## Neurobiological correlates of emotional intelligence in voice and

face perception networks

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**Abstract** 

Facial expressions and voice modulations are among the most important communicational

signals to convey emotional information. The ability to correctly interpret this information is

highly relevant for successful social interaction and represents an integral component of

emotional competencies that have been conceptualized under the term emotional

intelligence. Here, we investigated the relationship of emotional intelligence as measured

with the Salovey-Caruso-Emotional-Intelligence-Test (MSCEIT) with cerebral voice and face

processing using functional and structural magnetic resonance imaging. MSCEIT scores

were positively correlated with increased voice-sensitivity and gray matter volume of the

insula accompanied by voice-sensitivity enhanced connectivity between the insula and the

temporal voice area, indicating generally increased salience of voices. Conversely, in the

face processing system, higher MSCEIT scores were associated with decreased face-

sensitivity and gray matter volume of the fusiform face area. Taken together, these findings

point to an alteration in the balance of cerebral voice and face processing systems in the

form of an attenuated face-versus-voice bias as one potential factor underpinning emotional

intelligence.

(164 words)

Keywords:

Functional magnet resonance imaging (fMRI), emotional intelligence (EI), fusiform face area

(FFA), temporal voice area (TVA), voxel based morphometry (VBM).

Introduction

Emotions represent a major determinant of human behavior. In everyday life, they are in

large part communicated through signals from voice and face. In recent years, specialized

brain regions and networks underlying the cerebral processing of human voices and faces

have been identified. The temporal voice area (TVA; e.g., Belin et al., 2000; von Kriegstein

and Giraud, 2006; Ethofer et al., 2009; Pernet et al., 2015) and its counterpart the fusiform

face area (FFA; e.g., Kanwisher et al., 1997; Posamentier and Abdi, 2003; Kanwisher and

Yovel, 2006) are among the regions most consistently considered key functional modules in

voice and face processing, respectively. Additionally, limbic brain regions have been shown

to exhibit preferential responses to voices and faces also outside of the context of emotion

processing with a spatial overlap of voice and face-sensitivity in the amygdala (Mende-

Siedlecki et al., 2013; Pernet et al., 2015). As all these regions do not exclusively respond to

voices or faces, we use the terms voice-sensitive and face-sensitive to denote these cue

dependent preferences.

The ability to correctly interpret emotional information from voice and face is an integral

component of emotional competence. The model of emotional intelligence (EI) proposed by

Mayer and Salovey conceptualizes such competences as an ability for the "accurate

appraisal and expression of emotion in oneself and in others, the effective regulation of

emotion in self and others, and the use of feelings to motivate, plan, and achieve in one's

life" (Salovey and Mayer, 1990). According to this model, El encompasses an experiential

component (i.e., the perception and use of emotional states) and a strategic component (i.e.,

understanding and management of emotions). These domains are represented in the Mayer-

Salovey-Caruso-Emotional-Intelligence-Test (MSCEIT). It should be noted, however, that the

appropriateness of the term "intelligence" for the set of competences measured with the

MSCEIT has been questioned (for example, due to the fact that it does not represent a pure

maximum performance parameter (Petrides, 2011)). Despite this ongoing debate about the

construct of EI, this set of competences has been shown to be relevant to successful

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interaction. It was demonstrated that the MSCEIT is positively correlated with psychological

well-being (Lanciano and Curci, 2015), social competence (Brackett et al., 2006), quality of

social interaction (Lopes et al., 2004), perceived social support (Fabio, 2015) as well as

academic success (Chew et al., 2013; Lanciano and Curci, 2014), while it was negatively

correlated with loneliness (Wols et al., 2015). Finally, in incarcerated men, lower MSCEIT

scores were associated with higher scores for psychopathy (Ermer et al., 2012).

Furthermore, there is evidence not only for a link between emotional competence

encompassed as EI and effective behavioral nonverbal emotion processing (e.g., Dodonova

and Dodonov, 2012; Kniazev et al., 2013; Wojciechowski et al., 2014) but also for parallel

associations between EI and the cerebral activation during the processing of nonverbal

emotional signals (Killgore and Yurgelun-Todd, 2007; Kreifelts et al., 2010; Killgore et al.,

2013; Kniazev et al., 2013; Raz et al., 2014; Alkozei and Killgore, 2015; Quarto et al., 2016)

as well as during resting state (Takeuchi et al., 2013; Pan et al., 2014). At the structural level,

El and gray matter volume were found to be positively correlated in the insula and prefrontal

areas (Killgore et al., 2012; Tan et al., 2014).

However, it remains an open question if neurobiological correlates of the emotional

competences encompassed in the concept of EI can also be identified at the more basic

level of cerebral face and voice processing. As faces and voices constitute the most

prevalent means to express emotionally relevant information in human social communication,

it can be assumed that a high degree of emotional competence should be linked to the

sensitivity for vocal and facial cues within cerebral face and voice processing areas.

Thus, the present study aimed to clarify in a cohort of 85 healthy individuals, if and how El is

reflected in the neural responses and structure of canonical voice and face perception

networks. Based on the assumption of a link between the sensitivity to vocal and facial cues

and EI, we hypothesized a linear association of EI with voice- and face-sensitivity within the

respective voice- and face-sensitive brain regions and the amygdala as a central emotion

processing structure. We also investigated potential differential contributions of experiential

and strategic EI to cerebral voice- and face-sensitivity under the hypothesis of a stronger

association of experiential than strategic EI with cerebral voice- and face-sensitivity.

Furthermore, it was tested if beyond cerebral responses EI also modulates voice- and/or

face-sensitive functional connectivity of brain regions with El-associated voice- and/or face-

sensitivity. Here, we assumed that such EI-dependent modulations would involve the TVA for

voice-sensitive modulations of connectivity and the FFA for face-sensitive connectivity

modulations, respectively. Finally, we investigated if associations between EI and cerebral

voice- and face-sensitivity were also reflected at the structural level.

**Material and Methods** 

**Participants** 

85 healthy individuals (mean age 25.5 years, standard deviation (SD) = 3.1 years, 43 female)

participated at the Universities of Tübingen and Greifswald. All of the participants were native

German speakers and right-handed, as assessed with the Edinburgh Inventory (Oldfield,

1971). None of the individuals had a history of neurological or psychiatric illness or of

substance abuse or impaired hearing. Vision was normal or corrected to normal. None of the

individuals was taking regular medication. The study was performed according to the Code of

Ethics of the World Medical Association (Declaration of Helsinki) and the protocol of human

investigation was approved by the local ethics committee where the study was performed. All

individuals gave their written informed consent prior to their participation in the study.

Mayer-Salovey-Caruso-Emotional-Intelligence-Test (MSCEIT)

Following the MR-scanning procedure all participants were asked to complete the German

version of the Mayer-Salovey-Caruso-Emotional-Intelligence-Test (MSCEIT; Steinmayr et al.,

2011). The MSCEIT is a performance measure of emotional competences termed EI that

assesses how well people solve emotion-laden problems across several domains, including

perception, use, understanding and managing emotions. Perception and use of emotions is

subsumed as experiential EI, understanding and regulation of emotions as strategic EI.

