Exploration of the Neural Correlates of Ticklish Laughter by Functional Magnetic Resonance Imaging

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The burst of laughter that is evoked by tickling is a primitive form of vocalization. It evolves during an early phase of postnatal life and appears to be independent of higher cortical circuits. Clinicopathological observations have led to suspicions that the hypothalamus is directly involved in the production of laughter. In this functional magnetic resonance imaging investigation, healthy participants were 1) tickled on the sole of the right foot with permission to laugh, 2) tickled but asked to stifle laughter, and 3) requested to laugh voluntarily. Tickling that was accompanied by involuntary laughter activated regions in the lateral hypothalamus, parietal operculum, amygdala, and right cerebellum to a consistently greater degree than did the 2 other conditions. Activation of the periaqueductal gray matter was observed during voluntary and involuntary laughter but not when laughter was inhibited. The present findings indicate that hypothalamic activity plays a crucial role in evoking ticklish laughter in healthy individuals. The hypothalamus promotes innate behavioral reactions to stimuli and sends projections to the periaqueductal gray matter, which is itself an important integrative center for the control of vocalization. A comparison of our findings with published data relating to humorous laughter revealed the involvement of a common set of subcortical centers.

Keywords: fMRI, hypothalamus, PAG, tickle, vocalization

Introduction

Tickling involves the unpredictable stimulation (Darwin 1872) of vulnerable parts of the body (armpits, chest, side of the waist, and sole of the foot; Black 1984) by a familiar person. It is an ambivalent stimulus, evoking a mixture of pleasurable and unpleasant feelings (Plessner 1961) and leads to an involuntary stereotyped motor reaction (Provine 2000). Ticklish laughter is associated with sympathetic (Fry 1994; Szameitat et al. 2011) and emotional arousal (Bachorowski and Owren 2003). Tickling evolved from the rough-and-tumble play of animals; it is a primitive form of humor, which was referred to by Darwin as “protohumor” (Darwin 1872; Provine 2000). It provokes stimulus-driven emotionally valenced “Duchenne” laughter, which is classified as a prototypical nonserious social incongruity (Gervais and Wilson 2005).

Darwin (1872) and Hecker (1873) independently advanced the idea that tickling and humor are linked, in so far as both trigger laughter and share common qualities as stimuli (humor being considered as a “tickling of the mind”). Later, the tickle reaction was purported to be the primitive building block on which humor developed (Mc Gee 1979; Weisfeld 1993; Provine 2000; Gamble 2001). In support of this theory, Fridlund and Loftis (1990) have observed the existence of a close correlation between self-reported ticklishness and the tendency to laugh, giggle, or smile in everyday life.

However, other authors do not regard ticklish laughter as either a precursor or an imitation of humorous laughter (Gregory 1924; Bergler 1956; Plessner 1961; Harris 1999). According to Harris and Christensenfeld (1997), tickling and humor share a common final motor response but not an internal physiological state of mirth.

A wealth of data on humor has been gleaned by functional magnetic resonance imaging (fMRI) (Iwase et al. 2002; Mobbs et al. 2003; Osaka et al. 2003; Wild et al. 2006; Watson et al. 2007; Schwartz et al. 2008), whereas on tickling, only 2 such studies have been conducted (Blakemore et al. 2000; Carlsson et al. 2000). Moreover, in these 2 studies, the activated sensory network, not the vocal response, was evaluated, probably because of the difficulty that is experienced in controlling head movements during the ensuing laughter reaction, which obfuscate localization in the deep brain region (Iwase et al. 2002). Using state-of-the-art fMRI, we have now successfully explored the brain regions that are activated by ticklish laughter. Vocalization involves pathways stemming from both the motor cortex and the limbic system (for a review, see Jürgens 2009), each of which is implicated in laughter (Wild et al. 2003). The limbic pathway arises in the anterior cingulate gyrus and leads through the periaqueductal gray matter (PAG), wherein it receives information concerning the emotional state from the medial thalamus, the amygdala, and the hypothalamus (Jürgens 1998). The motor cortical and the limbic pathways converge at the level of the reticular formation of the pons and the medulla oblongata to interact with phonatory motor neurons. The motor cortical pathway is deemed to subserve the production of learned vocalizations, whereas the limbic one is believed to be critically involved in the production of nonverbal innate vocalizations (Jürgens 2009). The latter tenet is supported by data that has been gleaned from human patients with lesions in the motor cortex. Although speech and song production are compromised in these individuals, nonverbal vocalizations, such as coughing, crying, and even some forms of laughter, are unaffected (Grosnasser et al. 1988). Moreover, laughing (gelastic) seizures are observed in young patients developing hamartomas, irritating the lateral hypothalamus (Valdueza et al. 1994; Delalande and Fohlen 2003); in this situation, speech is not compromised.

