

Dopamine in the medial amygdala network mediates human bonding

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Edited by Linda M. Bartoshuk, University of Florida, Gainesville, FL, and approved January 6, 2017 (received for review July 26, 2016)

Research in humans and nonhuman animals indicates that social affiliation, and particularly maternal bonding, depends on reward circuitry. Although numerous mechanistic studies in rodents demonstrated that maternal bonding depends on striatal dopamine transmission, the neurochemistry supporting maternal behavior in humans has not been described so far. In this study, we tested the role of central dopamine in human bonding. We applied a combined functional MRI-PET scanner to simultaneously probe mothers' dopamine responses to their infants and the connectivity between the nucleus accumbens (NAcc), the amygdala, and the medial prefrontal cortex (mPFC), which form an intrinsic network (referred to as the "medial amygdala network") that supports social functioning. We also measured the mothers' behavioral synchrony with their infants and plasma oxytocin. The results of this study suggest that synchronous maternal behavior is associated with increased dopamine responses to the mother's infant and stronger intrinsic connectivity within the medial amygdala network. Moreover, stronger network connectivity is associated with increased dopamine responses within the network and decreased plasma oxytocin. Together, these data indicate that dopamine is involved in human bonding. Compared with other mammals, humans have an unusually complex social life. The complexity of human bonding cannot be fully captured in non-human animal models, particularly in pathological bonding, such as that in autistic spectrum disorder or postpartum depression. Thus, investigations of the neurochemistry of social bonding in humans, for which this study provides initial evidence, are warranted.

dopamine | maternal behavior | social affiliation | network connectivity | humans

Early social bonding with a primary caregiver is necessary for mental and physical health, whereas the absence of such bonding is a clear risk factor for adult illness (1). However, despite potentially enormous implications, to date the science of mother–infant bonding relies mostly on nonhuman animal models.

Research on nonhuman animals indicates that maternal bonding involves the nucleus accumbens (NAcc), amygdala, and medial prefrontal cortex (mPFC). In rodents, oxytocin and dopamine act in the amygdala and NAcc (2) to regulate maternal appetitive behaviors. In humans, functional MRI (fMRI) studies have verified that NAcc activity increases consistently when mothers gaze at their infants (3). Moreover, the NAcc and amygdala activity have been linked to the quality of maternal behavior (4). Mothers who were sensitive to their infants' cues for social engagement and who adjusted their own behavior to meet those needs (referred to as "mother–infant synchrony"), showed greater activations in the left NAcc and lower activation in the right amygdala when viewing films of their infants than did nonsynchronous mothers (4). In agreement with the animal studies, oxytocin has been implicated in human maternal behavior, so that synchronous mothers show a stronger link between levels of circulating plasma oxytocin and NAcc fMRI activations when viewing films of their infants (4). Moreover, oxytocin administration increased activations of the ventral tegmental

area, which sends dopaminergic projections to the NAcc as part of the mesolimbic system, in response to infant stimuli (5).

In this study, we extend our knowledge of the neural basis of bonding by demonstrating that dopamine is associated with human bonding. Bonding behavior was assessed in this study using indices of mother–infant synchrony. We also examined dopamine responses and intrinsic connectivity of the striatum with the broader medial amygdala network (Fig. 1) that connects the NAcc to the medial amygdala, rostral hypothalamus, ventromedial prefrontal cortex (vmPFC), subgenual anterior cingulate cortex (sgACC), and posterior cingulate cortex (PCC). This network's hubs were consistently linked to human maternal bonding (for review, see ref. 3; also see refs. 4 and 6). Moreover, atypical maternal behavior, as seen in patients with postpartum depression (PPD), is associated with attenuated maternal responses in the striatum (7), rapid striatal attenuation to reward (8), and disrupted connectivity between the right amygdala and the PCC (9). We examined the connectivity within the medial amygdala network using blood oxygenation level-dependent (BOLD) signals acquired during fMRI and examined dopamine function with the radiolabeled ligand [¹¹C]raclopride during PET imaging while a mother watched a film of her own infant and a film of an unfamiliar infant. Whole-brain network investigation during real experiences, using simultaneous probing of PET and fMRI, facilitates the mechanistic understanding of how multifaceted brain function relates to complex human behavior.

Significance

Early life bonding in humans has critical long-term implications for health, productivity, and well-being in society. Nonetheless, neural mechanisms of bonding are typically studied in rodents, and no studies to date had examined the neurochemistry of human social affiliation. This study utilizes a state-of-the-art technology to demonstrate that human maternal bonding is associated with striatal dopamine function and the recruitment of a cortico–striatal–amygdala brain network that supports affiliation. The simultaneous probing of neurochemical responses and whole-brain network function in mothers watching their infants provides a unique observation into an "affiliating brain." These results advance the mechanistic understanding of human social bonding and promote basic and clinical research in social neuroscience, development, and psychopathology.

Author contributions: S.A., B.C.D., C.C., and L.F.B. designed research; S.A., T.R., S.S., and J.M.H. performed research; S.A., A.T., T.R., R.F., B.C.D., C.C., and L.F.B. analyzed data; and S.A., T.R., C.C., and L.F.B. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

Freely available online through the PNAS open access option.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1612233114/-DCSupplemental.

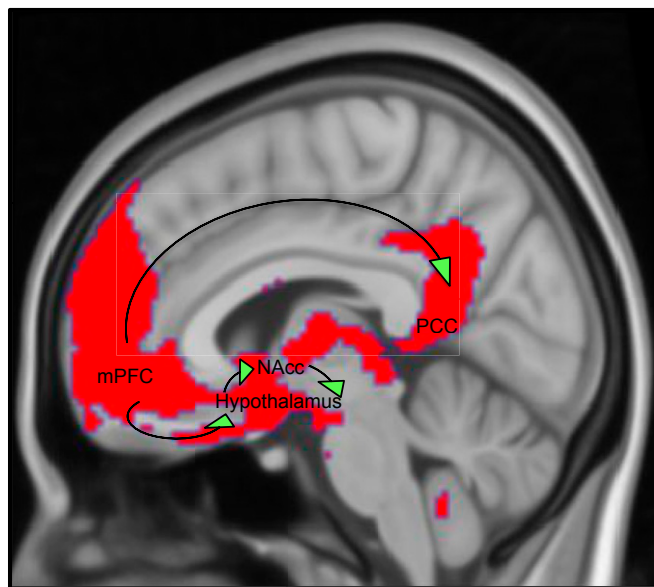


Fig. 1. The medial amygdala network (in red) includes the medial sector of the amygdala along with a set of connected regions in the NAcc, rostral hypothalamus, vmPFC, PCC, and sgACC (25). The map is based on 150 healthy adults and reflects a whole-brain intrinsic connectivity analysis seeded in the right medial amygdala. Previous research shows that stronger intrinsic connectivity within this network is associated with larger and more complex social networks (25). The connectivity and dopamine response within the network were tested for association with maternal bonding behavior and plasma oxytocin levels. We hypothesized that dopamine secretion within the medial amygdala network recruits the network to support human bonding (green arrowheads). We predicted that synchronous mothering would be associated with an enhanced dopamine response within the network that specifically signifies the own-infant over the unfamiliar-infant condition.

Nineteen mother–infant dyads completed the study. At the outset, we visited families at home. We filmed their unconstrained interactions and the infant’s solitary play. The interaction films were behaviorally coded to evaluate mother–infant synchrony, and the infant solitary play films were used as stimuli during brain imaging (4). Mothers then participated in two simultaneous PET–fMRI scanings in which we tracked the change in the binding of [¹¹C]raclopride to D2-type dopamine receptors while mothers viewed films of their own infant and films of an unfamiliar infant as control. We then measured task-independent changes in BOLD signal, from which we estimated the degree of intrinsic connectivity of each mother’s medial amygdala network. Just before the brain imaging session, blood was collected for oxytocin analysis.

