Functional specializations for music processing in the human newborn brain

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In adults, specific neural systems with right-hemispheric weighting are necessary to process pitch, melody, and harmony as well as structure and meaning emerging from musical sequences. It is not known to what extent the specialization of these systems results from long-term exposure to music or from neurobiological constraints. One way to address this question is to examine how these systems function at birth, when auditory experience is minimal. We used functional MRI to measure brain activity in 1- to 3-day-old newborns while they heard excerpts of Western tonal music and altered versions of the same excerpts. Altered versions either included changes of the tonal key or were permanently dissonant. Music evoked predominantly right-hemispheric activations in primary and higher order auditory cortex. During presentation of the altered excerpts, hemodynamic responses were significantly reduced in the right auditory cortex, and activations emerged in the left inferior frontal cortex and limbic structures. These results demonstrate that the infant brain shows a hemispheric specialization in processing music as early as the first postnatal hours. Results also indicate that the neural architecture underlying music processing in newborns is sensitive to changes in tonal key as well as to differences in consonance and dissonance.

auditory cortex | functional MRI | neonates | emotion

Music is a cultural phenomenon and an art; however, in recent years, it has also become a fruitful research tool in numerous fields of cognitive neuroscience such as auditory perception (1), learning and memory (2), brain plasticity (3), sensorimotor processing (4), and the mirror neuron system (5). Consistent results have described brain networks for music processing in adult nonmusicians involving the superior temporal gyrus, inferior frontal, and parietal areas, with a dominance of the right over the left hemisphere (1). These neural networks may be intrinsically related to the emergence of musical abilities in humans. It is, however, difficult to determine to what extent music processing skills, as observed in adults, are the result of an adaptation of the brain from exposure to the musical environment or to biological constraints that lead, with normal experience, to the typical trajectory of brain development. One way to address this issue is to study how the brain processes musical stimuli at a point where exposure to music has been minimal and presumably not sufficient to induce major shaping of the processing networks. We used functional MRI (fMRI) to investigate the neural correlates of music processing in neonates. In recent years, fMRI has been used successfully with pediatric and healthy infant populations (6–8), proving to be a noninvasive and reliable technique yielding valuable information about brain development. The babies who participated in the study were first exposed to music initially outside the uterine environment, allowing observation of the early developmental stages of a capacity that plays an important role for emotional, cognitive, and social development from the first days of life (9).

Music perception relies on sophisticated cognitive skills for the decoding of pitch, rhythm, and timbre and for the processing of sequential elements that form hierarchical structures and convey emotional expression and meaning. Despite the complexity of such cognitive operations, mounting evidence indicates that newborns and young infants are highly sensitive to musical information. Music modulates infants’ attention and arousal levels (10) and evokes pleasure or discomfort. Infants with casual exposure to music possess the abilities for relational processing of pitch and tempo; for the differentiation of consonant vs. dissonant intervals; and for the detection of variations in rhythm, meter, timbre, and tempo as well as duration of tones and musical phrases (11, 12). These musical competences of infants play a crucial role in early language learning, because the processing of speech prosody (e.g., speech melody, speech rhythm) provides important cues for the identification of syllables, words, and phrases (13–16). Neurophysiological studies using near-infrared optical topography (17) and magnetoencephalography (18) in 3-month-old infants showed more prominent activation to normal speech sounds than to flattened speech sounds in right temporo-parietal regions. This suggests that processing the slow-changing melodic components of natural speech relies on right-hemispheric neural resources already engaged at 3 months of age. Despite this evidence for the importance of musical competences in early childhood, the neural basis of music processing in infants has remained elusive and no previous study has investigated music processing in neonates with fMRI.

The present study addresses this issue, using excerpts of classical music pieces and counterparts of these excerpts that varied in their syntactic properties and in their degree of consonance and dissonance. The control conditions were chosen so as to be acoustically closely matched to the original stimuli and still musical. This allowed a more precise interpretation of any result observed in the original music condition. In other words, we asked whether the newborn brain would be sensitive to subtle variations of the musical stimuli or whether it would respond indiscriminately to any music-like stimulus.