Laughter that is evoked by tickling is an ontogenetically precocious form of this emotional manifestation (Leuba 1941; Poeck 1985). As such, it may involve innate mechanisms rather
than learned vocal behavior (Scheiner et al. 2006). And apropos of innate mechanisms, a comparison of ticklish laughter in humans with emotional vocalization circuits in other mammals is warranted. A wealth of data has been gleaned from studies with squirrel monkeys (Jürgens 1998). In this species, electrical stimulation of the mediodorsal thalamus, the amygdala, the hypothalamus, or other regions of the brain that are implicated in the limbic pathway, corresponds to vocalizations of various emotional states, including expressions of enjoyment. And in nonhuman primates, as well as in rats, laughter-like emotional responses have been repeatedly evoked during playful chasing and tickling (Darwin 1872; Matsusaka 2004; Panksepp 2007). Against the background of available evidence, we hypothesize that in humans, ticklish laughter—as an innate and “primitive” form of vocalization—is rooted in the limbic pathway.

In our fMRI experiment, 3 different situations were established: 1) tickling and laughter, which involved tickling healthy participants on the sole of the right foot and allowing them to laugh; 2) tickling and inhibition of laughter; and 3) voluntary laughter. Inhibition of laughter is believed to reduce brain activity in relays that are critical for the neuronal control of vocalization. Voluntary laughter generally lacks the emotional features of true laughter. To reveal the regions that are essential for the triggering of ticklish laughter, activation that was registered during tickling and laughter was compared with that elicited under the other 2 conditions and was correlated with the corresponding number of laughter events.

Materials and Methods

Subjects
Among the 27 healthy participants, 18 (11 females and 7 males, mean age: 24 years; age range: 21–29 years) were included in this study. Nine subjects were excluded because their head movements consistently exceeded 3 mm in 1 of the 3 evaluated directions using the Statistical Parametric Mapping (SPM) realignment procedure.

The informed consent of all participants was obtained, and the procedure was approved by the Ethical Committee of the University Hospital of Greifswald, Germany.

Behavioral Data
An fMRI-adapted fiber-optic microphone (MR confon, Magdeburg, Germany) was used to record laughter during the scanning procedure. Events of ticklish laughter and voluntary laughter were counted and classified as "strong" (more than one audible articulation) or "weak" (only one audible articulation). Strong vocalizations were awarded a 2-fold-high weighting than weak ones, which permitted a classification of the vocal response according to intensity as well as frequency.

Experimental fMRI Design
Before the onset of the experiment, both the tickler and the tickled person were instructed on the performance of each condition, and a trial run was conducted. During the scanning session, a friend (for 11 individuals) or the partner (for 7 individuals) of the participant stood in the scanning room and tickled or touched the right foot according to the particular stimulus condition (see Fig. 1). Each of the 3 tested conditions was indicated by a visual stimulus, consisting of a specific "smiley face" which was projected on separate screens for the tickler and for the tickled person. The latter watched the screen in a supine position via a mirror system. During tickling and laughter (T), the participants were manually tickled on the sole of the right foot and were encouraged to produce audible vocalizations. The same stimulation as for T was applied during tickling and inhibition of laughter (I), but the participants had to prevent themselves from it. During voluntary laughter (L), the subject was not tickled. Each of the 3 conditions was randomly presented 20 times, each lasting 6.2 s, and alternated with the presentation of a cross-signaled period of rest (11 s). To preserve the unpredictability of tickling, a red bar visible only to the tickler (F in Fig. 1) was randomly presented during T and I, urging stimulation by monotonous foot contact instead of tickling. This procedure ensured that the subjects were not prepared in advance for the real tickling stimulus, which was nevertheless applied 20 times. To minimize head movements and ensuing susceptibility artifacts in the fMRI signal, the participants held a wooden barbecue stick between their teeth during the course of the experiment, which did not interfere with laughter. After the session, the participants were asked to rate the mean sensation of tickling on a visual analogue scale (VAS, 1–10, 1 being the lowest score with no sensation of tickle and 10 the highest score).