Additionally, an overall MSCEIT score is calculated. The MSCEIT scoring was based on

consensus rating (normative sample of over 5,000 heterogeneous individuals; Mayer et al.,

2003).

Stimuli and experimental design

Two fMRI experiments were performed to localize face-sensitive (Kanwisher et al., 1997) and

voice-sensitive (Belin et al., 2000) brain areas.

The face localizer experiment included pictures from four different categories (faces, houses,

objects and natural scenes) using a block-design. All stimuli employed in the experiment

were black-and-white photographs and unknown to the participants. The face stimuli

included facial expressions which were primarily neutral with certain shifts from serious/

somewhat angry to friendly/ smiling. The house stimuli depicted different types of multilevel

buildings (e.g., brick, wooden, concrete). The object stimuli comprised common household

objects and items of clothing, while the natural scenes included pictures of different types of

panoramas (e.g., mountains, coast, river). Each block and category contained 20 stimuli.

Within blocks the stimuli were presented in random order for 300 ms interleaved with 500 ms

of fixation (1 block = 20 stimuli × (300 ms picture + 500 ms fixation) = 16 s). Eight blocks of

each category pseudorandomized within the experiment were shown separated by short ~

1.5 sec rest periods. To ascertain constant attention, a one-back task was employed in which

the participants had to press a button on a fiber optic system (LumiTouch, Photon Control,

Burnaby, Canada) with their right index finger when they saw a picture directly repeated.

Positions of repeated stimuli were randomized within blocks with the restriction that one

occurred during the first half of the block and one during the second half.

The voice localizer experiment was adapted from the seminal study by Belin et al. (2000))

and consisted of a passive-listening block design experiment with 24 stimulation blocks and

12 silent periods (each 8 s). Participants were instructed to listen attentively with their eyes

closed. The stimuli included 12 blocks of human vocal sounds (speech, sighs, laughs, cries),

6 blocks with animal sounds (e.g., various cries, gallops) and 6 blocks with environmental

sounds (e.g., doors, telephones, cars, planes). Stimuli were normalized with respect to mean

acoustic energy. The blocks were separated by 2 s of silence. Sound and silence blocks

were randomized across the experiment with the restriction that a block of silence was

always followed by at least one sound block.

Image acquisition

MRI was performed using a TRIO or VERIO 3T whole body scanner (Siemens, Erlangen,

Germany). At the TRIO structural T1-weighted images (176 slices, TR = 2300 ms, TE = 2.96

ms, voxel size: 1 x 1 x 1 mm<sup>3</sup>) and functional images (30 axial slices acquired in sequential

descending order, slice thickness 3 mm + 1 mm gap, TR = 1.7 s, TE = 30 ms, voxel size: 3 x

3 x 4 mm<sup>3</sup>, field of view 192 x 192 mm<sup>2</sup>, 64 x 64 matrix, flip angle 90°) were acquired. The

time series consisted of 336 images for the face localizer and 231 images for the voice

localizer. For correction of image distortions, a field map (36 slices, slice thickness 3 mm, TR

= 400 ms, TE(1) = 5.19 ms, TE(2) = 7.65 ms) was acquired. At the VERIO, structural T1-

weighted images (176 slices, TR = 1900ms, TE = 2.52 ms, voxel size: 1 x 1 x 1 mm<sup>3</sup>) and

functional images (34 axial slices acquired in sequential descending order, slice thickness 3

mm + 1 mm gap, TR = 2.0 s, TE = 30 ms, voxel size: 3 x 3 x 4 mm<sup>3</sup>, field of view 192 x 192

mm<sup>2</sup>, 64 x 64 matrix, flip angle 90°) were acquired. Time series consisted of 303 images for

the face localizer and 195 images for the voice localizer. A field map with 34 slices, TR = 488

ms, TE(1) = 4.92 ms, TE(2) = 7.38 ms was acquired.

Analysis of fMRI data

Data were analyzed with statistical parametric mapping software (SPM8, Wellcome Department of Imaging Neuroscience, London, UK, http://www.fil.ion.ucl.ac.uk/spm/). Preprocessing comprised the removal of the first five EPI images from each run to exclude measurements preceding T1 equilibrium, realignment, unwarping on the basis of a static field map, normalization into MNI space (Montreal Neurological Institute; Collins et al., 1994; resampled voxel size: 3 × 3 × 3 mm<sup>3</sup>) and spatial smoothing using a Gaussian filter with 8mm full width half maximum (FWHM). For the voice localizer experiment, three regressors were defined (vocal sounds [V], animal sounds [A], and environmental sounds [E]) using a box car function convolved with the hemodynamic response function (HRF) corresponding to the duration of the respective blocks of stimuli. In a similar fashion, four regressors (faces [F], houses [H], objects [O], and scenes [S]) were defined for the face localizer experiment. To remove low frequency components, a high-pass filter with a cutoff frequency of 1/128 Hz was employed. The error term was modeled as a first order autoregressive process with a coefficient of 0.2 and a white noise component to account for serial autocorrelations (Friston et al., 2002). The six motion parameters (i.e. translation and rotation on the x-, y- and z-axes) estimated during realignment were included in the models at single subject level as covariates to further reduce motion-related error variance. Voice-sensitivity was defined by the contrast V > (A, E) while face-sensitivity was defined by the contrast F > (H, O, S). The individual contrast images were calculated and statistically evaluated at the group level in a random-effects analysis using one-sample t-tests to define the face-sensitive fusiform face area (FFA) and the voice-sensitive temporal voice area (TVA) as functional regions of interest (ROI) for subsequent analyses. Statistical significance of activations was assessed at p < 0.001, uncorrected at voxel level, and FWE correction for multiple comparisons at cluster level with p < 0.05. For the definition of the FFA the fusiform gyrus was defined as a priori anatomical ROI and the temporal gyri and the temporal pole for the definition of the TVA. For the definition of the functional ROIs (i.e., FFA and TVA), FWE-cluster level correction was performed across these a priori anatomical ROIs using small volume

correction (SVC; Worsley et al., 1996) (see Tab. 1). Additionally, the amygdala served as

anatomically defined a priori ROI. The Automated Anatomic Labeling (AAL) toolbox

implemented in SPM (Tzourio-Mazoyer et al., 2002) was used for the definition of the

amygdala in MNI space.