Using fMRI, data were obtained from 18 healthy full-term newborns within the first 3 days of life. Babies heard 21-s musical stimuli alternating with 21-s blocks of silence (Fig. 1B). Three sets of musical stimuli were used (Fig. 1A): Set 1 (“original music”) consisted of original instrumental (piano) excerpts drawn from the corpus of major-minor tonal (“Western”) music of the 18th and 19th centuries. This experimental condition addressed a basic question about the neural network involved in music processing. From these original excerpts, two further sets (altered music: sets 2 and 3) were created. In


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set 2 (key shifts), all voices were infrequently shifted one semitone upward or downward, thus infrequently shifting the tonal center to a tonal key that was harmonically only distantly related to the preceding harmonic context (e.g., from C major to C# major). The recognition of such key changes involves that listeners sequence the musical information, establish and maintain a representation of a tonal center from the different tones of a musical passage, and realize when the representation of a tonal center differs from the tonal center of incoming musical information (19, 20). The extraction of a tonal center is a basis for the music-syntactic processing of harmonies (19, 20), and processing changes of the tonal center involves prefrontal cortical areas, particularly the pars opercularis of the inferior frontal lobar cortex (in the left hemisphere part of Broca’s area), ventrolateral premotor cortex, and dorsolateral prefrontal cortex in both adults and 10-year-old children (21, 22). Notably, the inferior frontal lobar cortex has particularly been shown to serve the processing of syntax in both music and language, although with right-hemispheric weighting in the music domain (21–23) and left-hemispheric weighting in the language domain (24, 25). We aimed at testing whether the brains of newborns are already sensitive to changes in tonal key and which brain areas would be involved in newborns for the processing of key changes.

In set 3 (dissonance), the upper voice (i.e., melody) of the musical excerpts was permanently shifted one semitone upward, rendering the excerpts permanently dissonant [similar to previous studies (26, 27)]. Dissonance and consonance are basic perceptual properties of tones sounding together, and the perception of consonance and dissonance relies on properties of the auditory pathway, presumably independent of extended auditory experience (11, 28, 29). Particularly in Western listeners, consonant tone combinations are perceived as more pleasant than permanently dissonant ones (26, 27, 29) and 2-month-old infants already show a preference of consonance over dissonance (11, 30, 31). We used the dissonant stimuli of set 3 to investigate whether neonates’ brains are sensitive to dissonance.

Stimulus sets 2 and 3 were also used to investigate neural correlates of emotional responses in newborns. In addition to studies showing that consonant music is usually perceived as more pleasant than permanently dissonant music, previous work with adults has shown that unexpected chord functions and changes in tonal key elicit emotional responses in listeners (32, 33). Up to now, however, the neural origins of human emotion in early life have remained unknown, and the knowledge about emotional processes obtained from adults cannot be extended to children, because the processes underlying the generation of emotions are heavily shaped by lifelong experiences. Studies investigating emotion using musical stimuli in adults revealed that music-evoked emotions involve core structures of emotional processing, such as the amygdala, nucleus accumbens, and orbitofrontal cortex (26, 27, 34). Our study also aimed at exploring the early sensitivity of such core structures of emotional processing, and the specificity of their responses, to original and altered music stimuli.

Note that all three stimulus sets were identical with regard to tempo, meter, rhythm, timbre, and contour and that the stimulus alterations (key shifts and dissonance) did not turn the original music into disorganized noise. Note also that newborns had little or no exposure to music except for the heavily filtered sounds reaching the fetus in utero (35). Thus, if the original music and the stimulus alterations elicit specific brain activity in newborns, this would also indicate predispositions for the processing of these musical features that are largely independent of culture-specific adaptations.

The present study uses fMRI to explore brain specialization for music processing in newborns with minimal exposure to external auditory stimuli.

### Results

The main effect of original music (set 1) vs. silence showed an extended right-hemispheric activation cluster focused in the superior temporal gyrus, with its peak activation being located in the primary auditory cortex (transverse temporal gyrius), extending into the secondary auditory cortex, and anteriorly into the planum polare, as well as posteriorly into the planum temporale, temporoparietal junction, and inferior parietal lobule (Fig. 2A); this observation was confirmed by region of interest (ROI) analysis (see below and Fig. 4). In addition, activation was observed in the right insula and the right amygdala-hippocampal complex (see Fig. 3).

