IL-17a promotes sociability in mouse models of neurodevelopmental disorders

A subset of children with autism spectrum disorder appear to show an improvement in their behavioural symptoms during the course of a fever, a sign of systemic inflammation. Here we elucidate the molecular and neural mechanisms that underlie the beneficial effects of inflammation on social behaviour deficits in mice. We compared an environmental model of neurodevelopmental disorders in which mice were exposed to maternal immune activation (MIA) during embryogenesis with mouse models that are genetically deficient for contactin-associated protein-like 2 (Cntnap2), fragile X mental retardation-1 (Fmr1) or Sh3 and multiple ankyrin repeat domains 3 (Shank3). We establish that the social behaviour deficits in offspring exposed to MIA can be temporarily rescued by the inflammatory response elicited by the administration of lipopolysaccharide (LPS). This behavioural rescue was accompanied by a reduction in neuronal activity in the primary somatosensory cortex dysgranular zone (S1DZ), the hyperactivity of which was previously implicated in the manifestation of behavioural phenotypes associated with offspring exposed to MIA. By contrast, we did not observe an LPS-induced rescue of social deficits in the monogenic models. We demonstrate that the differences in responsiveness to the LPS treatment between the MIA and the monogenic models emerge from differences in the levels of cytokine production. LPS treatment in monogenic mutant mice did not induce amounts of interleukin-17a (IL-17a) comparable to those induced in MIA offspring; bypassing this difference by directly delivering IL-17a into S1DZ was sufficient to promote sociability in monogenic mutant mice as well as in MIA offspring. Conversely, abrogating the expression of IL-17 receptor subunit a (IL-17Ra) in the neurons of the S1DZ eliminated the ability of LPS to reverse the sociability phenotypes in MIA offspring. Our data support a neuroimmune mechanism that underlies neurodevelopmental disorders in which the production of IL-17a during inflammation can ameliorate the expression of social behaviour deficits by directly affecting neuronal activity in the central nervous system.

The beneficial effects of infection and the ensuing inflammation on neurological disorders have previously been noted. For example, a subset of children with autism spectrum disorder (ASD) exhibit temporary but considerable improvements of their behavioural symptoms during episodes of fever, a sign of systemic inflammation. However, a mechanistic understanding of how fever-associated immune responses translate into behavioural relief—both at the molecular and neural level—is lacking. The aetiology of ASD includes both environmental and genetic risk factors. Mouse models for ASD, in which mice contain mutations in ASD risk genes (including Cntnap2, Fmr1, and Shank3), show behavioural abnormalities—including deficits in social interaction. Similar behavioural abnormalities are observed in mouse models for environmental risk factors for developing ASD, such as exposure to maternal inflammation. We sought to explore the mechanisms that enable the fever-associated rescue of sociability deficits, using both genetic and environmental mouse models for neurodevelopmental disorders.

LPS rescues sociability in MIA offspring

A febrile response can be exogenously induced by injecting mice with a low dose of LPS. Indeed, the intraperitoneal administration of LPS rescues sociability in MIA offspring.
Body temperature profile after injection of saline (vehicle) or LPS in offspring of mothers injected with PBS (PBS offspring) (vehicle, n = 11 and LPS, n = 11 mice, from 5 independent experiments). The initial spike in body temperature is due to handling stress.

Mice were tested for sociability (percentage of time spent investigating social object/total time spent investigating both social and inanimate objects) one day before LPS injection (pre-test; pre). Mice were then tested for sociability four hours after injection with vehicle or LPS (test). PBS offspring + vehicle, n = 10; PBS offspring + LPS, n = 9; MIA offspring + vehicle, n = 10; MIA offspring + LPS, n = 12; wild type (WT) + vehicle, n = 8; wild type + LPS, n = 11; Cntnap2 mutant + vehicle, n = 11; Cntnap2 mutant + LPS, n = 11; Fmr1 mutant + vehicle, n = 11; Fmr1 mutant + LPS, n = 15; Shank3 mutant + vehicle, n = 8; Shank3 mutant + LPS, n = 10; from 3 independent experiments. I.p., intraperitoneal.