Results

Behavioral Results. Bonding behavior was evaluated with two indices of mother–infant synchrony: mother–infant vocalization synchrony and maternal attunement. Mother–infant synchrony is the temporal contingency of behavior between a mother and her infant. To synchronize, a mother needs to be sensitive to her infant and adjust her behavior dynamically according to the infant’s cues.

Mother–infant vocalization synchrony measures the temporal matching of mother–infant vocalizations. As part of typical human bonding behavior, mothers and infants tend to synchronize their vocal communication (1). We calculated vocalization synchrony as the percentage of time during a 2-min unconstrained interaction in which mothers and infants were vocally synchronized. In our sample, mothers and infant synchronized vocalizations for 0–13% of the time, with a mean of 4% and median of 2.36%. Using the median, we divided the sample into two groups: high and low vocalization synchrony groups.

Maternal attunement is a broader measure of synchrony that traces how well the mother accommodates her vocal stimulation to the infant’s affect and social engagement. Maternal voice helps regulate the infant’s attention and affect (10). As such, mothers attune their vocal communication not only to the infants’ vocalizations but also to a broader set of affective cues, such as infants’ arousal levels and social engagement. We refer to this broader aspect of synchrony as “maternal attunement.” Maternal attunement measured the time in which mothers provided vocal stimulation while the infant was showing signs of positive affect (the infant is content and socially engaging). In our sample, maternal attunement ranged from 0–27% of the time in the 2-min unconstrained interaction, with a mean of 10.7% and a median of 10.8%. Age, race, or education level did not predict maternal attunement or mother–infant vocalization synchrony.

Imaging Results. Analyses with behavior and PET: Association between mother–infant vocalization synchrony and dopamine response. The imaging paradigm included two scans. Each mother viewed a film of her own infant during one scan (own-infant condition) and a film of an unfamiliar infant during another scan (unfamiliar-infant condition) as a control; the order was counterbalanced. To assess mothers’ endogenous dopamine response in each condition, we injected to the mothers with [¹¹C]raclopride 10 min into each film. Once injected, [¹¹C]raclopride binds specifically to free D2 receptors that are not occupied by endogenous dopamine molecules. Our primary outcome measure was [¹¹C]raclopride-nondisplaceable binding potential (BPnd) (11). BPnd indirectly indexes the levels of endogenous neural dopamine response: A decrease in [¹¹C]raclopride BPnd signifies a proportional increase in endogenous dopamine, and vice versa (12). Each mother’s relative dopamine response was calculated as the percentage change in endogenous dopamine response in the own-infant condition compared with the unfamiliar-infant condition using the formula $-\frac{[\text{raclopride BPnd}(\text{own infant}) - \text{raclopride BPnd}(\text{unfamiliar infant})]}{[\text{raclopride BPnd}(\text{unfamiliar infant})]} \times 100$.

Our results showed that mother–infant synchrony scores are associated with maternal dopamine responses and that high-synchrony mothers have a dopaminergic preference for their own infants. In a multivariate general linear model, relative dopamine responses in the high vocalization synchrony group were significantly higher in the right hemisphere regions of interest (main effect; $n = 19$, $F = 4.6$, $P < 0.02$), with greatest effects in the right pallidum ($F = 11.1$, $P < 0.004$) and right NAcc ($F = 9.3$, $P < 0.007$) (Fig. 2A; see Table S1 for a complete list of regions and effects). The left hemisphere contralateral regions did not display such a group difference in [¹¹C]raclopride BPnd. In the low-synchrony mothers, most mothers had a stronger dopamine response to the unfamiliar infant (for individual data see Fig. S1; Fig. S2 demonstrates that mothers with a dopaminergic preference in the right pallidum to their own infants are more synchronous). The scatterplots in Fig. 2B and C show the correlation between the individual differences in dopamine responses in the right pallidum and NAcc and vocalization synchrony scores.

Analyses with behavior and fMRI: Association between intrinsic connectivity within the medial amygdala network and maternal attunement. Unlike vocalization synchrony, which was linked to PET dopamine responses, maternal attunement was strongly associated with intrinsic connectivity in the medial amygdala network. Mothers who were more attuned to their infants had a more cohesive medial amygdala network (Fig. 3).

Infants’ ages were correlated with the dopamine responses in the right pallidum ($r = 0.49$, $P < 0.03$) and right amygdala ($r = 0.5$, $P < 0.01$) but were not correlated with the connectivity in the medial amygdala network ($r = -0.04$, $P < 0.8$), vocalization synchrony ($r = 0.3$, $P < 0.2$), or attunement ($r = 0.2$, $P < 0.5$). After controlling for infant age, vocalization synchrony was still correlated significantly with dopamine responses in the right pallidum ($r = 0.4$, $P < 0.03$). We also confirmed that infants’ age