We then addressed the differences in activation between the two altered conditions and silence: key shifts (set 2) vs. silence and dissonance (set 3) vs. silence. Both comparisons showed a very comparable pattern, namely, less extended activation in the right temporal regions (compared with the main effect of original music vs. silence) and, instead, left-hemispheric activation clusters in the superior and middle temporal regions (including primary auditory cortex), inferior frontal gyrus, amygdala, and ventral striatum (see Fig. 2B for pooled datasets). The direct comparison between the two altered conditions showed no significant differences in activation, except that blood oxygenation level-dependent contrast (BOLD) signal changes were stronger during the key shifts (set 2) in the left amygdala and ventral striatum. Because of the essentially identical activation patterns elicited by set 2 and 3, data of both sets were pooled, leading to an equal amount of scans for subsequent comparisons between altered music (sets 2 and 3 pooled) vs. silence and between original music (set 1) vs. altered music (details provided in Materials and Methods). The main effect of altered music (sets 2 and 3 pooled) vs. silence (Fig. 2B) illustrates that the right-hemispheric activation of the primary auditory cortex and posterior superior temporal gyrus (which was
observed up to the temporoparietal junction) was less pronounced compared with the main effect of original music vs. silence (Fig. 2A). The main effect of altered music vs. silence also shows activation of the left inferior frontal gyrus (which was not observed in the comparison of original music vs. silence) and activation of the left primary auditory cortex, which was comparable to that observed in the effect of original music vs. silence (see Fig. 3 for the direct contrast between original music and altered music).

The direct contrast of original music vs. altered music shows areas with significantly stronger hemodynamic responses for original music ($P < 0.05$, voxel level, uncorrected for cluster extent) in the superior and posterior part of the right auditory cortex and in the right amygdala-hippocampal complex (Fig. 3). Areas with stronger hemodynamic responses for altered music were observed in the left inferior frontal gyrus, and the left amygdala-hippocampal complex (a more precise identification of the medial temporal lobe structures, given their small size, exceeds the limits of 1.5 T fMRI spatial resolution).

To address specifically the level of activity in the right and left primary and secondary auditory cortices, ROI analysis was conducted on subjects’ average percent signal change for original music and altered music (Fig. 4 and Materials and Methods). ANOVA on these ROI values with the factors of hemisphere and stimulus type (two levels: set 1, pooled sets 2 and 3) indicated a significant interaction between hemisphere and stimulus type ($F = 4.74$, $P = 0.03$). This interaction was due to the difference in hemodynamic response between the two stimulus types in the right hemisphere, with stronger activation for original music compared with altered music (multiple comparison Tukey honestly significant difference, $P = 0.05$). In the left auditory cortex, no difference in activation was observed for the two stimulus types.

Discussion

Our data show that a hemispheric functional asymmetry for music perception is present at birth. Activation with right-hemispheric dominance was observed in the primary, secondary, and higher order auditory cortices in newborns, consistent with data obtained from adult nonmusicians, especially for the processing of pitch relations (36). This right-hemispheric dominance indicates that the newborn brain responds to musical information quite specifically, because the right primary auditory cortex is particularly involved in pitch analysis and integration (i.e., in the decoding of pitch height, pitch direction, and pitch chroma) (37), a prerequisite for the processing of music. Notably, the activations extended beyond Heschl’s gyrus toward the right planum polare as well as toward the planum temporale and the inferior parietal lobule. Previous functional neuroimaging studies with adults showed that the right superior temporal gyrus (including the planum polare) becomes increasingly involved with increasing melodic complexity (38) and that right-hemispheric areas of auditory cortex outside the primary zone are specialized in the processing of pitch patterns, in the encoding and recognition of melodies, and in auditory-motor transformation (39, 40). Our data show that these areas are already recruited with right-hemispheric dominance by newborns for the processing of musical information.