The associations between MSCEIT scores and individual cerebral face- and voice-sensitivity

were investigated using linear regression analyses. We first calculated hypothesis-based

ROI analyses centered on FFA, TVA and amygdala (p < 0.001 at voxel level with FWE

correction [p < 0.05] for multiple comparisons across the respective ROI volume). In view of

the lack of previous studies on the association of emotional competences or intelligence and

cerebral voice- and face-sensitivity, the ROI analyses were complemented with an

explorative whole-brain analysis (p < 0.001 at voxel level with FWE correction [p < 0.05] for

multiple comparisons at cluster level). In clusters with a significant association of MSCEIT

scores and cerebral face- and/or voice-sensitivity, mean contrast estimates were extracted

and the regression coefficients obtained for the association with experiential and strategic El

were tested for differences when using the matrix approach described by A. Paul Beaulne

(http://www.spsstools.net/Syntax/RegressionRepeatedMeasure/CompareRegressionCoeffici

ents.txt). Effect sizes for such differential associations with experiential and strategic EI are

given as Cohen's d. Additionally, validation analyses were performed where face- and voice-

sensitivity were defined as minimum difference contrasts (i.e., F – max(H, O, S), V – max(A,

E)) to ensure that observed associations with MSCEIT scores unequivocally originated from

specific modulations of cerebral responses to voices or faces, respectively. Finally, another

validation analysis was performed comparing the regression coefficients obtained for the

association of MSCEIT scores with the responses to voices, or faces, respectively, with

those regression coefficients obtained for the associations of MSCEIT scores and each of

the other stimulus classes within the respective localizer experiment. Age, gender and the

MRI scanner in which the experiments were performed were included as covariates in all

group analyses. As it has been repeatedly demonstrated that the TVA is not uniform but

contains several distinct peaks of voice-sensitivity with assumedly distinct functional profiles,

all significant functional effects observed in the TVA in the present study were spatially

referenced by their Euclidian distance in MNI space to the TVA voice-sensitivity peaks in the

present study (see Tab. 1 and Supplementary Tab. 1) and to the distinct voice-sensitivity

peaks/ clusters observed in the seminal study by Belin et al. (2000)) as well as the recent

large scale study by Pernet et al. (2015)), which included 218 individuals.

To minimize the risk of missing significant associations between cerebral face-sensitivity and

MSCEIT scores occurring in other major face processing areas (i.e., the posterior superior

temporal sulcus [pSTS] and the occipital face area [OFA]), complementary ROI analyses

focused on these regions were performed. For the definition of the face-sensitive areas in the

pSTS, the superior and middle temporal gyri and the angular gyrus were selected as the a

priori anatomical ROI, and for the definition of the OFA, the inferior occipital gyrus was

selected as the a priori anatomical ROI. Parallel to the definition of TVA and FFA, the

statistical significance of face-sensitive activations in these anatomical a priori ROIs was

assessed with a voxel-wise threshold of p < 0.001 and FWE correction for multiple

comparisons at cluster level with p < 0.05 across the anatomical ROIs using SVC (see

Supplementary Tab. 2).

Psychophysiological interaction analyses (PPI, Friston et al., 1997) were performed to

assess the relationship of EI and voice-/ face-sensitive modulations of functional connectivity

(FC). Areas with significant associations between El and voice-/ face-sensitivity were

selected as seed regions for the PPI analyses. In these analyses, the time-course of the

BOLD response, based on a sphere with a radius of 3 mm around the peak-activation voxel

within the respective seed region of the contrast of interest (e.g. V - (A,E)) was extracted in

each individual participant and was defined as the physiological variable. The

psychophysiological interaction was calculated as the product of the deconvolved activation

time course (Gitelman et al., 2003) and the vector of the psychological variable (i.e., the

voice- or face-sensitivity defining contrasts V - (A,E) and F - (H,O,S), respectively). The

relationships between EI and individual face- and voice-sensitive FC-modulations (i.e., PPI

estimates) were, again, investigated using linear regression analyses. The sequence of

group level analyses was parallel to the analyses performed for the cerebral activation

patterns as described above. Again, differential associations of experiential and strategic El

with voice-/ face-sensitive connectivity patterns were tested post-hoc by comparing the

respective regression slopes using Beaulne's matrix procedure.

Voxel-based morphometry (VBM)

The VBM8 toolbox (http://dbm.neuro.uni-jena.de/vbm.html) implemented in SPM8 was used

for the preprocessing of the T1-weighted structural images applying the default settings: The

images were segmented into gray matter, white matter and cerebrospinal fluid, DARTEL-

normalized to MNI space (resampled voxel size: 1.5 × 1.5 × 1.5 mm³) and modulated with

the nonlinear components enabling the comparison of the absolute amount of tissue

corrected for individual brain sizes. The gray matter segments were smoothed with a

Gaussian kernel (8 mm FWHM). The association between MSCEIT scores and gray matter

volume was then analyzed as described for the functional images. Primarily, analyses

focused on regions where voice- and/ or face-sensitivity were associated with El. In these

regions, mean gray matter volumes were extracted and analyzed. Additionally, the

regression on MSCEIT scores was performed in an a priori ROI including the insula, the

orbitofrontal and the anterior mediofrontal cortex (Killgore et al., 2012) as defined using the

AAL toolbox with a voxel-wise threshold of p < 0.001, uncorrected, and FWE correction (p <

0.05) at cluster level for multiple comparisons within the ROI.

Results

Participant sample data

In our study population of 85 healthy individuals (mean age 25.5 years, standard deviation

[SD] = 3.1 years, 43 female) the mean overall MSCEIT score was 105.7 (SD = 13.1), the

mean experiential El subscore was 104.3 (SD = 14.8) and the mean strategic El subscore

was 105.1 (SD = 10.9). The covariates age, gender and MRI scanner were not substantially

correlated with the MSCEIT scores (all p > 0.05, all abs(r)  $\leq$  0.09).

fMRI analysis

Cerebral activation

In the ROI analyses, a significant positive linear association between individual MSCEIT

scores and cerebral voice-sensitivity was detected in the left amygdala (Fig. 1; peak MNI

coordinate: -18 -3 -18, cluster size 54 mm<sup>3</sup>, t = 3.4, p<sub>FWEcorr</sub> = 0.01) but not in the other ROIs.

At whole brain level, an additional significant positive linear relationship between MSCEIT

scores and voice-sensitivity was observed in the left anterior insula extending into the inferior

frontal gyrus (left insula/ IFG; see Fig. 1, Tab. 2; p<sub>FWEcorr</sub> = 0.045).

--- Figure 1 about here ---

In both regions, i.e. left insula/ IFG and left amygdala, the relationship between MSCEIT

scores and voice-sensitivity was driven by increased responses to voices in individuals with

greater MSCEIT scores (r ≥ 0.32, p ≤ 0.003) whereas there was no effect of MSCEIT scores

on cerebral responses to animal and environmental sounds (all abs(r)  $\leq$  0.16, all p > 0.05;

Fig. 2A and 2B).