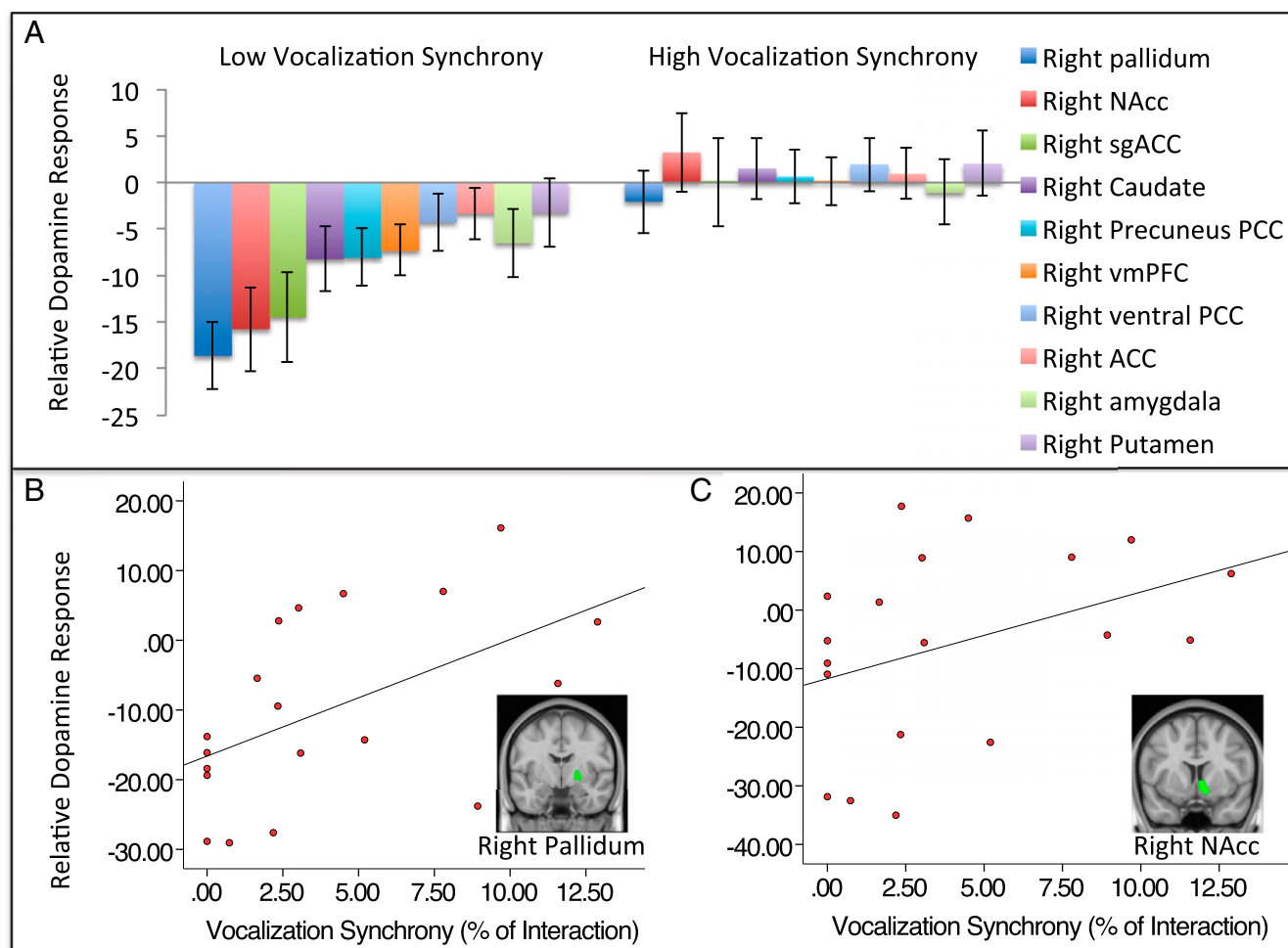


Fig. 2. Evidence that behavioral synchrony between mothers and infants is associated with maternal striatal dopamine responses to the infant. (A) A general linear model analysis demonstrates that high-synchrony mothers have higher relative dopamine responses than low-synchrony mothers in the own- vs. unfamiliar-infant comparison (indexed by percent [^{11}C]raclopride BPnd change). Regions of interest are presented according to their effect size (see the full list of regions in Table S1). Error bars represent the SEM. Mothers were assigned to a high- or low-synchrony group based on the vocalization synchrony median (2.36%). (B and C) Individual differences in vocalization synchrony in mothers are positively correlated with dopamine responses (indexed by the percent [^{11}C]raclopride BPnd change) in the right pallidum (one-tailed, $n = 19$, $r = 0.517$, $P < 0.012$) (B) and right NAcc (one-tailed, $n = 19$, $r = 0.378$, $P < 0.055$, trending) (C).

did not mediate the relationship between vocalization synchrony and right pallidum dopamine (Sobel test not significant, $z = 1.03$, $P < 0.3$). These analyses confirm that mother–infant vocalization synchrony is associated with maternal dopamine responses across infants' ages.

Analyses with fMRI and PET: Association between medial amygdala network connectivity and dopamine responses in mothers. Connectivity within the medial amygdala network was associated with in-network dopamine responses. Mothers with stronger medial amygdala network connectivity showed increased in-network endogenous dopamine responses in the right sgACC, right amygdala, and right NAcc while watching their own infants but not while watching an unfamiliar infant, (Fig. 4). A trend of correlation also was evident for the right PCC ($P < 0.06$). Dopamine responses in the contralateral left regions were not correlated with intrinsic connectivity in the left hemisphere medial amygdala network.

Analyses with plasma oxytocin levels. Peripheral oxytocin was measured in circulating plasma because there is still no reliable specific radiotracer for human oxytocin receptors (13). Central oxytocin is secreted as a neurotransmitter via axon terminals in the brain, whereas peripheral oxytocin is secreted via the pituitary gland into the blood circulation as a hormone. Studies in nonhuman animals (14) showed no direct correlation between levels of peripheral and

central oxytocin. Nonetheless, numerous studies reported a link between plasma oxytocin and human behavior (for review see ref. 15). In this study peripheral oxytocin and its relations to central dopamine and behavior were evaluated in an exploratory way. Plasma for oxytocin analysis was available for 17 subjects and ranged from 43–370 pg/mL with a mean of 195 pg/mL. Levels of plasma oxytocin predicted medial amygdala network connectivity with a negative correlation coefficient (Fig. 5). Plasma oxytocin was positively correlated with vocalization synchrony ($r = 0.5$, $P < 0.03$) and showed a trend of correlation with dopamine responses in the left NAcc in the own-infant condition ($r = 0.36$, $P < 0.07$).

Discussion

The results of this study demonstrate that human maternal bonding is associated with dopamine responses in the NAcc and pallidum and with the strength of intrinsic connectivity within the medial amygdala network. Moreover, stronger intrinsic connectivity in the medial amygdala network is associated with increased within-network dopamine levels tested simultaneously and with lower plasma oxytocin levels. Together, these data provide evidence indicating that dopamine is involved in human bonding.

When humans interact, their appetitive behaviors synchronize (1). Synchrony has been shown to improve prosocial behavior in

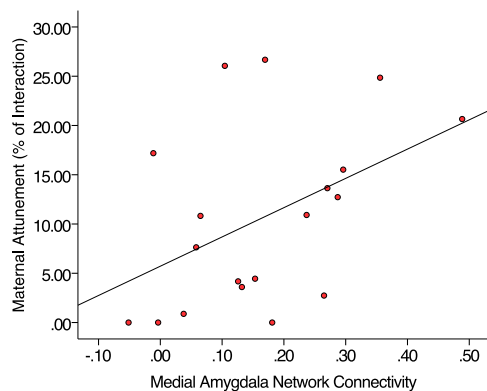


Fig. 3. Mothers with a stronger medial amygdala network are more attuned to their infants ($n = 19$, $r = 0.46$, $P < 0.02$, one-tailed). Medial amygdala network connectivity is represented as Fisher's r -to- z transformed Pearson correlation coefficients between the right medial amygdala seed and the rest of the network's nodes. Maternal attunement was measured as the percent of time during a 2-min interaction in which mothers provided positive vocal stimulation to their infants while the infants were content and socially engaged.

infant macaques (16, 17), among healthy children (1), and in children with autistic spectrum disorder (18–21). The evidence presented here, which links synchrony to dopamine, provides initial evidence that dopamine is involved in the prosocial effects of synchrony. Behavioral synchronization demands dynamic adjustment of one's behavior to the dyadic partner. In this study, synchronous mothers had stronger dopamine responses to their own infant in

the NAcc and pallidum. In rodents, dopamine in the NAcc regulates appetitive maternal behaviors (2). In response to a salient event, dopamine in the NAcc releases the pallidum from GABA inhibition to disinhibit (or activate) motor pathways that execute appetitive behavior toward the pups (2). The results of this study, linking human behavioral synchronization to dopamine, join the rodent literature and mark dopaminergic function in the NAcc and pallidum as a regulatory pathway of appetitive bonding behavior in humans.