Data Acquisition
Imaging was performed on a 3 T Scanner (VERIO, Siemens, Erlangen, Germany) with a 12-channel head coil. Functional images were obtained using a $T_2^*$-weighted echo planar imaging (EPI) sequence (repetition time: 2.2 s; echo time: 30 ms; flip angle: 90°), which embraced almost the entire brain in 24 contiguous axial slices (resolution: 3.0 x 3.0 x 3 mm, with a 1-mm gap). Images were additionally tilted by 30° relative to the anterior/posterior commissure (AC-PC line) to minimize susceptibility artifacts. During the functional session, 805 volumes were measured. Thirty-four phase and magnitude images were acquired in the same field of view and slice orientation, using a gradient echo (GRE) sequence with time repetition (TR) = 488 ms, time echo, TE(1) = 4.92 ms, TE(2) = 7.38 ms, and α = 60° to calculate a field map in which geometric distortions in the EPI images were nullified. An anatomical $T_1$-weighted, 3D Magnetization Prepared Rapid Gradient Echo image was acquired for each subject. The total number of sagittal anatomical slices amounted to 176 (TR = 1900 ms; TE = 2.52 ms; α = 90°; voxel size = 1 x 1 x 1 mm³).

Data Analysis
Data were analyzed using SPM5 software (Wellcome Department of Cognitive Neuroscience, London, England), running on Matlab version 7.4 (MathWorks Inc; Natick, MA). Unwrapping of geometrically distorted EPIs was performed in the phase-encoding direction using...
the FieldMap Toolbox for SPM5. Each individual scan was realigned to the first scan to correct for movement artifacts. EPIs were coregistered with the T1-weighted anatomical image. The coregistered T1-image was segmented and normalized to the Montreal Neurological Institute (MNI) template; the EPIs were resliced at 3 × 3 × 3 mm³. The resulting images were smoothed with a 9 × 9 × 9 mm³ Gaussian Kernel Filter (full-width at half maximum) to increase the signal-to-noise ratio. A temporal high-pass filter (128 s) was applied to eliminate slow-signal drifts. Movement parameters estimated during the realignment procedure were introduced as covariates into the model to control for variance due to head displacements. An event-related analysis was used to separately identify neuronal activity following stimulation for T, I, and L in each subject. The onset of the response to tickling was estimated to be 1 s after the visually presented start signal. This time delay included the presumed mean reaction time for the motor response of the tickler (approximately 200 ms) and the mean delay of the tickled person in discerning the tickling stimulus (approximately 800 ms). Since pain fibers are involved during tickling (Zotterman 1939; Lahuerta et al. 1990), the conduction may include slower pathways than after mere sensory stimulation. The beginning of the tickling sensation was chosen as the event of interest rather than the onset of audible laughter because we were interested in the specific brain processes that led to the outburst of laughter. The data were thus analyzed by modeling neuronal activity 1 s after the visual presentation of the respective condition. They were then adapted to the hemodynamic response function as supplied by SPM5. This procedure was adopted in the evaluation of all 3 conditions, even L. Voluntary laughter was continuous, and an evaluation of the early event was likely to capture the preparatory activity, which was considered to be most comparable to the T event. A one-sample t-test was implemented for the random-effect analysis at the group level. Main-effect of conditions T, I and L were considered to be statistically significant at a family-wise error rate of P < 0.05, using a correction for multiple comparisons across the entire brain volume. To demonstrate activity in the motor cortex, a region of interest (ROI) analysis was applied for T and L. Brain activity that corresponded specifically to ticklish laughter was established by performing a conjunction analysis (global mean) on a full factorial model of the contrast T versus I and L. Furthermore, each participant’s (intensity weighted) number of vocal responses to tickling was correlated with its corresponding fMRI-signal magnitude (as represented by the contrast image) during the condition T. This analysis was calculated by a simple regression (implemented in SPM5) for the whole brain. The thresholds for the 2 latter analyses were set at P < 0.001 (uncorrected). Parameter estimates (beta-values) for each experimental condition (T, I, and L) were derived from peaks of activation in the hypothalamus (centered at 3, −12, −15 (x, y, z) for the right, and −6, −6, −15 for the left hemisphere).