In adult brains, the right-hemispheric preference for music processing has been attributed to different specializations of the left and right auditory cortices for the processing of temporal and spectral aspects of acoustic stimuli (41, 42). These cortices might have evolved a complementary specialization, with the left hemisphere having better temporal resolution (crucial for speech analysis) and the right hemisphere having a better frequency resolution (required for pitch processing) (43). These functional hemispheric differences correspond to anatomical findings of volume differences in the left and right auditory cortices (41, 44).

The lateraled specialization of brain structures is not a uniquely human phenomenon, and it is not determined by specific exposure to environmental sounds (45). Gross anatomical asymmetries around the Sylvian fissure have been observed in fetuses from midgestation and in newborns (46). Recently, studies have identified a large number of genes expressed differently in the left and right perisylvian fetal cortices at different weeks of gestation corresponding with periods of neural proliferation and migration (47, 48). These hemispheric asymmetries, may result in early functional asymmetries, even before the onset of hearing (49).

The right-preponderant auditory cortex activation observed in the present study parallels the left-hemispheric auditory cortex activations shown for infants hearing speech (8). Taken together, these results demonstrate the presence of early propensities in the way the auditory nervous system processes sound (8, 50). Notably, the areas

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Fig. 2. Activations elicited by the musical stimuli in newborns ($n = 18$, random effects group analysis; $P < 0.0002$ at the voxel level and $P < 0.05$ at the cluster level) overlaid over a T2-weighted image from a single newborn subject (note that the spatial resolution of the functional group data is lower compared with the anatomical image). (A) Mean activations for original music vs. silence are shown for six axial slices. Note the right-hemispheric predominance of temporal activation (yellow arrows). (B) Mean activations for altered music (key shifts and dissonance pooled) vs. silence. Note the left-hemispheric activation in the inferior frontal gyrus (orange arrows) and the reduced activation in the right temporal lobe (compared with the contrast of original music vs. silence, white arrow). (Details are provided in Materials and Methods.)

Fig. 3. Direct contrast of original music vs. altered music in healthy newborns ($n = 18$, random effects group analysis; $P < 0.05$ at the voxel level, uncorrected) overlaid on a T2-weighted image from a single newborn (note that the spatial resolution of the functional group data is lower compared with the anatomical image). Regions more active for original music are shown in orange/yellow, and regions more active for altered music are shown in blue. Two axial slices show a stronger activation of the left inferior frontal gyrus in response to altered music. The slices also show a stronger activation of (posterior) auditory cortex in response to original music. The two coronal slices show activation of the left amygdala-hippocampal complex (and of the ventral striatum) for altered music and activation of the right amygdala-hippocampal complex for original music. The two sagittal slices show the larger right superior temporal activation for original music.
activated by hearing music were not confined to primary auditory cortices. This is reminiscent of auditory cortex activations reported for infants hearing sentences (51) and presumably implies that the complex hierarchical organization of auditory language processing is paralleled by a similar hierarchical organization of music processing very early in life.

In addition to the asymmetry of auditory cortex activation for the processing of the original music, we showed that BOLD signals were modulated by alterations of the musical stimuli. In particular, in the BOLD signal changes in response to the altered music (compared to the original music) were smaller in the right primary and secondary auditory cortex (Fig. 2 and Fig. 4), but larger in left-hemispheric superior temporal and inferior-frontal cortices (Fig. 2B). When newborns perceived stimuli that contained manipulations of the acoustic structure (i.e., with regard to dissonance) and manipulation of the music-syntactic structure (i.e., with regard to the shift of tonal key), we observed rather symmetrical activity of the primary auditory cortices (Fig. 4). These findings indicate that newborns’ neural responses to musical stimuli can be modulated by structural variations of the stimuli. Importantly, a significant difference in the right auditory cortex activity was observed when directly comparing original music vs. altered music, that is, two conditions in which musical stimuli were presented (Fig. 4). Thus, the right-lateralized auditory cortex activation was attributable to the specific features of the (mainly consonant and structured) original music and was not simply an unspecific response that could have been elicited by any sound in general. These results are in agreement with behavioral evidence from neonates and older babies showing that infants are highly sensitive to pitch alterations and that this ability is subserved by neural predispositions present from birth (52).