The infants' films are salient social stimuli that elicit striatal dopamine responses. However, unlike high-synchrony mothers, low-synchrony mothers showed increased dopamine responses in the unfamiliar-infant condition. One possible explanation for this finding could be the novelty of the unfamiliar infant (22). Instead, in synchronous mothers dopamine responses to the mother's own infant are stronger than those to a novel infant, possibly because the mother's own infant is extremely salient to her (1). In the clinical realm, such salience-specificity is known as "primary maternal preoccupation" (23), which describes a mother's complete focus on her infant while disregarding all distractions (23). Low maternal preoccupation is linked to PPD and nonsynchronous parenting (23). In our sample none of the mothers had a psychiatric diagnosis; however, low-synchrony mothers did not show differential dopamine responses that favor their own infants. This lack of a differential response could represent a relative deficit in their infant saliency with regard to other infants. However, beyond increased saliency, the dopaminergic patterns measured here might represent the selectivity of maternal attachment. Primate mothers and sheep show a selective bonding to their own young (24). Additionally, D2 receptors in the NAcc are important for the

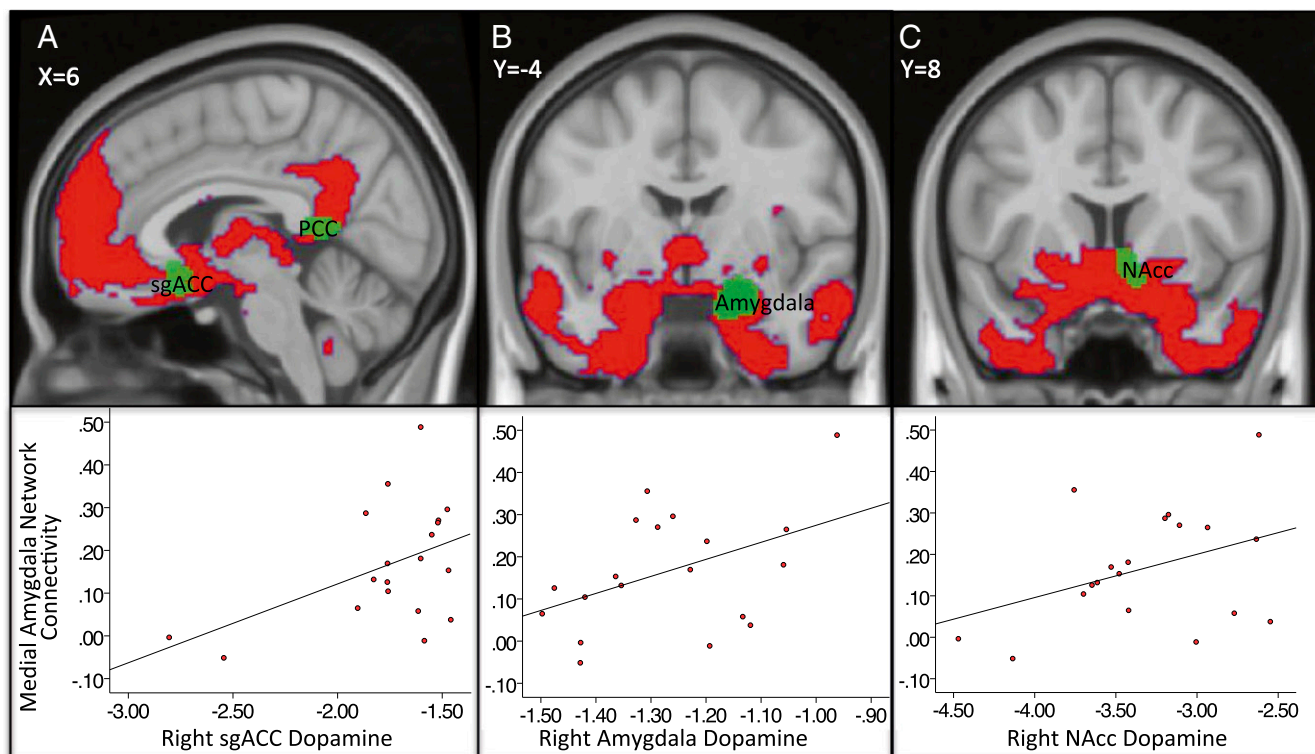


Fig. 4. Stronger intrinsic connectivity in the medial amygdala network is predicted by increased in-network dopamine responses during the own-infant condition. (A–C, Upper) Intrinsic connectivity maps of the medial amygdala network (in red), overlaid with regions of interest for PET analysis [manually illustrated in green, according to FreeSurfer segmentation atlases (34)] in which [^{11}C]raclopride BPnd is correlated with the network connectivity. (A–C, Lower) The Pearson one-tailed correlation graphs ($n = 19$). (A) Right sgACC ($r = 0.45$, $P < 0.03$). (B) Right amygdala ($r = 0.455$, $P < 0.02$). (C) Right NAcc ($r = 0.38$, $P < 0.05$). In the x axes, an increase in dopamine responses during the own-infant condition is indexed by a decrease in [^{11}C]raclopride BPnd. In the y axis, medial amygdala network connectivity is represented as Fisher's r -to- z transformed Pearson correlation coefficients between the right medial amygdala seed and the rest of the network's nodes.

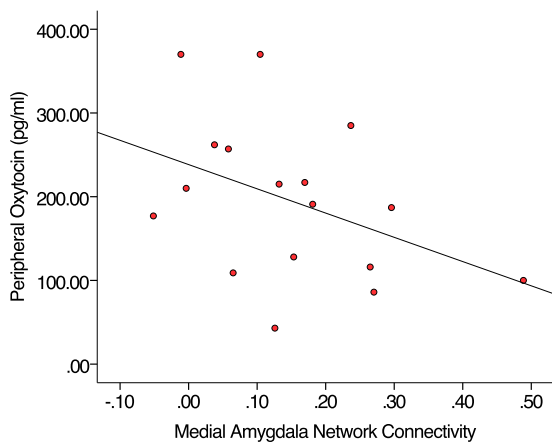


Fig. 5. Plasma oxytocin negatively predicts connectivity in the medial amygdala network (two-tailed, $n = 17$, $r = -0.415$, $P < 0.049$). In the x axis, medial amygdala network connectivity is represented as Fisher's r -to- z transformed Pearson correlation coefficient values between the right medial amygdala seed and the rest of the network's nodes.

formation of selective social bonds (2). Accordingly, in humans the increased D2-mediated dopaminergic responses in the NAcc in the own-infant condition as compared with an unfamiliar-infant condition may reflect the involvement of D2 receptors in a mother's selective attachment to her own child.

This study reports a finding for a second key element in human maternal bonding: the medial amygdala network. Previous studies have shown that the strength of connectivity within the medial amygdala network is a reliable predictor for social affiliation (25). Our results extend those findings by demonstrating that the network is specifically involved in maternal bonding. Moreover, our results extend previous studies on the neural basis of maternal bonding (4, 6) by showing that bonding behavior relies not on the discrete function of the NAcc, amygdala, and mPFC (3) but instead on the synchronous firing in these regions as a network. The medial amygdala network includes the medial-rostral hypothalamus (6), which contains the medial preoptic area (MPOA) (26). The MPOA has a dominant role in maternal behavior in every mammalian species examined experimentally (24). Our study provides evidence for the involvement of this region in human bonding. This evidence helps integrate animal and human studies conceptually and suggests that homologous striatal circuitry has evolved to include broader neural connections with the human cortex. The medial amygdala network possibly supports synchronous affiliation by coordinating two functions: reward and mentalization. Synchrony appeared to be intrinsically rewarding to humans and nonhuman primates and to activate reward circuitry (27, 28). In addition to reward, synchrony relies on mentalization (i.e., the ability to represent the partner's intentions and anticipate behavior) (4). The medial amygdala network includes subcortical reward regions, such as the NAcc, hypothalamus, and amygdala, which are potentially important for motivating and regulating behavior. The network also includes cortical regions, such as the vmPFC, sgACC, and PCC, which are consistently reported to support mentalization (9). Both the subcortical and cortical regions, and particularly the connectivity between them, promote synchronous bonding by coordinating cortical circuitry that supports mentalization to striatal reward circuitry that supports behavioral regulation. Linking behavior to dopamine in cortical regions of the medial amygdala network extends the contribution of this study beyond the rodent literature because it marks a possible role for dopamine in the higher social cognition needed for human bonding.