The relationship between MSCEIT scores and voice-sensitivity can also be further illustrated

by splitting the participants into three subsamples with regard to their MSCEIT scores (i.e.,

"low" El [n = 28, mean MSCEIT score 90.9 (SD = 1.6)], "average" El [n = 29, mean MSCEIT

score 107.0 (SD = 0.5)] and "high" EI [n = 28, mean MSCEIT score 119.2 (SD = 1.1)]). Only

for individuals with "high EI" the left insula/ IFG exhibited significant voice-sensitivity (t = 3.6,

p = 0.001) whereas in "average" and "low" El individuals this region did not exhibit voice-

sensitivity (both abs(t)  $\leq$  1.3, both p > 0.05; see Fig. 2C). In contrast, in the left amygdala only

individuals with "low" EI did not exhibit significant voice-sensitivity (t = 1.4, p > 0.05) while the

other two groups did comparably do so (both  $t \ge 4.4$ , both p < 0.001; see Fig. 2D).

--- Figure 2 about here ---

A significant negative relationship of MSCEIT scores and face-sensitivity was found in the

right FFA at ROI level (see Fig. 3, Tab. 2, p<sub>FWFcorr</sub> = 0.02) whereas we did not observe any

significant associations between MSCEIT scores and face-sensitivity within the other ROIs or

at the whole brain level.

--- Figure 3 about here ---

The negative correlation between MSCEIT scores and face-sensitivity (i.e., the differential

response to faces as compared to the other classes of stimuli) in the right FFA was driven by

a combination of decreased responses to faces and increased responses to the other

stimulus classes with increasing MSCEIT scores. Statistical post-hoc decomposition

indicated, however, that these effects were non-significant when analyzed separately (all

 $abs(r) \le 0.2$ , all p > 0.05; see Fig. 4A). In the group comparison, individuals with "low" El

exhibited greater face-sensitivity in the right FFA than the "average" and "high" El groups

(both  $t \ge 2.2$ , both  $p \le 0.04$ ; see Fig. 4B).

--- Figure 4 about here ---

Moreover, in the left insula/ IFG and right FFA, experiential EI contributed more strongly to

the observed association between EI and voice- and face-sensitivity, respectively, than did

strategic EI (both  $t \ge 2.0$ , both  $p \le 0.03$ , both  $d \ge 0.43$ ; see Suppl. Fig. 1A and 1B). This

effect was marginally significant in the left amygdala (t = 1.6, p = 0.06, d = 0.35).

Validation analyses using minimum difference contrasts evidenced that all observed

associations between EI and voice- and face-sensitivity were driven by increased responses

to voices (left insula/ IFG: r = 0.40, p < 0.001; left amygdala: r = 0.33, p = 0.002) and weaker

responses to faces (right FFA: r = -0.32, p = 0.003). Further validation analyses targeting

differential associations between MSCEIT scores and the cerebral responses to the different

stimulus categories confirmed the regression coefficient difference in the left insula/ IFG for

voice vs. animal sounds (t = 4.2, p < 0.001, d = 0.93) and voice vs. environmental sounds (t

= 3.6, p < 0.001, d = 0.78). In the left amygdala, differences for voice vs. animal sounds (t =

3.1, p < 0.005, d = 0.68) and voice vs. environmental sounds (t = 2.3, p < 0.05, d = 0.50)

were corroborated. Also the regression coefficient differences in the right FFA for faces vs.

houses (t = -3.7, p < 0.001, d = 0.81), faces vs. objects (t = -3.1, p < 0.005, d = 0.69) and

faces vs. scenes (t = -3.3, p < 0.005, d = 0.73) were statistically significant.

The complementary ROI analyses targeting potential associations between MSCEIT scores

and face-sensitivity in other major face processing areas (i.e. the pSTS and the OFA, see

Supplementary Tab. 2) did not produce any significant results: the voxel-wise statistical

threshold of p < 0.001 was not reached in any voxel included in the analyses.

Voice- and face-sensitive modulations of functional connectivity

Psychophysiological interaction (PPI) analyses using areas with significant associations

between MSCEIT scores and voice-/ face-sensitivity as seed regions demonstrated a

significant positive linear association of MSCEIT scores with voice-sensitive FC increases

between the left insula/ IFG and a region in the middle part of the right TVA (Tab. 3 and Fig.

5, p<sub>FWEcorr</sub> = 0.049). Spatial comparison revealed that the observed PPI effect is situated

closest to the middle STS voice-sensitivity peaks found in the present and previous studies

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(see Supplementary Tab. 1). Using the left amygdala and the right FFA as seed regions no

significant relationships between MSCEIT scores and voice-sensitive (left amygdala) or face-

sensitive (right FFA) FC modulations were observed (Tab. 3).

--- Figure 5 about here ---

Voxel based morphometry

The relationship between EI and voice-sensitivity in the left insula/ IFG and face-sensitivity in

the right FFA was paralleled by concurrent associations between MSCEIT scores and gray

matter volume in these areas (i.e., left insula/ IFG: r = 0.19, p = 0.04, one-tailed; right FFA: r

= -0.27, p = 0.01; see Fig. 6). No such correlation was observed in the left amygdala (r =

0.15, p > 0.05). Apart from the functional ROIs, a positive correlation of MSCEIT scores and

gray matter volume was observed in the right OFC (see Tab. 4 and Fig. 7, p<sub>FWEcorr</sub> = 0.04).

Differential relationships between gray matter volume and experiential versus strategic El

were not observed (all abs(t) < 1.9, all p > 0.05, d < 0.39).

--- Figures 6 and 7 about here ---

**Discussion** 

Our findings demonstrate that the complex set of emotional competences termed EI is linked

to the cerebral processing of faces and voices already at the level of sensory voice- and

face- as well as limbic emotion-processing areas rather than in higher cognitive brain

regions. Notably, this link was observed irrespective of an experimentally inherent cognitive

focus on emotional information as this is the case e.g. in an emotion evaluation task.

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The first main finding, a positive relationship between EI and voice-sensitivity in the anterior insula extending into the IFG, fits several neuroimaging studies which reported an association of insular responses and El during the processing of emotional cues (i.e., emotional faces; Killgore and Yurgelun-Todd, 2007; Alkozei and Killgore, 2015; Quarto et al., 2016). These latter findings have been discussed with reference to the somatic marker hypothesis (Damasio, 1994), in which especially the anterior part of the insula plays a major role as a neural structure integrating the emotional salience of a stimulus and the individual's own affective state (e.g., Phillips et al., 2003) during decision making. However, as the relationship between EI and insular voice-sensitivity exists outside the context of explicit emotion processing, it may be necessary to interpret this preference for one of the major carrier signals of emotionally relevant information in human social life in relation to the general salience processing function of the insula (e.g., Bartra et al., 2013; Hayes et al., 2014) within the so-called salience network (Seeley et al., 2007). This notion of increased salience of human voices in emotionally competent individuals is also consistent with the finding of a corresponding relationship of El and voice-sensitivity in another central part of the network subserving salience processing, namely the amygdala (e.g., Adolphs, 2010; Fernando et al., 2013). Following this conception, the El-associated voice-sensitive FC increase between the insula and right TVA may be a correlate of more pronounced parsing of vocal signals for emotionally and socially relevant information. This observation potentially reflects a neural mechanism of how effective voice processing supports emotional competences. It dovetails with current findings of increased FC between the TVA and the anterior insula/ IFG during the task-irrelevant extraction of emotional information from vocal cues (Frühholz and Grandjean, 2012) as well as decreased FC between these areas in psychiatric conditions with perceptual deficits for vocally communicated emotional information (i.e., schizophrenia [Kantrowitz et al., 2015] and autism spectrum disorders [Abrams et al., 2013]). The spatial proximity of the El-associated voice-sensitive FC increase to the middle STS voice-sensitivity peaks observed in the present and previous studies (Belin et al., 2000; Pernet et al., 2015) indicates that this effect may reflect voice-specific acoustic

processing (e.g., Kriegstein and Giraud, 2004; Charest et al., 2013; Latinus et al., 2013; Giordano et al., 2014).