The pattern of activation observed for the altered music condition presumably reflects a sensitivity of the newborn brain to dissonance (both altered sets contained a higher degree of sensory dissonance compared with the original sets: set 2 consisted of more dissonant intervals than the original music, and the key shifts in set 3 also compared with the original sets: set 2 consisted of more dissonant intervals than the original music, and the key shifts in set 3 also), introduced a subtle sensory dissonance attributable to the semitone shift of notes (SI Text). Neuropsychological and neuroimaging studies from adults as well as behavioral studies with older babies (12) showed that human infants tested at 7 months perceive sensory dissonance similar to adults (53), that infants prefer to listen to consonant intervals (11, 30) and appear to discriminate consonant and dissonant music shortly after birth (28). The perception of sensory dissonance is a function of the physical properties of auditory stimuli, as well as those of basic physiological and anatomical constraints, resulting from limitations of the auditory system in resolving tones that are too proximal in pitch (54). This phenomenon is independent of specific experience and is observable across species (55, 56). Although the specific weighting of genetic and experiential factors in shaping early sensitivity to consonance and dissonance remains to be specified, our results indicate sensitivity of the auditory cortex to consonance and dissonance already at birth.

In conclusion, our results show that a neuronal architecture serving the processing of music is already present at birth. Right-lateralized auditory cortex activity was observed for the processing of original music, indicating that neonates already show a right-hemispheric predominance for the processing of musical information. This activation was modulated by contrasting original music with altered but still musical stimuli, indicating that the response we obtained was not generically driven by auditory stimulation. Our data show that the
newborn’s brain is sensitive to changes in the tonal key of music and to differences in the sensory dissonance of musical signals. Such structural manipulations activated the inferior frontal cortex, and thus possibly Broca’s region, which becomes fundamentally important for the learning of language during later stages of development. Finally, the activity changes within limbic structures suggest that newborns engage neural resources of emotional processing in response to musical stimuli. One inherent limitation of our study is that it cannot estimate the extent to which uterine input has already shaped the neural systems at birth. The fetus perceives auditory information in the final weeks of gestation, and salient features of this input can be recognized after birth (62–64). Also, our findings do not imply that the specific response we observe when the newborn brain is exposed to musical information evolved for, and is exclusively involved in, instrumental music processing (similar responses are likely to be elicited by song, and by the melodic aspects of speech). Future studies should explore the responses at birth to auditory stimuli with different degrees of musical content, such as melodic singing, child-directed speech, nonlexical prosodic speech, and with minimal prosodic content. This study demonstrates competencies of newborns for the processing of music, providing insight into the neural origins of music, a universal human capability.

Materials and Methods

Subjects. Eighteen healthy, full-term, nonsedated newborns (8 female, Apgar score ≥8) within the first 3 days of life participated in the study. Gestation and birth histories were normal for all subjects. Data from 3 other newborns were not used because of large movement artifacts. The majority of the subjects’ immediate family members were right-handed (88%) (65), with no history of learning disabilities or psychiatric and neurological disorders, and of mono- liguinal Italian background (one bilingual English-Italian), and they were not musicians. Infants’ behavior during scanning was recorded to monitor changes in the sensory dissonance of musical signals. Such structural manipulations activated the inferior frontal cortex, and thus possibly Broca’s region, which becomes fundamentally important for the learning of language during later stages of development. Finally, the activity changes within limbic structures suggest that newborns engage neural resources of emotional processing in response to musical stimuli. One inherent limitation of our study is that it cannot estimate the extent to which uterine input has already shaped the neural systems at birth. The fetus perceives auditory information in the final weeks of gestation, and salient features of this input can be recognized after birth (62–64). Also, our findings do not imply that the specific response we observe when the newborn brain is exposed to musical information evolved for, and is exclusively involved in, instrumental music processing (similar responses are likely to be elicited by song, and by the melodic aspects of speech). Future studies should explore the responses at birth to auditory stimuli with different degrees of musical content, such as melodic singing, child-directed speech, nonlexical prosodic speech, and with minimal prosodic content. This study demonstrates competencies of newborns for the processing of music, providing insight into the neural origins of music, a universal human capability.