Previous findings have suggested a possible mechanistic role for dopamine in intrinsic network function. For example, several

studies showed that patients with Parkinson's disease, who suffer from a dopaminergic deficiency, have abnormal default-mode network connectivity, which is restored after treatment with the dopaminergic precursor L-dopa (29). In healthy humans, L-dopa increased the connectivity in several networks, and the dopaminergic antagonist haloperidol decreased the connectivity in those networks (30). This evidence converges with the results of this report to support a possible role for dopamine in network regulation and thus in human cognition. This finding could have important implications for neuropathologies of the dopaminergic system, including Parkinson's disease, schizophrenia, addiction (30), and possibly social dysfunction and PPD.

The analysis of plasma oxytocin yielded intriguing results. Oxytocin, which is considered a "prosocial" hormone (2), was negatively correlated to the medial amygdala network, which is considered a prosocial network (25). Animal models revealed that central oxytocin projections from the hypothalamus to the striatum have both inhibitory and excitatory pathways affecting dopaminergic receptors in the NAcc and amygdala [summarized in figure 3 from Numan and Young (2)]. The negative correlation between oxytocin and the medial amygdala network requires further investigation of the oxytocin-dopamine interplay in humans.

Studying postpartum mothers in a behavioral PET-fMRI paradigm involved some limitations. Mothers' responses to their own infants were compared with their responses to unfamiliar infants and are thus relative. Future studies should administer an additional control condition measuring baseline levels of raclopride BPnd. Moreover, previous studies have observed both left and right NAcc involvement in maternal synchronization, but this study only observed the right NAcc. Recent studies suggest a role for D2 asymmetry in motivation, with D2 receptors in the right hemisphere involved in behavioral regulation, which is important for synchrony. Future mechanistic studies should evaluate a possible lateralization in striatal function. Additionally, animal models do not show that dopamine acts in the pallidum to regulate maternal behavior, but there is strong evidence that dopamine acts on the MPOA to stimulate rodent maternal behavior (31). The human medial-rostral hypothalamus, where the MPOA is located (26), is medially adjacent to the pallidum, so it is possible that the pallidal dopamine observed here actually reflects dopamine transmission into the rostral hypothalamus. Future investigation of the rostral hypothalamus in human bonding, using neuroimaging with improved spatial resolution, is of high importance because of this brain area's homology to rodents' MPOA. Importantly, these methods are inherently correlational and cannot provide information about the causal nature of the reported relationships. Moreover, our study focused on positive interactive behavior and did not model maternal dopamine responses to an infant's distress, a subject for future research. Furthermore, although mother-infant synchrony associated with maternal dopamine responses across ages, age could account for some neural variability and may be an important aspect to consider in future research.

The evidence reported here encourages future research on the neurochemistry of human bonding, including additional neurotransmitters such as central oxytocin and opioids. Moreover, our results may be useful for clinical research testing the involvement of dopamine and the medial amygdala network in PPD and developmental psychopathology. Dopamine within the medial amygdala network potentially promotes human bonding and thus could play a considerable role in optimal human development.

Materials and Methods

Participants. Nineteen mothers (age range 21–42 y) and their infants (age range 4–24 mo) completed the study. Participants had no psychiatric history and were not breastfeeding or pregnant. The Massachusetts General Hospital Institutional Review Board approved the study, and all mothers signed an informed consent before participating.

Procedure. During a visit to a subject's home, study staff collected video recordings of the mother and the infant. Mothers were then invited to participate in two consecutive PET-fMRI scans during which the mother viewed films of her own infant and an unfamiliar infant, in changing order. While lying in the scanner, mothers passively watched footage of infants playing. The stimuli included a 20-min movie of their own infant during solitary play followed by a 5-min rest and then a 20-min movie of the unfamiliar infant. The radiotracer was injected 10 min into the first film, and PET data collection continued for 90 min. During the second scan, mothers watched the same components of the stimuli with the order of the infants reversed. The initial order of the movies was randomized across participants.

Combined PET-fMRI Scanner. PET data were acquired using the Siemens BrainPET scanner. This prototype device consists of a head-only PET insert (BrainPET) that fits in the bore of the 3-T Total Imaging Matrix Trio MRI scanner (Siemens Healthcare). Each of the 192 BrainPET detector modules consists of a 12×12 array of $2.5 \times 2.5 \times 20$ mm lutetium oxyorthosilicate (LSO) crystals read out by a 3×3 array of magnetic field-insensitive avalanche photodiodes. A PET-compatible circularly polarized (CP) transmittal coil and an eight-channel receive array coil were used to acquire the MR data simultaneously.

MRI and fMRI. Structural data were acquired using a T1-weighted magnetization-prepared rapid acquisition with a gradient echo (MPRAGE) sequence [repetition time (TR) = 2,530 ms, echo time (TE) = 1.63 ms, inversion time (TI) = 1,200 ms, flip angle = 7° , and 1-mm isotropic voxels]. MRI data analysis was performed using FreeSurfer (surfer.nmr.mgh.harvard.edu) and included unpacking, reconstruction, motion correction, intensity normalization, spatial normalization, white matter segmentation, registration, segmentation, and labeling of cortical and subcortical

structures. For intrinsic connectivity analysis, whole-brain fMRI data were acquired with an echo-planar sequence during 6-min resting-state periods (TR = 3,000 ms; TE = 30 ms; 3.0-mm isotropic voxels, 47 slices). To analyze the resting-state fMRI data, a temporal bandpass filter removed frequencies >0.08 Hz. To examine the intrinsic functional connectivity strength of the medial amygdala network, we created spherical volumes around the right medial amygdala seed and the rest of the nodes (25). (For a complete list of the bilateral network coordinates see Table S2.) For each participant, we computed pairwise correlation coefficients between the mean BOLD signal time course of the medial amygdala seed and every target region. The pairwise correlation coefficient values were averaged in each hemisphere to represent a composite measure of connectivity across the network. Then Fisher's r -to- z transformations were calculated and used to assess the correlations between the strength of intrinsic network connectivity and maternal attunement, central dopamine, and plasma oxytocin.

Behavioral Coding of Mother–Infant Synchrony. To measure bonding behavior, 2-min interaction videos were coded for mother–infant synchrony by trained coders (For a detailed description of the coding scheme, see Table S3) (4). Four categories of behavior (vocalization, gaze, affect, and touch) were coded for each dyadic partner, and then the temporal synchronization of those behaviors was computed. We chose to operationalize synchrony using behavioral contingencies of vocalization because vocalization is an outgoing appetitive behavior central to bonding (10, 32), which relates to dopamine (33), and can be measured accurately in both mothers and infants (1).

ACKNOWLEDGMENTS. This research was funded by National Institute of Child Health and Human Development Grant R21HD07164 and National Institute of Biomedical Imaging and Bioengineering Grant R01EB014894.