Results

Behavioral Results

During I, the vocal response upon request was strong (as defined in the Materials and Methods section), with an average number of 20 laughter events per participant. During T, the average score per subject was 9.75 (“weak” laughter: 11.5, weighted as 5.25; “strong” laughter: 4.50). In this situation, each participant laughed at least once during the scanning session. The average tickling stimulus was gauged as moderate with a mean rating of 6.7 on a scale of 1–10 (range: 3.3–9). No statistically significant correlation existed between the participants’ rating of tickling and the corresponding number of weighted laughter events (r(9) = 0.16; P = 0.52). During L, some of the participants adapted a strategy that evokes emotionally driven laughter, such as imagining a comical situation (2 participants) or recalled true laughter on their attempt to induce voluntary laughter (1 participant). This may have decreased the signal for ticklish laughter relative to that for voluntary laughter. During I, all participants were able to prevent themselves from laughing.

FMRI Results: Activation during Tickling and Laughter, Tickling and Inhibition of Laughter, and Voluntary Laughter

During T, stimulation of the sole of the right foot was associated with increased activation in the left primary sensory-motor cortex (representing the foot) and bilaterally in the secondary sensory-motor parietal operculum (including SII) and in the supplementary motor area (SMA) (Fig. 2, Table 1). ROI analysis within the S1/M1 revealed bilateral activation in the primary sensory-motor area (representing mimic musculature, larynx, pharynx, and diaphragm). Activity was also recorded in the frontal operculum, the thalamus, the putamen/pallidum, the hypothalamus (bilaterally), the anterior and posterior lobe of the right cerebellum, and the PAG. Activation in the visual regions was likewise enhanced as a response to the projected “smiley faces.” During I and L, activation occurred in similar regions of the brain to those described for T, with the exceptions that during I none was registered in the midbrain PAG, the cerebellum, the visual cortex, or the auditory cortex. During I, the cerebellum was activated bilaterally, but only in the posterior lobe. In contrast to the irregular laughter that was evoked during T, a vocal event was consistently recorded during L. Consequently, activities in the primary and secondary auditory cortex were recorded only in the latter situation. Neither I nor L activated the hypothalamus.

Increased Activation during Tickling and Laughter

To characterize the brain activity that is specifically associated with ticklish laughter the results for T were compared with those for the other 2 conditions: I and L (Fig. 4 and Table 2). A conjunction analysis of the 2 comparisons revealed a network for sensory integration of the right foot stimulation, which included the right anterior cerebellar lobe (lobules I–IV), the bilateral parietal operculum (including the left SII) and the left primary sensory-motor foot area. This analysis also revealed heightened activation for T in brain areas corresponding to the limbic system, namely, in the midbrain substantia nigra, the hippocampus and bilaterally the tuberolateral part of the hypothalamus, and the amygdala.

Correlation Analysis of Tickling and Laughter with Vocalizations

During T, a linear positive correlation between each participant’s (intensity weighted) number of vocalizations and its corresponding fMRI-signal magnitude occurred in the lateral-ventral portion of the PAG (Fig. 3 and Table 3, for localization see also Carrive and Morgan 2004). Vocalization-related activity was also observed in the right primary sensory-motor region for facial expression, in the right frontal and the left parietal operculum, in the bilateral medial thalamus, and in the right posterior pallidum. It has to be noted that the observed effect in the above-mentioned regions was specific for T. Namely, at the applied threshold of 0.001 (uncorrected), the number of vocalizations emitted during T did not correlate with fMRI activity during I or L, except in the left primary sensory-motor cortex during I.
Discussion

A comparison of the imaging signals for T with those for I, and L revealed hypothalamic activity exclusively in the former situation. This finding confirms our hypothesis that the limbic pathway of vocalization is critically involved in ticklish laughter. Tickling and laughter is also associated with specific activities in higher-order sensory-motor areas (SII and the cerebellum), possibly paving the way to the deliberate control of the ensuing vocalization. Unexpectedly for such a primitive involuntary reaction, the same primary sensory-motor and premotor regions for the control of vocalization are activated in all 3 situations (T, I, and L). This finding indicates that the spontaneous laughter elicited by tickling is also under the control of the voluntary motor cortical pathway.