Sounds were presented via piezoelectric, European Union-certified, MRI-compatible headphones custom-made to fit newborns’ ears and incorporating analog noise-canceling technology to eliminate passive dead-ening, leading to noise reduction on the order of 30–40 dB above 600 Hz (MR Confon). With noise reduction, the scanner noise had an intensity of about 66 dB sound pressure level (SPL) at the eardrum of the infant. The sound pre-sentation was adjusted to a comfortable volume level (about 84 dB SPL) allowing the music to be clearly audible above residual scanner noise. A block design was used with 21 y-blocks alternating between conditions (music, silence, altered music, silence) in a pseudorandom order, such that two versions of the same excerpt never followed each other for a total scan time of 7 min (Fig. 1B). Each sequence contained only one kind of altered music. Two 7-min sequences were presented in alternate order to each successive infant: one containing music and key shifts alterations and one containing music and dissonance alterations. The two kinds of alteration were obtained from two different sets of five excerpts so as to avoid repeating the same nonaltered stimulus twice.

Data Analysis. Images were processed within the framework of the general linear model in AFNI (66).

After EPI time series reconstruction, every brain volume of each participant’s functional runs was examined to identify artifacts attributable to either subject head movement or MRI scanning system properties. Sequences from 3 subjects were excluded from analysis because of motion movement during scanning. Eighteen of the remaining 18 newborns provided usable data for two sequences. For the two other babies, only one sequence (with original music and key shifts) was usable. For within-subject motion correction and realignment, both functional time series were aligned to a “base” image free of artifacts. Motion correction and image realignment was performed with a weighted linear least-square algorithm (3dvolreg) (67, 68) with gradient descent, using six parameters of rigid body movement. In total, 198 volumes were obtained. In addition, large and motion correction were eliminated from further analyses. On average, 198 volumes were retained per subject (SD = 68). A spatial filter with an rms width of 4 mm was applied to each EPI volume.

To control for individual differences in absolute values of activation, for each voxel, we computed the mean intensity value of the sequence time series and divided it by the values within that voxel to obtain the percent signal change at each time point. The two functional sequences, if usable, were concatenated.

Template Creation and Image Registration. Because a suitable newborn template is currently unavailable, we created an ad hoc template from the subjects’ whole-head EPI scans (3.75 × 3.75 × 3 mm resolution) (SI Text). For each subject, a 12-parameter affine-general transformation algorithm (3d WarpDrive) was then used to coregister the subject’s average functional volume to the infant template. The same transformation matrix was subsequently applied to the subject’s functional scans to align them to the infant template.

Image Analysis: Single-Subject Level. Multiple linear regression implemented in AFNI’s 3DDeconvolve was used to fit stimulus reference vectors to the MRI time series at each voxel for each participant, obtaining an estimate of the BOLD response to each condition of interest (69). For the 16 subjects for whom two sequences were usable, the regression was run with nine regressors of no interest (mean, linear trend, second-order polynomial within each sequence to account for slow changes in MRI signal, and six outputs from the motion-correction algorithm to account for residual variance attributable to subject motion not corrected by registration) and three regressors of interest (one per condition: music, key shifts alterations, and dissonance alterations). Each regressor of interest was obtained by convolving a square wave for each stimulation block of that condition with a gamma variate function approximating the hemodynamic response (70). For each subject, the regression models provided estimates of the response (% signal change of the BOLD response) to each stimulus type in each voxel.

Multiple regression was also run on 18 subjects, pooling dissonance and key shifts alterations, because both conditions were modeled as “altered music.” This comprehensive regression was run with nine regressors of no interest (as above) and two regressors of interest (one per condition: music and altered music).

Group Analysis. A random effects model implemented in AFNI’s 3DANOVA was used to create group maps in the common template space. The estimates obtained through the regression were entered in two-way mixed-effects ANOVAs performed on each voxel in template space, with stimulus type as a fixed variable and subjects as random variables. For the analysis on 16 subjects who had usable data from both sessions, three stimulus types were included: music, key shifts alterations, and dissonance alterations. For the comprehensive analysis on 18 subjects, music and altered music were used as variables. The ANOVAs generated activation maps for each stimulus


19. Koelsch S, Fritz T, V Cramon DY, Müller K, Friederici AD (2006) Investigating emotion processing, the single-subject statistical maps that were the basis for the group ANOVAs were used. Subjects’ average (unthresholded) percent signal change in each ROI was entered into random effects ANOVA with hemispheres (left/right) and stimulus type (music and altered noise) as fixed variables and subjects as random variables.