Surprisingly, the FFA, as one of the most central modules of the cerebral face processing network exhibited a negative relationship between its sensitivity for faces and MSCEIT scores. Functionally, this might be explained by greater neural efficiency during general face processing in emotionally competent individuals as has been suggested for emotional facial expressions (Killgore and Yurgelun-Todd, 2007). Yet, the decreased face-sensitivity of the FFA in highly emotionally competent individuals is structurally mirrored by a reduction in gray matter volume. While a negative correlation between EI and gray matter volume in the fusiform gyrus has previously been reported in a large scale study (Tan et al., 2014), such findings stand in contrast to training (e.g., Kreifelts et al., 2013) and learning (e.g., Gimenez et al., 2014) studies in which increased neural efficiency was not only accompanied by decreased cerebral activation but also by an increase in gray matter volume. Specifically, a decrease in FFA responses to emotional cues following an emotion communication training was associated with increased gray matter volume in this region (Kreifelts et al., 2013). Nevertheless, the novel and somewhat surprising finding of a negative relationship between emotional competence and FFA gray matter volume is validated by positive correlations between gray matter volume in the OFC and insula and El observed in the very same analysis. These results correspond with those of previous studies in healthy individuals (Killgore et al., 2012; Tan et al., 2014) and individuals with brain injuries (Barbey et al., 2014) and converge with current concepts of these structures in emotional perception, evaluation and decision making (Gutierrez-Cobo et al., 2016). Thus, further discussion is warranted. Alternatively, the opposite associations between MSCEIT scores and voice-sensitivity on the one hand and face-sensitivity on the other might be an indicator of a reduced visual bias in emotionally competent individuals. Generally, adults exhibit a visual preference during the processing of audiovisual signals, both in the abstract (Robinson and Sloutsky, 2004) but also in the emotional (i.e., facial and vocal expressions; Santorelli, 2006) domain. Such a bias does not exist in children or is even reversed towards the auditory modality (Robinson

and Sloutsky, 2004). One could hypothesize that those individuals who develop a high

degree of emotional competence in the sense of effective emotional learning during

childhood also develop a weaker visual bias, or in other words, process voices and faces in a

more balanced manner. Again, it is in line with our results that the experiential domain of El

should be more strongly associated with such altered voice and face processing than the

strategic domain of El.

The idea of a reduced visual bias in voice and face processing therefore appears as a very

interesting topic for future research on the bases of El. If it could be demonstrated that El is

indeed positively correlated with a reduced face bias at the behavioral level, this would open

up the avenue to a line of research investigating the possibility of fostering the development

of emotional competences through "voice bias trainings", with the potential to clarify a causal

relationship. Based on findings that voice and face processing are at least partially

genetically determined (Brown et al., 2012; Koeda et al., 2015), genetic imaging might be

employed to elucidate the potential genetic foundations of alterations in voice and face

processing associated with El. Jointly, such investigations could clarify if the increased

cerebral voice-sensitivity and reduced face-sensitivity are the consequence or rather the

cause of well-developed emotional competences. Future studies are needed to determine if

certain vocal and facial features are more strongly associated with EI than others in analogy

to the observation of differential associations between different emotional facial expressions

and EI in the anterior insula (Quarto et al., 2016).

Finally, one should keep in mind that constructs of EI share a considerable amount of

variance with other interindividual characteristics (e.g. personality and cognitive ability;

Joseph and Newman, 2010; Joseph et al., 2015). This precludes any definitive conclusions

about the specificity of the observed associations with regard to the construct of EI and is

reflected in the relatively low incremental validity of El constructs over measures of

personality and cognitive ability (e.g., Joseph and Newman, 2010). Thus, to determine El-

specific neural correlates, large scale studies are needed incorporating all of the partially co-

linear interindividual characteristics.

Limitations

Due to the fact that the canonical voice and face localizer experiments employed in the

present study vary with regard to attentional load and processing effort (i.e., passive listening

vs. one-back working memory task), an influence of these factors, e.g., in the form of voice-

sensitive automatic attention and evaluation processes or face-sensitive reduced effort in the

working memory task, on our results can't be excluded and certainly further research on the

relations of EI with attention and processing effort during voice and face perception is

warranted.

In addition, the auditory and visual stimuli are not perfectly comparable with regard to

emotional content and stimulus-inherent dynamics. As the potential emotional content of the

face and voice stimuli has not been explicitly quantified and compared, it cannot be ruled out

completely, that the emotional information incidentally included in the stimulus material might

have contributed differently to the TVA (as determined by the voice localizer) than to the FFA

(as determined by the face localizer) activations. Moreover, the visual stimuli are presented

as static pictures, whereas a dynamic is inherent in auditory stimuli. Even if the FFA

responds to static and dynamic faces in the same way (Pitcher et al., 2011), an influence of

this factor can't be fully excluded. These potential interfering factors should be addressed in

more detail in further experiments evaluating a shift of voice- and face sensitivity related to

EI.

On the other hand, the employment of these canonical designs allows us to relate our

findings directly to the extensively investigated voice and face processing networks

described on the basis of commonly used localizer experiments. Additionally, it appears

more than unlikely that the opposite relationships between voice- and face-sensitivity and

MSCEIT scores should be solely due to differences in attention, effort and stimulus material

as these functional associations were found to be accompanied by concurrent associations

between gray matter volume and El which also exist outside the context of the respective

experimental settings.

Conclusion

In the present study, we demonstrated that emotional competence measured as EI and

cerebral voice- as well as face-sensitivity are inversely associated in healthy individuals. The

concordant positive correlations between EI and the voice-sensitivity and gray matter volume

of the right anterior insula as well voice-sensitively increased functional connectivity between

the insula and the TVA can be interpreted as correlates of a generally increased salience of

voices in emotionally competent individuals. This notion is further supported by the

comparable positive association of El and voice-sensitivity in the left amygdala. In contrast,

the right FFA, as one central functional module of the cerebral face processing system,

exhibits a negative correlation between EI and face-sensitivity as well as gray matter volume.