**Fig. 2** Immune stimulation reduces hyperactivation in the S1DZ of MIA offspring. **a**, Representative images illustrating FOS (green) expression in the S1DZ and CeA, after injection with vehicle or LPS. Scale bars, 200 μm. Numerals indicate cortical layers. **b**, Quantification of FOS-expressing cells in the S1DZ (b) and CeA (c). For experiments in **a**–**c**, PBS offspring + vehicle, n = 8; PBS offspring + LPS, n = 9; MIA offspring + vehicle, n = 13; MIA offspring + LPS, n = 11; from 3 independent experiments. **d**, AAV encoding either EYFP or EYFP fused to enhanced halorhodopsin (NpHR) was bilaterally injected into the S1DZ of monogenic mutant mice. Scale bar, 500 μm. **e**, Performance on sociability was assessed in the presence (on) and absence (off) of optical inhibition. For experiments in **d**–**f**, wild type + EYFP, n = 7; wild type + NpHR, n = 8; Cntnap2 mutant + EYFP, n = 11; Cntnap2 mutant + NpHR, n = 9; Fmr1 mutant + EYFP, n = 8; Fmr1 mutant + NpHR, n = 12; Shank3 mutant + EYFP, n = 8; Shank3 mutant + NpHR, n = 10; from 6 independent experiments. **g–i**, Quantification of FOS-expressing cells after injection of vehicle or LPS in the S1DZ (h) and CeA (i). For experiments in **g–i**, Cntnap2 mutant + vehicle, n = 9; Cntnap2 mutant + LPS, n = 10; Fmr1 mutant + vehicle, n = 7; Fmr1 mutant + LPS, n = 9; Shank3 mutant + vehicle, n = 6; Shank3 mutant + LPS, n = 8; from 3 independent experiments. All n values refer to the number of mice used. Statistics calculated by two-way ANOVA with Sidak’s (c) or Tukey’s (e) post hoc tests, or one-way repeated-measures ANOVA with Tukey’s post hoc test (f). Graphs are mean ± s.e.m.
Acute fever does not restore sociability

We next probed whether fever has a role in the observed behavioural rescue. Unlike in control offspring, LPS treatment did not induce changes in body temperature in MIA mice (Extended Data Fig. 4a), which suggests that the febrile response might not be the main factor that contributes to the rescue. To directly test whether fever is dispensable, we sought to increase the body temperature of the mice without inducing systemic inflammation by targeting inhibitory designer receptors exclusively activated by designer drugs (DREADDs)\(^1\) to GABAergic neurons in the ventral part of the lateral preoptic nucleus of \(\mathrm{vgat}^{\text{Cre}}\) (Methods) mice (Fig. 1d). As previously reported\(^\text{2}\),\(^\text{3}\),\(^\text{4}\), inhibition of these neurons led to an increase in body temperature of about 1 \(^\circ\)C (Fig. 1e). However, the induction of febrile response alone did not promote social preference in MIA offspring (Fig. If, g, Extended Data Fig. 4b–e), confirming that fever per se is not the main driver of the LPS-induced rescue. Of note, \(\mathrm{vgat}^{\text{Cre}}\) MIA offspring exhibited sociability deficits comparable to those of wild-type MIA offspring, were unaffected by treatment with clozapine \(N\)-oxide (CNO) and showed an increase in sociability after treatment with LPS (Extended Data Fig. 4f–j).

