults. Creativity Research Journal, 29, 71–77.
- Shpiro, A., Moreno-Bote, R., Rubin, N., & Rinzel, J. (2009). Balance between noise and adaptation in competition models of perceptual bistability. *Journal of Computational Neuroscience*, 27, 37–54.
- Simpson, A., & Riggs, K. J. (2005). Inhibitory and working memory demands of the day-night task in children. *British Journal of Developmental Psychology*, 23, 471–486.
- Stuart, M., Masterson, J., Dixon, M., & Quinlan, P. T. (1993–1996). Children's Printed Word Database. Retrieved from https:// www1.essex.ac.uk/psychology/cpwd/.

- Sussman, E., Wong, R., Horvath, J., Winkler, I., & Wang, W. (2007). The development of the perceptual organization of sound by frequency separation in 5–11-year-old children. *Hearing Research*, 225, 117–127.
- Troyer, A. K., Moscovitch, M., & Winocur, G. (1997). Clustering and switching as two components of verbal fluency: Evidence from younger and older healthy adults. *Neuropsychology*, *11*, 138–146.
- van Ee, R. (2009). Stochastic variations in sensory awareness are driven by noisy neuronal adaptation: Evidence from serial correlations in perceptual bistability. *Journal of the Optical Society of America A*, 26, 2612–2622.
- van Ee, R. (2011). Percept-switch nucleation in binocular rivalry reveals local adaptation characteristics of early visual processing. *Journal of Vision*, 11(2). https://doi.org/10.1167/11.2.13.
- van Ee, R., van Boxtel, J. J. A., Parker, A. L., & Alais, D. (2009). Multisensory congruency as a mechanism for attentional control over perceptual selection. *Journal of Neuroscience*, 29, 11641–11649.
- van Noorden, L. P. A. S. (1975). Temporal coherence in the perception of tone sequences Retrieved from. Eindhoven, Netherlands: Technische Hogeschool Eindhoven http://alexandria.tue.nl/extra1/PRF2A/7707058.pdf.

Wallach, H., & O'Connell, D. N. (1953). The kinetic depth effect. Journal of Experimental Psychology, 45, 205–217.

- Wallach, M. A., & Kogan, N. (1965). A new look at the creativity-intelligence distinction. *Journal of Personality*, 33, 348–369. Warren, R. M., & Gregory, R. L. (1958). An auditory analogue of the visual reversible figure. *American Journal of Psychology*, 71,
- 612–613. Warren, R. M., & Warren, R. P. (1966). A comparison of speech perception in childhood, maturity and old age by means of the verbal transformation effect. *Journal of Verbal Learning and Verbal Behavior*, 5, 142–146.
- Watson, C. (2018, May 17). Laurel or Yanny explained: Why do some people hear a different word? The Guardian. Retrieved from https://www.theguardian.com/technology/2018/may/16/yanny-or-laurel-sound-illusion-sets-off-ear-splittingarguments.
- Wimmer, M. C., & Doherty, M. J. (2011). The development of ambiguous figure perception. Monographs of the Society for Research in Child Development, 76(1, Serial No. 298).
- Wimmer, M. C., & Marx, C. (2014). Inhibitory processes in visual perception: A bilingual advantage. Journal of Experimental Child Psychology, 126, 412–419.
- Wiseman, R., Watt, C., Gilhooly, K., & Georgiou, G. (2011). Creativity and ease of ambiguous figural reversal. British Journal of Psychology, 102, 615–622.