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Supporting Information

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SI Materials and Methods

Stimuli. The musical excerpts were taken from the following pieces: (i) J.S. Bach, French Suite No.1 in D, BWV812, Allemagne; (ii) Scarlatti, Sonata in E, K.162, L.21; (iii) Schubert, Six Moments Musicaux, D.780, Op.94, No.3 in F--; (iv) Schubert, Andante in C, D.29; (v) Schubert, 34 Valses sentimentales, D.779, Op.50; (vi) J.S. Bach, Two-Part Inventions, No.5 in Eb, BWV776; (vii) J.S. Bach, Two-Part Inventions, No.11 in G, BWV782; (viii) Scarlatti, Sonata in D, K.10, L.370; (ix) Scarlatti, Sonata in D, K.21, L.363; and (x) Mozart, Piano Sonata in G, K.283.

Analysis of Acoustical Deviance in Key Shifts. To estimate the degree of acoustical deviance introduced by the changes of tonal key in the excerpts with key changes, we performed auditory modeling of the acoustic congruency of all key shifts with auditory sensory memory traces established by the previous chords. Snippets 4 s in duration were entered into the analysis; half of the snippets contained a key shift after 3 s, and the other half consisted of the corresponding snippets without a key shift. The modeling estimates pitch images of the auditory sensory memory; acoustic information decays but is kept in the auditory sensory memory for a certain time. The results of the modeling are correlations of the non-key shifts with the key shifts. The modeling also estimates pitch images of the auditory sensory memory; acoustic information decays but is kept in the auditory sensory memory for a certain time. The results of the modeling are correlations of the information stored in the auditory sensory memory than the regular sequences without key shifts (Fig. S1). Auditory modeling was performed with the Institute for Psychoacoustics and Electronic Music (IPEM) toolbox. The length of the local context integration window was 0.1 s, and that of the global context integration window was 1.5 s.

Additional Newborn Data Analysis. Analyses revealed that a small group of subjects (n = 3) showed consistent stimulus-related negative BOLD responses for all conditions compared with baseline. A reversed BOLD response has been observed in a number of fMRI studies involving infants and has been attributed to sedation, sleep, and neural or vascular immaturity (1, 2). In particular, Anderson et al. (1) reported signal decrease in bilateral primary auditory cortex in nonsedated sleeping infants in conditions similar to our experimental setup. To avoid rejecting the subjects with these “negative activations” or “deactivations,” we modeled their BOLD response by convolving a negative gamma function (simulating inverted BOLD) with a square wave. The inclusion of these three subjects did not change the lateralization and overall pattern of the activation for all conditions.

Newborn Template. Because a suitable newborn template is currently unavailable, we created an ad hoc template from the subjects’ whole-head EPI scans (3.75 × 3.75 × 3 mm resolution). First, a subject whose head was well positioned in the imaging space and symmetrical in shape was selected. We subsequently registered one scan per subject to the reference scan, after applying spatial filtering with an rms width of 4 mm. The chosen scans were free of artifacts and were used for within-subject motion correction and alignment. The across-subject registration algorithm (3DWarpedDrive) used a 12-parameter affine-general transformation, correcting for scaling and shearing differences in addition to rigid body transformations. The registered scans were then averaged to create a group average volume. We then repeated the registration of each subject’s original volume to the group volume. Finally, the scans registered to the group volume were averaged to create the infant template.

For visualization of brain activations, statistical maps in the common template space were interpolated to 1 mm3 resolution and overlapped onto an equally interpolated anatomical T2-weighted scan from a single newborn subject.