Together, these results indicate a shifted balance of voice and face processing systems in

the form of an attenuated face-versus-voice bias as a neural correlate of El. The present

study offers a starting point for future research projects aimed to further elucidate the direct

behavioral relevance of the observed El-associated voice-versus-face processing shift within

the framework of more elaborate voice-face processing studies. Moreover, it remains an

open question for future studies if the alteration in the balance of cerebral voice and face

processing is rather a genetically determined precondition for the development of a high

degree of emotional competence, or if it is the consequence of learning mechanisms

underlying the individual development of emotional competence.

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## Tables

**Table 1.** Functional regions of interest (ROI): voice-sensitive area in the temporal lobe (temporal voice area, TVA) and face-sensitive area in the fusiform gyrus (fusiform face area, FFA) in 85 healthy individuals.

Voice- and	Peak MNI coordinate	t peak voxel	Cluster size	p value
face- sensitive	(x y z)	(df = 81)	(mm³)	
areas				
Right TVA	60 -12 -3	20.2	32670	<0.001
	57 -21 0	17.9		
Left TVA	-57 -18 -3	19.2	32400	<0.001
	-60 -30 3	16.6		
Right FFA	42 -48 -21	15.0	2214	0.002
Left FFA	-42 -54 -21	9.6	1701	0.005

**Table 2.** Linear associations between EI and cerebral voice- and face-sensitivity.

Peak MNI coordinate region	Peak	MNI	coordinate	t peak voxel	Cluster	size
	(x y z)			(df = 80)	(mm³)	
Voice-sensitivity						
L insula / L inferior frontal gyrus	-36 15	6		4.5	2511 *	
partes triangularis et opercularis						
R thalamus	6 -6 0			3.9	432	
R inferior frontal gyrus pars	54 24	15		3.6	297	
triangularis						
R putamen	30 18	0		3.5	135	
R superior temporal gyrus	66 -39 18		3.5	216		
Face-sensitivity						
L precuneus / L superior parietal	-15 -57 42		-4.0	351		
gyrus						
R fusiform gyrus / R cerebellum	36 -60	-18		-3.7	432 **	

The initial whole-brain analysis was performed at a threshold of p < 0.001 at voxel level with FWE correction [p < 0.05] for multiple comparisons at cluster level; significant results are marked with \*. ROI analyses centered on FFA, TVA and amygdala were performed at a threshold p < 0.001 at voxel level with FWE correction [p < 0.05] for multiple comparisons across the ROI volume based on small volume correction (SVC) and significant results are marked with \*\*. Only clusters  $\geq$  135 mm³ (voxel size 3 x 3 x 3 mm³) are reported. df = degrees of freedom, R = right, L = left.

Table 3. Linear associations between MSCEIT scores and voice- as well as face-sensitive modulations of functional connectivity (FC). Psychophysiological interaction (PPI) analysis.

Peak MNI coordinate region	Peak	MNI	coordinate	t peak vox	el Cluster size
	(x y z)			(df = 80)	(mm³)
Seed region 1: L inferior frontal					
gyrus (IFG)/ insula					
R superior frontal gyrus	21 54	36		4.4	486
R middle and inferior temporal	60 -27	-12		4.2	756 **
gyrus					
L fusiform gyrus	-30 -3	0 -24		3.8	243
R middle frontal gyrus	33 24	51		3.8	756
R anterior cingulum	6 21 1	5		3.8	162
R midbrain	9 -21 -	-9		3.7	135
L superior frontal gyrus	-27 57	27		3.7	270
L cerebellum	-9 -84	-39		3.6	459
R cerebellum	12 -48	-18		3.5	351
R middle frontal gyrus	48 36	30		3.5	135
L midbrain	-6 -30	-12		3.5	162
Seed region 2: L amygdala					
L middle temporal gyrus	-48 -3	-24		4.2	459

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Seed region 3: R fusiform face

area (FFA)

No suprathreshold clusters.

Results are shown at a threshold of *p*< 0.001, uncorrected, at voxel-level; cluster significance

was assessed using FWE correction [p < 0.05] for multiple comparisons across the a priori

ROIs based on small volume correction (SVC) (significant clusters marked with \*\*). Only

clusters ≥ 135 mm³ (voxel size 3 x 3 x 3 mm³) are reported. df = degrees of freedom, R =

right, L = left.

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**Table 4.** Linear associations between MSCEIT scores and gray matter volume. Voxel based morphometry (VBM).

Peak MNI coordinate region	Peak	MNI	coordinate	t peak	voxel	Cluster size
	(x y z)			(df = 80	)	(mm³)
Positive relationship						
R middle and superior frontal gyri,	27 48	-20		4.0		1596 **
partes orbitales						
L fusiform gyrus/ inferior temporal	-28 -4	-41		3.8		486
gyrus						
Negative relationship						
L cuneus	-4 -90	36		3.9		358
L superior partietal gyrus/ superior	-10 -82	2 46		3.7		186
occipital gyrus						

Results are shown at a threshold of p< 0.001, uncorrected, at voxel-level; cluster significance was assessed using FWE correction [p < 0.05] for multiple comparisons across the a priori anatomical ROI based on small volume correction (SVC; significant clusters marked with \*\*). Only clusters  $\geq$  135 mm³ are reported (voxel size: 1.5 x 1.5 x 1.5 mm³). df = degrees of freedom, R = right, L = left.

Supplementary Table 1. Euclidian distances between the peak of the area in the right TVA which exhibited a positive linear association of its voice-sensitive FC modulations with the individual MSCEIT scores (60 -27 -12) and the location of TVA voice-sensitivity peaks in the present study as well as previous reference studies by Pernet et al. (2015)) and Belin et al. (2000)).

		Peak MNI coordinate	Euclidian
		(x y z)	distance (mm)
This publication	R TVA anterior peak	60 -12 -3	17.5
	R TVA posterior peak	57 -21 0	13.7
Pernet et al.	R TVA anterior peak	60 -14 0	17.7
2015			
	R TVA posterior peak	60 -26 0	12.0
Belin et al. 2000	R TVA anterior STS peak	60 -1 -4	27.2
	R TVA middle STS peak	63 -13 -1	18.1
	R TVA posterior STS peak	56 -30 6	18.7
	R TVA MTG peak	52 -19 -1	15.8
Pernet et al.	R anterior STS	55 -2 -7	26.0
2015			
	R middle STS	53 -18 -3	14.5
	R posterior STS	42 -35 3	24.8

The Euclidian distances are calculated by the square roots of the sum of the squared distances between the x-, y- and z-coordinates of the mentioned peak voxels to the peak of the right TVA cluster which exhibited a positive linear association of its voice-sensitive FC

modulations with the individual MSCEIT scores (60 -27 -12). R = right, TVA = temporal voice area, STS = superior temporal sulcus, MTG = middle temporal gyrus.

**Supplementary Table 2.** Functional regions of interest (ROI) for complementary validation analyses: face-sensitive areas in the pSTS and the inferior occipital gyrus (occipital face area, OFA) in 85 healthy individuals.