**Tickling Activates the Motor Cortical Pathway of Vocalization**

In all 3 situations (T, I, and L), a common activation of the primary sensory-motor cortex predictably involved brain areas that represent the face, the tongue, larynx and pharynx (Rolandic operculum), and the diaphragm (Maskill et al. 1991; Lotze et al. 2000; Brown et al. 2009). In contrast to previous observations relating to voluntary smiling or laughter (Wild et al. 2003), which were restricted to the somatotopic facial area, our study reveals the broad pattern of effectors that is activated during ticklish laughter. Consistent with the findings of Wild et al. (2003), activity occurred in the SMA and the frontal operculum, which have been previously shown to encode various other orofacial functions, such as song production (Kleber et al. 2007) and speech (Brown et al. 2008). Taken together, our findings indicate that the motor cortical pathway is involved not only in voluntary but also in involuntary laughter, such as is evoked during tickling, and even in inhibition of laughter. The result for the latter 2 situations deserves a special comment. Inappropriate, uncontrollable laughter has been observed in patients with lesions of the motor cortical pathway, thereby implicating this system in inhibitory control of vocalization (Wild et al. 2003). Activation of the motor cortical pathway during T and I signalizes inhibitory control which may regulate the vocal response according to the situational context. Moreover, Table 1 demonstrates that the neuronal response during I is even higher than the one during T: since laughter must be actively inhibited, cortical motor control is augmented. Our data reveal no evidence of prefrontal activity during I (see Table 1), although inhibitory control at the cognitive level is a key function of this region (Konishi et al. 1999; de Zubicaray et al. 2000). In a previous imaging study on humorous smiling and laughter, involvement of the left prefrontal cortex was reported and attributed to the perception and cognitive processing of stimulus characteristics (Wild et al. 2006). Inspection of our data at a lower threshold level (P < 0.0001 [uncorrected]) reveals though, activity in the left inferior prefrontal cortex (brain region [BA] 10, MNI coordinates: -39, 45, 12 (x, y, z), z-value = 4.25) during I that was not observed during T and L. Hence, we may conclude that, during I, cognitive strategies are adopted to inhibit the reaction to

![Figure 2. Activation patterns during T, I, and L. Three-dimensional top (first row) and lateral (from the right: R) views (second row) of the brain, and sagittal slices through the right hemisphere (third row). Activity is apparent in the primary sensorimotor areas (S1/M1) representing the diaphragm (D), mimic musculature and larynx (MIM), and the pharynx (Rolandic operculum: RO), as well as in the SMA and in the frontal operculum (FO). Activity in the left-foot area (FOOT) and in the parietal operculum (PO) is observed only during T and I. The auditory cortex (Heschl’s gyrus: HG) is activated only during L. T: P < 0.0001 (uncorrected); I: P < 0.0001 (uncorrected); L: P < 0.05 (FWE-corrected). On each scale, the threshold level (T-value) for observed activity is indicated.](image-url)
Ticklish Laughter Activates Specific Components in the Limbic Pathway of Vocalization

The neuronal equivalent of ticklish laughter was analyzed by correlating the activity during T with its corresponding laughter scores (Table 2 and Fig. 3) and by comparing T with the conditions in which laughter was either forbidden I or voluntary L (Table 2 and Fig. 4). A positive correlation occurred in the midbrain PAG, thereby confirming the role of this region as a crucial relay station in the limbic pathway of vocalization (Graham Brown 1915; Dabby et al. 2004; Jürgens 2009), since the PAG was also activated during the main conditions T and L (see Table 1). Our data support existing evidence of this region’s involvement in initiating and controlling the intensity of all forms of vocal reaction (Larson 1991; Davis et al. 1996).