- Feldman R (2007) Parent–infant synchrony and the construction of shared timing: physiological precursors, developmental outcomes, and risk conditions. *J Child Psychol Psychiatry* 48(3–4):329–354.
- Numan M, Young LJ (2016) Neural mechanisms of mother–infant bonding and pair bonding: Similarities, differences, and broader implications. *Horm Behav* 77: 98–112.
- Swain JE, et al. (2014) Approaching the biology of human parental attachment: Brain imaging, oxytocin and coordinated assessments of mothers and fathers. *Brain Res* 1580:78–101.
- Atzil S, Hendler T, Feldman R (2011) Specifying the neurobiological basis of human attachment: Brain, hormones, and behavior in synchronous and intrusive mothers. *Neuropsychopharmacology* 36(13):2603–2615.
- Gregory R, Cheng H, Rupp HA, Sengelaub DR, Heiman JR (2015) Oxytocin increases VTA activation to infant and sexual stimuli in nulliparous and postpartum women. *Horm Behav* 69:82–88.
- Atzil S, Hendler T, Zagoory-Sharon O, Winetraub Y, Feldman R (2012) Synchrony and specificity in the maternal and the paternal brain: Relations to oxytocin and vasopressin. *J Am Acad Child Adolesc Psychiatry* 51(8):798–811.
- Laurent HK, Ablow JC (2013) A face a mother could love: Depression-related maternal neural responses to infant emotion faces. *Soc Neurosci* 8(3):228–239.
- Moses-Kolko EL, et al. (2011) Rapid habituation of ventral striatal response to reward receipt in postpartum depression. *Biol Psychiatry* 70(4):395–399.
- Chase HW, Moses-Kolko EL, Zevallos C, Wisner KL, Phillips ML (2014) Disrupted posterior cingulate–amygdala connectivity in postpartum depressed women as measured with resting BOLD fMRI. *Soc Cogn Affect Neurosci* 9(8):1069–1075.
- Golinkoff RM, Can DD, Soderstrom M, Hirsh-Pasek K (2015) (Baby)talk to me: The social context of infant-directed speech and its effects on early language acquisition. *Curr Dir Psychol Sci* 24(5):339–344.
- Innis RB, et al. (2007) Consensus nomenclature for in vivo imaging of reversibly binding radioligands. *J Cereb Blood Flow Metab* 27(9):1533–1539.
- Laruelle M (2000) Imaging synaptic neurotransmission with in vivo binding competition techniques: A critical review. *J Cereb Blood Flow Metab* 20(3):423–451.
- Freeman SM, Young LJ (2016) Comparative perspectives on oxytocin and vasopressin receptor research in rodents and primates: Translational implications. *J Neuroendocrinol* 28(4).
- Kendrick KM, Keverne EB, Hinton MR, Goode JA (1991) Cerebrospinal fluid and plasma concentrations of oxytocin and vasopressin during parturition and vaginocervical stimulation in the sheep. *Brain Res Bull* 26(5):803–807.
- Bartz JA, Zaki J, Bolger N, Ochsner KN (2011) Social effects of oxytocin in humans: Context and person matter. *Trends Cogn Sci* 15(7):301–309.
- Sciafani V, Paukner A, Suomi SJ, Ferrari PF (2015) Imitation promotes affiliation in infant macaques at risk for impaired social behaviors. *Dev Sci* 18(4):614–621.
- Simpson EA, et al. (2014) Inhaled oxytocin increases positive social behaviors in newborn macaques. *Proc Natl Acad Sci USA* 111(19):6922–6927.
- Slaughter V, Ong SS (2014) Social behaviors increase more when children with ASD are imitated by their mother vs. an unfamiliar adult. *Autism Res* 7(5):582–589.
- Landa RJ, Holman KC, O'Neill AH, Stuart EA (2011) Intervention targeting development of socially synchronous engagement in toddlers with autism spectrum disorder: A randomized controlled trial. *J Child Psychol Psychiatry* 52(1): 13–21.
- Nadel J, et al. (2000) Do children with autism have expectancies about the social behaviour of unfamiliar people? A pilot study using the still face paradigm. *Autism* 4(2):133–145.
- Contaldo A, Colombi C, Narzisi A, Muratori F (2016) The social effect of “being imitated” in children with autism spectrum disorder. *Front Psychol* 7:726.
- Schultz W (1998) Predictive reward signal of dopamine neurons. *J Neurophysiol* 80(1):1–27.
- Leckman JF, et al. (2004) Primary parental preoccupation: Circuits, genes, and the crucial role of the environment. *J Neural Transm (Vienna)* 111(7):753–771.
- Numan M (2017) *Reference Module in Neuroscience and Biobehavioral Psychology: Parental Behavior* (Elsevier, Amsterdam).
- Bickart KC, Hollenbeck MC, Barrett LF, Dickerson BC (2012) Intrinsic amygdala–cortical functional connectivity predicts social network size in humans. *J Neurosci* 32(42): 14729–14741.
- Lechan RM, Toni R (2013) Functional anatomy of the hypothalamus and pituitary. *Endotext*, eds De Groot LJ, et al. (MDText.com, Inc., South Dartmouth, MA).
- Kühn S, et al. (2010) Why do I like you when you behave like me? Neural mechanisms mediating positive consequences of observing someone being imitated. *Soc Neurosci* 5(4):384–392.
- Atzil S, Hendler T, Feldman R (2014) The brain basis of social synchrony. *Soc Cogn Affect Neurosci* 9(8):1193–1202.
- Delaveau P, et al. (2010) Dopaminergic modulation of the default mode network in Parkinson's disease. *Eur Neuropsychopharmacol* 20(11):784–792.
- Cole DM, et al. (2013) Dopamine-dependent architecture of cortico-subcortical network connectivity. *Cereb Cortex* 23(7):1509–1516.
- Stolzenberg DS, et al. (2007) Dopamine D1 receptor stimulation of the nucleus accumbens or the medial preoptic area promotes the onset of maternal behavior in pregnancy-terminated rats. *Behav Neurosci* 121(5):907–919.
- Murray L, Kempton C, Woolgar M, Hooper R (1993) Depressed mothers' speech to their infants and its relation to infant gender and cognitive development. *J Child Psychol Psychiatry* 34(7):1083–1101.
- Simonyan K, Horwitz B, Jarvis ED (2012) Dopamine regulation of human speech and bird song: A critical review. *Brain Lang* 122(3):142–150.
- Fischl B, et al. (2002) Whole brain segmentation: Automated labeling of neuroanatomical structures in the human brain. *Neuron* 33(3):341–355.
- Farde L, et al. (1985) Substituted benzamides as ligands for visualization of dopamine receptor binding in the human brain by positron emission tomography. *Proc Natl Acad Sci USA* 82(11):3863–3867.
- Byars LG, et al. (2005) Variance reduction on randoms from delayed coincidence histograms for the HRRT. *Proceedings of the IEEE Medical Imaging Conference (IEEE, New York)*, Vol 5, pp 2622–2626.
- Izquierdo-Garcia D, et al. (2014) An SPM8-based approach for attenuation correction combining segmentation and nonrigid template formation: application to simultaneous PET/MR brain imaging. *J Nucl Med* 55(11):1825–1830.
- Watson CC (2000) New, faster, image-based scatter correction for 3D PET. *IEEE Trans Nucl Sci* 47(4):1587–1594.
- Catana C, et al. (2011) MRI-assisted PET motion correction for neurologic studies in an integrated MR-PET scanner. *J Nucl Med* 52(1):154–161.
- Muzic RF, Jr, Cornelius S (2001) COMKAT: Compartment model kinetic analysis tool. *J Nucl Med* 42(4):636–645.
- Logan J, et al. (1996) Distribution volume ratios without blood sampling from graphical analysis of PET data. *J Cereb Blood Flow Metab* 16(5):834–840.