Face-sensitive	Peak MNI coordinate	t peak voxel	Cluster size	p value
areas	(x y z)	(df = 81)	(mm³)	
Right pSTS	57 -60 12	12.9	29349	<0.001
Left pSTS	-57 -66 21	9.6	20385	<0.001
Right OFA	42 -81 -12	9.1	1917	0.001
Left OFA	-39 -54 -21	5.1	162	0.031

For the definition of the pSTS the middle and superior temporal gyri as well as the angular gyrus were defined as a priori anatomical ROI, and the inferior occipital gyrus for the definition of the OFA. Results are based on a threshold of p < 0.001, uncorrected at voxel level, and FWE correction for multiple comparisons at cluster level with p < 0.05 across these a priori ROIs. Voxel size was  $3 \times 3 \times 3$  mm³. df = degrees of freedom. P values given in the table are FWE corrected.

Figure captions

Figure 1. Linear associations between MSCEIT scores and cerebral voice-sensitivity.

Correlations of MSCEIT scores with voice-sensitivity (red) rendered onto a standard brain

(A) and coronal as well as transversal slices of the study population mean anatomical scan

(B, C). Functional (TVA, FFA) and anatomical regions of interest are rendered in different

colors (TVA = dark blue; FFA = green; amygdala = yellow). Results shown at a threshold of p

< 0.001, uncorrected, at voxel-level; cluster significance was assessed using FWE-correction

for multiple comparisons across the whole brain (marked with \*) and the a priori ROIs (small

volume correction, SVC), respectively (marked with \*\*) with a threshold of p < 0.05. The

diagram (D) illustrates the direction of the relationship between the MSCEIT scores and

voice-sensitivity in the left IFG/ insula).

Figure 2. Linear associations between MSCEIT scores and absolute contrast estimates

in voice-sensitive left IFG/ insula and amygdala.

Correlations of MSCEIT scores with corrected contrast estimates for voices, animals, and

environmental sounds in the (A) left IFG/ insula and (B) left amygdala. There is a positive

correlation of increasing MSCEIT scores and higher contrast estimates for voices (red line),

whereas there is obviously no clear effect on animal (brown line) and environmental sounds

(blue line). In the left IFG/ insula only the third of subjects with the highest EI showed a

positive correlation, whereas subjects with medium and low El showed a slightly negative

correlation (C). A similar, although only marginal effect was observed in the left amygdala

(D).

Figure 3. Linear associations between MSCEIT scores and cerebral face-sensitivity.

Correlation of MSCEIT scores with face-sensitivity (red) rendered onto a standard brain (A)

and a coronal slice of the study population mean anatomical scan (B). Functional (TVA, FFA)

regions of interest are rendered in different colors (TVA = dark blue; FFA = green). Amygdala

ROI not displayed in this figure. Results shown at a threshold of p < 0.001, uncorrected, at

voxel-level; the cluster marked with two asterisks is found to be significant using FWE-

correction for multiple comparisons across the FFA as a priori ROIs (small volume correction,

SVC), with a threshold of p < 0.05. The diagram (D) illustrates the direction of the relationship

between the MSCEIT scores and face-sensitivity in the right FFA).

Figure 4. Linear associations between MSCEIT scores and absolute contrast estimates

in face-sensitive right FFA.

Correlations of MSCEIT scores with corrected contrast estimates for faces, houses, objects,

and scenes in the right FFA (A). Decreased face-sensitivity in individuals with higher

MSCEIT scores is due to a decrease in contrast estimates for faces (red line) as well as an

increase in contrast estimates for houses (green line), objects (brown line) and scenes (blue

line). Subjects with low EI showed higher face-sensitivity-contrast estimates than did subjects

with medium and high EI (B).

Figure 5. Linear associations between MSCEIT scores and voice-sensitive

modulations of FC (PPI).

Correlations of MSCEIT scores with voice-sensitive (red) modulations of connectivity (A-C).

Functional (TVA, FFA) and anatomical regions of interest are rendered in different colors

(TVA = dark blue; FFA = green; amygdala = yellow). Results shown for the left IFG/ insula

seed at a threshold of p < 0.001, uncorrected, at voxel-level (A, B); cluster significance was

assessed using FWE-correction for multiple comparisons across the a priori ROIs (SVC;

marked with \*\*) with a threshold of p < 0.05. The diagram (D) illustrates the direction of the

relationship between the MSCEIT scores and voice-sensitive FC modulations between the

left IFG/ insula and the right TVA.

Figure 6. Convergence of negative relationships between MSCEIT scores and face-

sensitivity and gray matter volume in the right FFA.

Negative correlations of MSCEIT scores with face-sensitivity (green) and gray matter volume

(red) rendered onto a standard brain (A) and a transversal slice of the study population mean

anatomical scan (B). Results shown at a threshold of p < 0.01, uncorrected, for illustration

purposes. The diagram (C) illustrates the direction of the relationship between the MSCEIT

scores and gray matter volume in the part of the right FFA which exhibited a negative

correlation between MSCEIT scores and face-sensitivity.

Figure 7. Positive relationship between MSCEIT scores and gray matter volume in the

right OFC.

Positive correlation of MSCEIT scores with gray matter volume (red) rendered onto a

standard brain (A) and a transversal slice of the study population mean anatomical scan (B).

Results shown at a threshold of p < 0.001, uncorrected, cluster significance was assessed

using FWE-correction for multiple comparisons across the a priori anatomical ROI (small

volume correction, SVC) with a threshold of p < 0.05. The diagram (C) illustrates the direction

of the relationship between the MSCEIT scores and gray matter volume in the right OFC.

Supplemental Figure 1. Linear associations between experiential and strategic El and

voice- and face-sensitive regions in the left IFG/ insula and right FFA.

In the left IFG/ insula (A) and in the right FFA (B) the experiential EI (red line) contributed

more strongly to the voice- and face-sensitivity than did strategic EI (blue line).

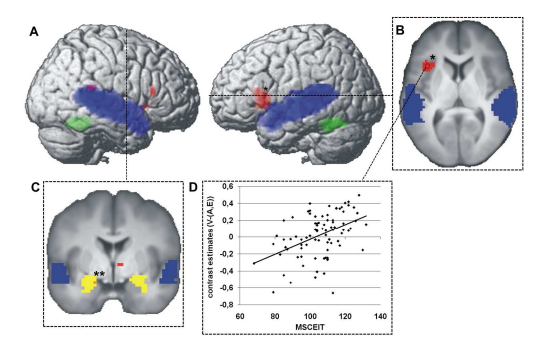


Figure captions
Figure 1. Linear associations between MSCEIT scores and cerebral voice-sensitivity.

Correlations of MSCEIT scores with voice-sensitivity (red) rendered onto a standard brain (A) and coronal as well as transversal slices of the study population mean anatomical scan (B, C). Functional (TVA, FFA) and anatomical regions of interest are rendered in different colors (TVA = dark blue; FFA = green; amygdala = yellow). Results shown at a threshold of p < 0.001, uncorrected, at voxel-level; cluster significance was assessed using FWE-correction for multiple comparisons across the whole brain (marked with \*) and the a priori ROIs (small volume correction, SVC), respectively (marked with \*\*) with a threshold of p < 0.05. The diagram (D) illustrates the direction of the relationship between the MSCEIT scores and voice-sensitivity in the left IFG/ insula).