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In contrast, our comparative analysis revealed hypothalamic activity to be specific for ticklish laughter alone. Behavioral and visceral reactions are intrinsically regulated in the hypothalamus, which is anatomically connected to the PAG (Veazey et al. 1982; Semenenko and Lumb 1992; Saper 2004). Specifically,

tickling. The absence of activity in the corresponding area of the right hemisphere remains unexplained because older data suggest that naturally arising motor responses to emotional stimuli, notably laughter and smiling, are under the control of the right prefrontal cortex (Shammi and Stuss 1999). Wild et al. (2006) suggested that action of the right inferior prefrontal cortex is essential to restrain laughter but not to effect it. Hence, our results indicate that, during I, the right prefrontal cortex is not involved in the suppression of the vocal response to tickling; the seat of this inhibitory response lies at other cortical levels.

Finally, activity in the posterior cerebellar lobe occurred only in the 2 conditions that involved laughter (T and L) but not during I. This finding is consistent with the role of the cerebellum in the on-line adjustment of ongoing motor responses rather than in preparatory aspects (during I) relating to these (Bloedel 1992).

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hypothalamic nuclei comprise the largest input to vocalization areas, and activity stemming therefrom stimulates subregions of the PAG to evoke calls of hedonic or aversive character (Altafullah et al. 1988; Dujardin and Jürgens 2006). The hypothalamus can also regulate physiological phenomena, such as heart rate that accompany genuine laughter (Kuniecki et al. 2003) and emotional (Bachorowski and Owren 2003) as well as sympathetic arousal (Hecker 1873; Fry 1994). Activation of the hypothalamus coincided with that of the amygdala and the hippocampus (Table 2), which are believed to form part of a network that integrates sensory information and promotes appropriate visceral and behavioral reactions (Price 2005). Both structures project to the hypothalamus and from there to lower brainstem regions, such as the PAG (Leonard and Scott 1971; Poletti et al. 1973). The hypothalamus thus appears to hold a pivotal position between the cortical perception of stimuli and the regulation of behavior (Steams 1972). It may either facilitate or trigger the stereotyped pattern of ticklish laughter via the PAG, and modulate the physiological bodily changes that are associated with it, if tickling was applied under appropriate consensual conditions.

Interestingly, the increase in hypothalamic activity that was evoked during ticklish laughter was confined to the tuberal-lateral portion (Fig. 4), which accords with the findings of several recent studies relating to humor processing in both normal persons (Mobbs et al. 2003; Watson et al. 2007) and narcoleptic/cataplectic patients (Reiss et al. 2008; Schwartz et al. 2008). Since the measured response occurs at an anatomical location that contains the hypocretin/orexin-secreting neurons, their involvement has been suspected (Schwartz et al. 2008).

In this connection, it is worth mentioning that a wealth of neurological data suggests that the hypothalamus generates laughter (Davison and Kelman 1939; Martin 1950). Benign tumors (hamartomas) developing in the tuberal part of the hypothalamus of children, often lead to gelastic (laughing) seizures (Valdueza et al. 1994; Delandale and Fohlen 2003) and also vascular pathologies, such as aneurysm and bleeding in the region of the circle of Willis (Martin et al. 2003). Whilst hamartoma may generate intrinsic epileptic activity (Kuzniecky et al. 1997), vascular pathologies probably trigger laughter via a mechanical irritation or compression of the tuberal-lateral hypothalamus. This contention is supported by the observation that a swabbing of the floor of the third ventricle or pressing the infundibulum during the resection of ependymal cyst or craniopharyngeoma produced high mood and induced the patient to burst into laughter (Foerster and Gagel 1934).

The increased blood oxygen level-dependent (BOLD)-signal in the tuberal-lateral part of the hypothalamus could therefore be localized in the hypocretin/orexin cells rich region, explaining the arousal state typically accompanying tickling (Fry 1994). But the signal seems to occur more laterobasally, closer to the lateral tuberal nucleus (LTN), an entity of unknown function particularly well developed in humans (Le Gros Clark 1938), which is target of neuropathological changes in Huntington and Pick disease (Braak and Braak 1998). The human LTN may have its homologous in the rodent PVT, a newly recognized nucleus in the lateral hypothalamus of rodents (Gerig and Cello 2007; Girard et al. 2011; Meszar et al. 2012). In the future, the question of the exact intrahypothalamic localization of the BOLD signal could be resolved by high-resolution fMRI using the ticklish-laughter paradigm.