Supporting Information

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SI Materials and Methods

[¹¹C]raclopride, a selective D2 receptor antagonist, was synthesized from the O-desmethyl raclopride precursor and [¹¹C]methyl iodide. The synthesis and subsequent purification by HPLC were performed according to Farde et al. (35) with minor modifications. [¹¹C]raclopride (10.2 ± 1.67 mCi) was injected i.v. as a manual bolus for each scan. Relative dopamine response (depicted in Fig. 2) was indexed as $-\text{[}^{11}\text{C]raclopride BPnd(own infant)} - \text{[}^{11}\text{C]raclopride BPnd(unfamiliar infant)} \times 100 / \text{[}^{11}\text{C]raclopride BPnd(unfamiliar infant)}$. The endogenous dopamine response in the own-infant condition (depicted in Fig. 4) was indexed as $-\text{[}^{11}\text{C]raclopride BPnd(own infant)}$.

Blood for oxytocin analysis was drawn before the MR-PET scans. Samples were spun right after extraction at $1,163 \times g$ for 10 min and stored at -20°C until assayed. Plasma oxytocin was assayed at the Brigham Research Assay Core Laboratory using the Enzo Life Sciences Immunoassay ELISA Kit (catalog no. AD1-900-153A).

Emission data were acquired in list-mode format for 90 min. For each coincidence event, the line of response joining the two crystals in which the two 511 keV photons were detected was rebinned into sinogram space using nearest neighbor approximation and axial compression (span 9 and maximum ring difference 67). Each sinogram consisted of 192 angular projections and 256 radial elements. The calculation of random coincidences was performed by sorting the delayed coincidences into delayed single maps from which the total singles rate as well as the variance reduced random were estimated (36). The sensitivity data were acquired with a plane source scanned in 16 positions (with a 22.5° angular step), 4 h per position, and the normalization sinogram was derived from these data. The head attenuation map was generated from the MR data (37). The scatter coincidences sinogram was obtained using a calculated method based on the

single scatter estimation method (38). To correct for head motion over the duration of the scan, head motion estimates were derived offline from the MR data for all the echo planar imaging-based acquisitions and were used to correct the dynamic PET frames before image reconstruction (39). For this purpose, the list-mode dataset was first divided into frames of variable duration according to the pharmacokinetic protocol (i.e., 8×10 s, 3×20 s, 2×30 s, 1×60 s, 1×120 s, 1×180 s, 8×300 s, and 4×600 s). Motion correction was subsequently applied separately to each of these frames. The head attenuation and scatter correction sinograms were estimated only in the reference frame. The motion-corrected PET volumes corresponding to each of the frames were reconstructed using the standard Ordinary Poisson Ordered Subset Expectation Maximization (OP-OSEM) 3D algorithm from motion-corrected prompt and random coincidences, normalization, attenuation, and scatter coincidences sinograms using 16 subsets and six iterations. The reconstructed volume consisted of 153 slices with 256×256 pixels ($1.25 \times 1.25 \times 1.25$ mm). The BPnd (11) of [¹¹C]raclopride was the primary outcome measure of the PET scan. “BPnd” refers to the radioligand molecules that are specifically attached to a neuroreceptor (as opposed to free radioligand in the tissue) and is the typical measure when using a reference tissue method (11). Regional analyses were performed using COMKAT (40) and a reference tissue model with the time-activity curve derived from the cerebellar cortex as a reference (41). The cerebellum was chosen as reference region because it does not contain specific D2 receptor-like binding sites and can be used for the determination of nonspecific binding and free radioligand in the brain. BPnd values for all subjects were imported to SPSS for group analyses and correlated (in Pearson correlation analyses) to indices of behavioral synchrony, resting-state BOLD in the medial amygdala network, and plasma oxytocin levels.