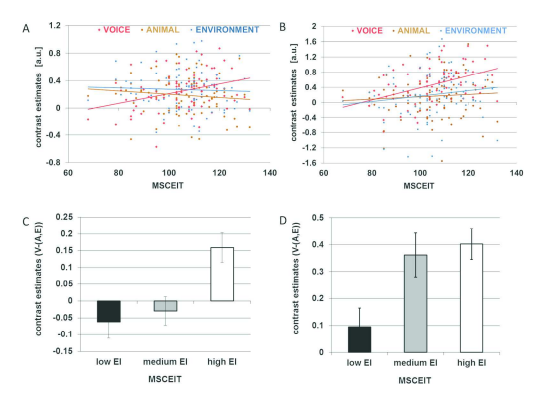


Figure 2. Linear associations between MSCEIT scores and absolute contrast estimates in voice-sensitive left IFG/ insula and amygdala.

Correlations of MSCEIT scores with corrected contrast estimates for voices, animals, and environmental sounds in the (A) left IFG/ insula and (B) left amygdala. There is a positive correlation of increasing MSCEIT scores and higher contrast estimates for voices (red line), whereas there is obviously no clear effect on animal (brown line) and environmental sounds (blue line). In the left IFG/ insula only the third of subjects with the highest EI showed a positive correlation, whereas subjects with medium and low EI showed a slightly negative correlation (C). A similar, although only marginal effect was observed in the left amygdala (D).

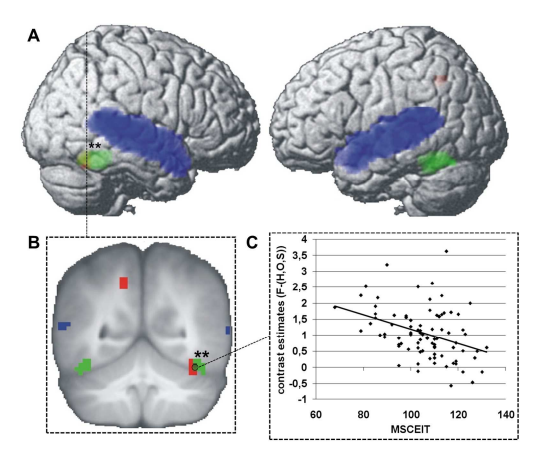


Figure 3. Linear associations between MSCEIT scores and cerebral face-sensitivity. Correlation of MSCEIT scores with face-sensitivity (red) rendered onto a standard brain (A) and a coronal slice of the study population mean anatomical scan (B). Functional (TVA, FFA) regions of interest are rendered in different colors (TVA = dark blue; FFA = green). Amygdala ROI not displayed in this figure. Results shown at a threshold of p < 0.001, uncorrected, at voxel-level; the cluster marked with two asterisks is found to be significant using FWE-correction for multiple comparisons across the FFA as a priori ROIs (small volume correction, SVC), with a threshold of p < 0.05. The diagram (D) illustrates the direction of the relationship between the MSCEIT scores and face-sensitivity in the right FFA).

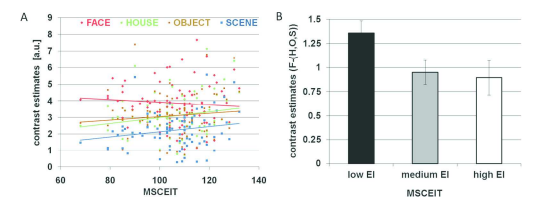


Figure 4. Linear associations between MSCEIT scores and absolute contrast estimates in face-sensitive right

Correlations of MSCEIT scores with corrected contrast estimates for faces, houses, objects, and scenes in the right FFA (A). Decreased face-sensitivity in individuals with higher MSCEIT scores is due to a decrease in contrast estimates for faces (red line) as well as an increase in contrast estimates for houses (green line), objects (brown line) and scenes (blue line). Subjects with low EI showed higher face-sensitivity-contrast estimates than did subjects with medium and high EI (B).

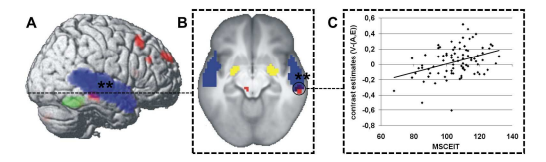


Figure 5. Linear associations between MSCEIT scores and voice-sensitive modulations of FC (PPI). Correlations of MSCEIT scores with voice-sensitive (red) modulations of connectivity (A-C). Functional (TVA, FFA) and anatomical regions of interest are rendered in different colors (TVA = dark blue; FFA = green; amygdala = yellow). Results shown for the left IFG/ insula seed at a threshold of p < 0.001, uncorrected, at voxel-level (A, B); cluster significance was assessed using FWE-correction for multiple comparisons across the a priori ROIs (SVC; marked with \*\*) with a threshold of p < 0.05. The diagram (D) illustrates the direction of the relationship between the MSCEIT scores and voice-sensitive FC modulations between the left IFG/ insula and the right TVA.

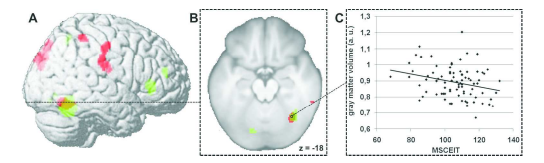


Figure 6. Convergence of negative relationships between MSCEIT scores and face-sensitivity and gray matter volume in the right FFA.

Negative correlations of MSCEIT scores with face-sensitivity (green) and gray matter volume (red) rendered onto a standard brain (A) and a transversal slice of the study population mean anatomical scan (B). Results shown at a threshold of p < 0.01, uncorrected, for illustration purposes. The diagram (C) illustrates the direction of the relationship between the MSCEIT scores and gray matter volume in the part of the right FFA which exhibited a negative correlation between MSCEIT scores and face-sensitivity.

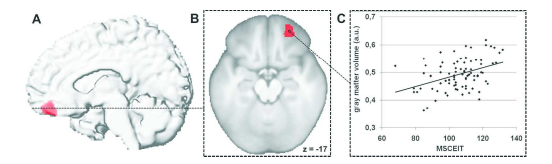


Figure 7. Positive relationship between MSCEIT scores and gray matter volume in the right OFC. Positive correlation of MSCEIT scores with gray matter volume (red) rendered onto a standard brain (A) and a transversal slice of the study population mean anatomical scan (B). Results shown at a threshold of p < 0.001, uncorrected, cluster significance was assessed using FWE-correction for multiple comparisons across the a priori anatomical ROI (small volume correction, SVC) with a threshold of p < 0.05. The diagram (C) illustrates the direction of the relationship between the MSCEIT scores and gray matter volume in the right OFC.