Activation of the anterior cingulate gyrus occurred during L and I, but not during T. Moreover, during L it was coactivated with the PAG (Table 1). fMRI investigations (Liu et al. 2010)
and case studies in humans (Rubens 1975; Jürgens and Kirzinger 1982), as well as lesioning experiments in monkeys (Sutton et al. 1974), afford evidence that the anterior cingulate gyrus is involved in motivated vocal output. And the findings of another study with monkeys indicate that this region controls vocalization via the PAG (Jürgens and Pratt 1979). The activity in the anterior cingulate gyrus that was observed during L and I reflects the participant’s efforts to control vocalization. Ticklish laughter, on the other hand, appears to be produced independently from these motivational circuits and to involve unconscious pathways relying on the hypothalamus.

**Ticklish Laughter Activates Higher-Order Sensory Regions**

Although tickled persons may enjoy the experience, they typically engage in a defensive reaction, which may include a withdrawal of the tickled body part, a feeling of slight discomfort (Ruggieri and Milizia 1983), and laughing or giggling. The idea of a protective mechanism is supported by the finding that tickle-induced laughter depends partially on an intact pain sensation (Zotterman 1939; Lahuerta et al. 1990). Our comparison of T with the 2 other conditions (I and L) revealed the brain structures that could be involved in such specific processes, for example, the right anterior cerebellar lobe and the left parietal operculum SII (lateralized in accordance to tickling on the right foot, see Table 2). Our findings accord with those of previous imaging studies that focused on the sensory representation of tickling (Blakemore et al. 2000; Carlsson et al. 2000). Increased activity in the secondary sensory SII has been associated with the conscious perception of both skin contact (Burton et al. 1997) and pain (Coghill et al. 1999). The cerebellum is integrating sensory feedback into an online correction of movement (Bloedel 1992). There are also indications that cerebellar activity is augmented in the presence of a mismatch between predicted and actual sensory perceptions (Restuccia et al. 2007; Bubic et al. 2009) and that it is implicated in the processing and anticipation of pain (Ploghaus et al. 1999). Furthermore, the main input nucleus to the cerebellum—the inferior olive—cannot distinguish between tactile sensory and pain-fibers inputs (Oscarsson and Sjolund 1977), and may thus transmit ambiguous information via the cerebellar climbing fibers during tickling. Additional cortical processing may be necessary to qualify the character of the sensory stimulation. If so, then a functionally relevant mechanism of mismatch detection and resolution could lie within the sequence of actual tickling and the prediction of pain and would entail activity in the cerebellum and the SII. Ticklish laughter would occur when this mechanism was activated to a sufficient degree. Such an idea is reminiscent of models of humor processing which have proposed that the recognition and resolution of cognitive incongruities between stored (anticipated) and actual information is necessary for the detection of humor and the ensuing burst of laughter (Suls 1972; Attardo 1997).

Previous studies relating to humorous laughter have revealed an involvement of the same limbic-related regions that we have observed in the context of ticklish laughter (Mobbs et al. 2003;
Wild et al. 2003; Watson et al. 2007), thereby indicating a tendency to recruit a similar motivational network in both situations. However, the medial prefrontal cortex and the nucleus accumbens are activated only during the processing of humor (Goel and Dolan 2001; Samson et al. 2008; Samson et al. 2009). This network appears to be essential for self-referential processes (Luan Phan et al. 2005; van der Meer et al. 2010). In monkeys, its output feeds the hypothalamus and the PAG (An et al. 1998; Ongur et al. 1998). Hence, humorous stimuli probably generate a subjective sensation of mirth, which contributes to trigger the laughter response that is mediated by the hypothalamus and the PAG. The tactile stimulation that produces the specific sensation of tickling (possibly in the SII and the cerebellum) would activate the same subcortical centers.

Our study highlights the pattern of brain activity that characterizes ticklish laughter. Several cortical and subcortical centers are implicated, and the pattern of activity resembles that evoked by humorous laughter, excepting an involvement of the medial prefrontal cortex and the nucleus accumbens in the latter situation only. Hence, whilst the subjective feeling of mirth and reward may be relevant in humorous laughter, they play but a subsidiary role in ticklish laughter. Nevertheless, the hypothesis of Darwin (1872) and Hecker (1873), that ticklish laughter forms the primitive building block supporting humorous laughter, is confirmed by our results.

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