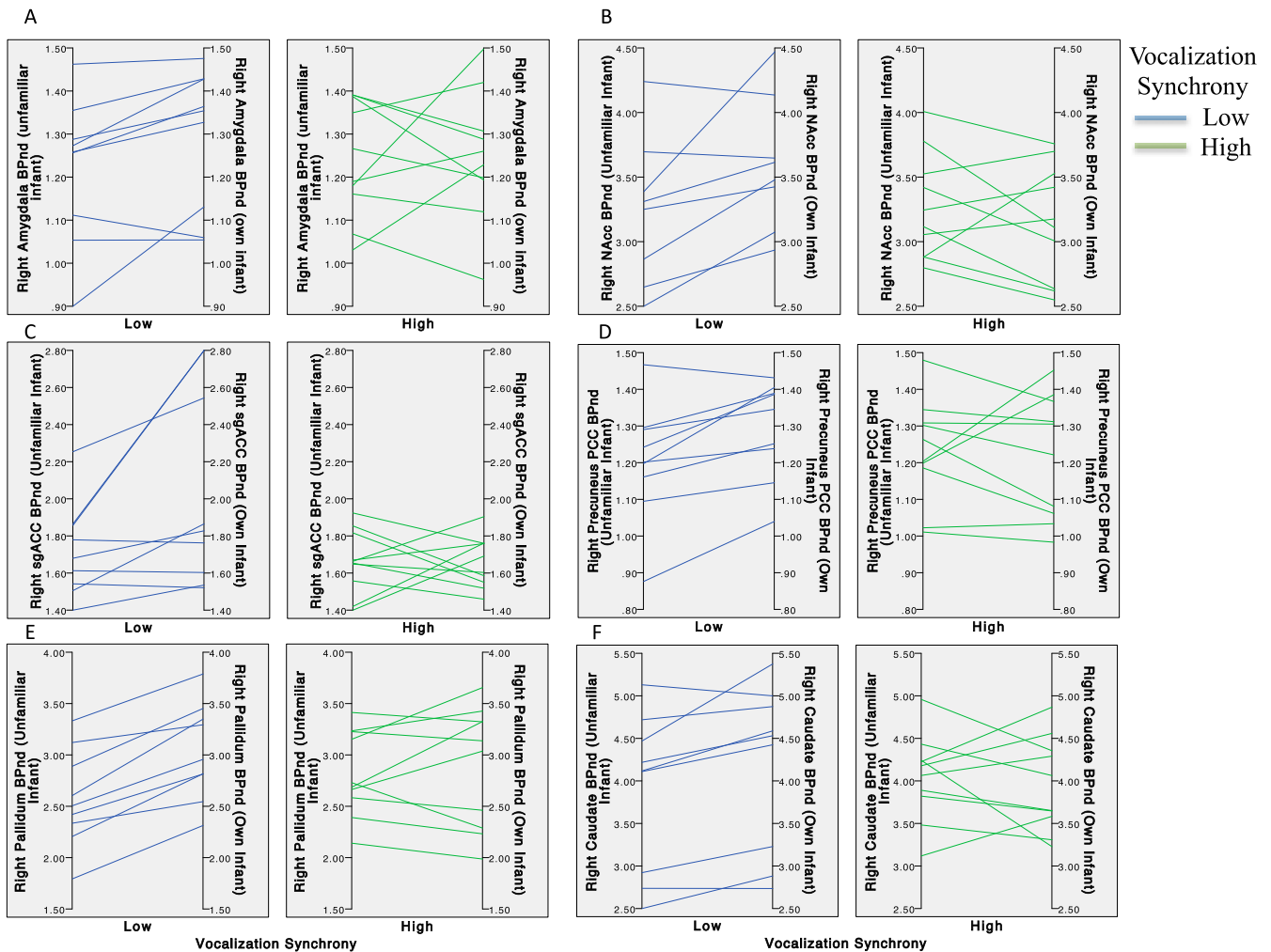


Fig. S1. Individual [^{11}C]raclopride BPnd data showing that more high-synchrony mothers than low-synchrony mothers have a dopamine preference to their own infants. Lower [^{11}C]raclopride BPnd represents increased endogenous dopamine. High-synchrony mothers are depicted in green. Low-synchrony mothers are depicted in blue. (A) Right amygdala: Six of 10 mothers in the high-synchrony group have lower raclopride BPnd in the own-infant condition (indexing increased endogenous dopamine) compared with one of nine mothers in the low-synchrony group. (B) Right NAcc: Six of 10 mothers in the high-synchrony group have lower raclopride BPnd in the own-infant condition (indexing increased endogenous dopamine) compared with two of nine mothers in the low-synchrony group. (C) Right sgACC: Six of 10 mothers in the high-synchrony group have lower raclopride BPnd in the own-infant condition (indexing increased endogenous dopamine) compared with three of nine mothers in the low-synchrony group. (D) Right PCC: Six of 10 mothers in the high-synchrony group have lower raclopride BPnd in the own-infant condition (indexing increased endogenous dopamine) compared with one of nine mothers in the low-synchrony group. (E) Right pallidum: Six of 10 mothers in the high-synchrony group have lower raclopride BPnd in the own-infant condition (indexing increased endogenous dopamine) compared with zero of nine mothers in the low-synchrony group. (F) Right caudate: Six of 10 mothers in the high-synchrony group have lower raclopride BPnd in the own-infant condition (indexing increased endogenous dopamine) compared with two of nine mothers in the low-synchrony group.

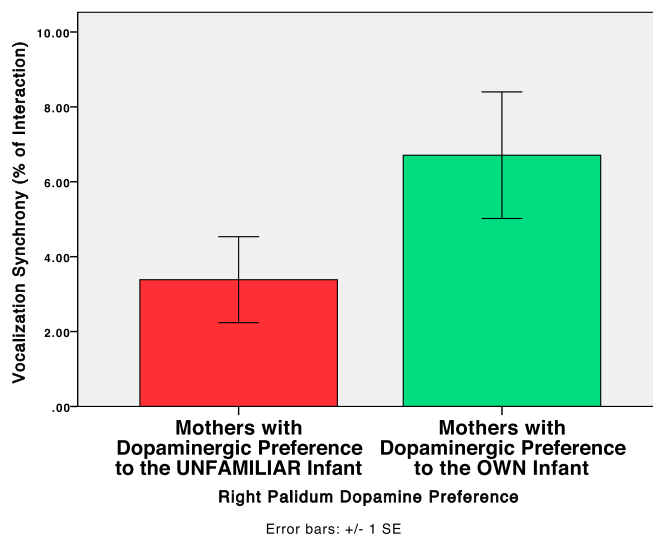


Fig. S2. Differences in synchrony by endogenous dopamine preference. Mothers whose right pallidum response showed an endogenous dopaminergic preference for their own infant (green bar, $n = 6$) had significantly higher vocalization synchrony scores than mothers whose right pallidum dopaminergic response was higher for the unfamiliar infant (red bar, $n = 13$) ($F = 4.3$, $P < 0.05$).

Table S1. Differences in the percentage of dopamine responses comparing mothers with high ($n = 10$) or low ($n = 9$) vocalization synchrony

Region of interest	Low vocalization synchrony (mean of percent change in dopamine signal)	High vocalization synchrony (mean of percent change in dopamine signal)	F	P
Right pallidum	-18.7	-2.1	11.12	0.004
Right NAcc	-15.8	3.2	9.3	0.007
Right caudate	-14.5	0.0	4.15	0.058
Right sgACC	-8.2	1.5	4.2	0.055
Right precuneus PCC	-8.0	0.6	4.1	0.058
Right vmPFC	-7.3	0.1	3.8	0.066
Right ventral PCC	-4.3	1.9	2.2	0.159
Right ACC	-3.3	0.9	1.2	0.294
Right amygdala	-6.5	-1.1	1.06	0.3
Right putamen	-3.2	2.0	1.02	0.327
Left putamen	-4.1	2.6	3.06	0.098
Left precuneus PCC	-3.4	2.2	2.65	0.122
Left vmPFC	-6.3	-1.4	1.732	0.206
Left ACC	-4.3	0.1	1.232	0.283
Left Pallidum	3.4	0.2	0.325	0.576
Left ventral PCC	-1.7	0.3	0.251	0.623
Left caudate	-0.3	2.3	0.239	0.631
Left sgACC	-1.3	-3.7	0.215	0.649
Left amygdala	0.1	1.6	0.112	0.742
Left NAcc	-0.1	-0.6	0.008	0.929

Mothers with higher vocalization synchrony have increased percentage of endogenous dopamine signal change in the contrast between her own and an unfamiliar infant, indexed by a decrease in the change in [^{11}C]raclopride BPnd. Group differences are calculated using a multivariate general linear model. Regions are presented in each hemisphere according to their effect sizes.

Table S2. Nodes of the medial amygdala network

Medial amygdala network nodes	x	y	z	Sphere radius, mm
Left anterior hippocampus	-22	-14	-20	2
Right anterior hippocampus	24	-14	-20	2
Left nucleus accumbens	-8	8	-8	2
Right nucleus accumbens	8	8	-8	2
Left vmPFC	-2	44	-10	4
Right vmPFC	2	40	-8	4
Left dorsomedial PFC	-2	60	4	4
Right dorsomedial PFC	2	60	4	4
Left ventromedial hypothalamus	-4	4	-8	2
Right ventromedial hypothalamus	6	4	-12	2
Left temporal pole	-34	20	-34	4
Right temporal pole	40	22	-38	4
Left thalamus	-2	-6	4	2
Right thalamus	2	-6	4	2
Left substantia innominata	-14	0	-12	2
Right substantia innominata	14	0	-12	2
Left posterior cingulate cortex	-4	-52	22	4
Right posterior cingulate cortex	4	-50	20	4
Left subgenual anterior cingulate cortex	-2	30	-4	4
Right subgenual anterior cingulate cortex	2	26	-6	4
Left anterior medial temporal gyrus	-60	-6	-20	4
Right anterior medial temporal gyrus	62	-6	-22	4

We used a hypothesis-driven, seed-based resting-state fMRI analysis. Coordinates were previously identified in independent samples (25). We created spherical regions of interest around the right medial amygdala seed and the rest of the nodes. For each participant, we computed pairwise correlation coefficient values between the mean BOLD signal time course of the medial amygdala seed and every target region. The pairwise correlation coefficient values then were averaged in each hemisphere to represent a composite measure of connectivity across the medial amygdala network.

Table S3. Coding bonding behavior

Subject	Behavior	Code
Mothers	Vocalization	Motherese (high-pitched speech and sing-song vocalization)
		Adult-voice vocalization to the infant
		Talk to the experimenter
	Gaze	No vocalization
		To infant
		To object
		To environment
	Affect	Gaze aversion
		Negative
		Neutral
	Touch	Positive
Affectionate touch		
Functional touch		
Infants	Vocalization	No touch
		Cry
		Fuss
		Positive vocalizations (e.g., cooing, giggling, or laughing)
		No vocalizations
	Gaze	To parent
		To object
		To environment
	Affect	Gaze aversion
		Negative
		Neutral
		Positive

Trained coders coded second-by-second behaviors for 2-min films of mother–infant interactions. The trained coder chose the correct predefined code for each behavior for every second of the interaction. Each behavior is coded separately for each dyadic partner. Thereafter, the software (Noldus) allows the computation of temporal nesting of events. The vocalization synchrony variable summarizes the percentage of time during the 2-min film in which mother and infant simultaneously performed positive vocalizations, i.e., the mother was performing motherese while the infant was cooing, giggling, or laughing. The vocalization attunement variable summarizes the percentage of time during the 2-min film in which mothers performed motherese while the infant was showing positive affect (content and socially engaged).