Fetal Acoustic Exposure. There are some limits on fetal acoustic exposure. The maternal abdomen acts as a low-pass filter, with high-frequency energy rejected at a rate of ~6 dB per octave (3). The sound energy that filters through the abdominal wall, through the uterus, and into the amniotic fluid is further filtered by the skull of the fetus, as the sound is transmitted through a bone conduction route rather than through the external and middle ear systems. This results in the fetus being able to hear the low-frequency components of speech and music (<500 Hz) but only when the airborne signal exceeds 60 dB. Voices would be detected at normal conversational levels, but many of the speech sounds would not be discriminated. In the same way, the higher frequencies of musical stimuli would be filtered out and timbre would be modified. The acoustic environment of the late-term fetus is largely composed of a stream of maternal heartbeats, maternal breathing, and words spoken by the mother (4). Nevertheless, the auditory information that gets through to the fetus is sufficient to induce recognition not only of the maternal voice (5) but of speech and musical stimuli heard during the fetal period (6, 7).

Sensory Dissonance Attributable to Key Shifts. The key shifts introduced a higher amount of sensory dissonance than in-key chords because of the semitone shift of notes. The first tones of the key shifts formed more dissonant intervals with the preceding tones than tones of the non-key shifts [the term “sensory dissonance” is used with regard to the key shifts in the broader sense, referring to the relations between successive sounds, as reported by Parnicu (6)]. In other words, the tones of key shifts had a lower pitch commonality with the preceding notes compared with the non-key shifts (with pitch commonality being the major component of sensory dissonance). The sensory dissonance of the permanently dissonant excerpts (without key shifts) is attributable to the simultaneous sound of tones forming dissonant intervals. We use sensory dissonance to make clear that we do not refer to “musical dissonance” in the sense that chords are perceived as unexpected because of their syntactic irregularity.

Acoustic Analysis of Stimuli With and Without Key Shifts. Part of the auditory system stores incoming auditory information in the auditory sensory memory, establishes a model of acoustic regularities inherent in the auditory input, and automatically detects acoustic changes that deviate from the regularities of the established model (9). We performed an analysis modeling the acoustic congruency of key shifts with the acoustic information of the previous musical information stored in the auditory sensory memory (details are provided in Fig. S1). Fig. S1 shows that the key shifts introduced pitch information that was less congruent with the information stored in the auditory sensory memory (compared with the pitch information of the regular sequences without key shifts). Therefore, it is likely that such acoustic deviations were automatically detected by the auditory system.

Previous functional neuroimaging studies showed that such automatic change detection mainly involves the auditory cortex [reviewed in (10)], with additional contributions from the inferior frontolateral cortex, which has been suggested to play a role in the judgment of novelty for the allocation of attentional resources (10). Note that these studies showed an increase of BOLD signal in...
the temporal (and frontal) cortex in response to acoustic changes (in adults, fMRI studies with infants on automatic auditory change detection are, to our knowledge, not available to date). This suggests that, in our study, changes in the auditory cortex of infants in response to the key changes were not attributable to auditory sensory memory operations related to the detection of different pitches, because key shifts led to a reduction of BOLD signal in the auditory cortex in the right hemisphere for altered music (compared with original music) and no difference in BOLD response was observed in the left auditory cortex between altered and original music.


Fig. S1. Modeling of the acoustic congruency of key shifts with the acoustic information of the previous musical information stored in the auditory sensory memory. Stimulus snippets of all key shifts (and of the corresponding snippets without key shifts) were analyzed using the Contextuality Module of the Institute for Psychoacoustics and Electronic Music (IPEM) Toolbox; the length of the local context integration window was 0.1 s, and that of the global context integration window was 1.5 s, as suggested by Leman et al. [Leman M, Lesaffre M, Tanghe K (2005) IPEM toolbox for perception-based music analysis (version 1.02). Available at http://www.ipem.ugent.be/toolbox/index.htm]. The abscissa represents the time line (the key shift occurs at 3 s), and the ordinate depicts the correlation values. The data show the correlation of the local context (pitch representation of the current chord) with the global context (pitch representation of the previous musical information stored in the auditory memory), separately for key shifts (red line) and for the corresponding passages without key shifts (blue line; thin dotted lines indicate SEM). Note the drop of correlation for key shifts (red line), showing that key shifts were less congruent with the information stored in the auditory sensory memory than the regular sequences without key shifts (blue line).