LPS reduces FOS induction in the S1DZ

We previously established that adult MIA offspring display cortical abnormalities that are preferentially localized in the S1DZ\(^7\), a subregion of the primary somatosensory cortex (SI) that is cyto-architecturally defined by the absence of a discernible fourth layer (Extended Data Fig. 5). The cortical phenotype is characterized by an overall increase in neural activity that, when reduced, can acutely rescue MIA-induced deficits in social behaviours\(^5\). We therefore investigated whether the LPS-induced behavioural rescue in MIA offspring is accompanied by changes in neural activity in the S1DZ. MIA offspring exhibited an increase in the number of Cells in the S1DZ that express FOS (a marker for neuronal activation), relative to control offspring. However, in LPS-treated MIA offspring, the number of FOS\(^+\) neurons in the S1DZ was reduced to the level of control offspring (Fig. 2a, b, Extended Data Fig. 6a–c). LPS injections did not elicit a generalized, brain-wide effect.
**IL-17a rescues sociability deficits**

The injection of LPS is known to increase the production of inflammatory cytokines\(^1\). We observed that administration of LPS results in a robust increase in the plasma levels of IFNγ, IL-6 and TNF-α (also known as TNF-α) (Fig. 3a). IL-17a—the orthologue of which in *Caenorhabditis elegans* has previously been implicated in modulating sensory responses\(^2\)—was prominently upregulated in MIA offspring, but not in monogenic mutant mice or in control mice (Fig. 3a). Furthermore, we noted that the receptor subunit A for IL-17a (IL-17Ra) is expressed in cortical neurons, including in the S1DZ (Fig. 3b–d, Extended Data Fig. 8a–c). These data suggested that the increased levels of IL-17a after treatment with LPS in MIA offspring, but not in monogenic mutant mice, may directly affect the S1DZ and thus restore sociability. Consistent with this idea, the direct administration of recombinant IL-17a into the S1DZ was sufficient to increase sociability not only in MIA offspring but also in Cntnap2 and Fmr1 mutant mice (Fig. 3e, Extended Data Fig. 8d–h).

**LPS-induced rescue requires IL-17a**

To further determine whether IL-17a mediates the LPS-driven behavioural rescue, we inhibited IL-17a activity in the brain via intracerebroventricular injection of blocking antibodies. Antibodies against IL-17a prevented both the LPS-induced rescue of sociability (Fig. 4a, Extended Data Fig. 9a–e) and the reduction of FOS expression in the S1DZ of MIA offspring (Extended Data Fig. 9f). To directly assay the effects of LPS on neural activity, we used multi-electrode arrays to measure the firing rate of neurons in the S1DZ in awake mice as defined by Tukey. PBS offspring + vehicle + isotype, \(n = 65\) cells; PBS offspring + LPS + isotype, \(n = 42\) cells; PBS offspring + LPS + anti-IL-17a, \(n = 40\) cells; MIA offspring + vehicle + isotype, \(n = 75\) cells; MIA offspring + LPS + isotype, \(n = 48\) cells; MIA offspring + LPS + anti-IL-17a, \(n = 43\) cells; two PBS offspring and 2 MIA offspring in 12 independent experiments. e, Lentivirus encoding either EYFP or enhanced green fluorescent protein (EGFP) fused to nuclear Cre (nCre) was bilaterally injected into the S1DZ of IL-17Ra\(^{-/-}\) MIA offspring. Scale bar, 200 μm. f, g, Mice were tested for sociability one day before injection (pre). The following day, mice were tested for sociability four hours after LPS injection (test). For experiments in e, f, IL-17Ra\(^{-/-}\)EYFP, \(n = 9\) and IL-17Ra\(^{-/-}\)EGFP:nCre, \(n = 10\); from 5 independent experiments. Unless otherwise indicated, \(n\) values refer to the number of mice used. Statistics calculated by two-way repeated-measures ANOVA with Bonferroni’s post-hoc test (a, f) or two-way ANOVA with Tukey’s post-hoc test (d). Graphs are mean ± s.e.m.
Upon treatment with LPS, we observed a decrease in the overall firing rate that was prevented by blocking IL-17a in MIA offspring. Treatment with LPS or IL-17a blocking antibody did not change the neuronal firing rate in the S1DZ of control offspring (Fig. 4b–d, Extended Data Fig. 10c). Furthermore, LPS injection did not restore sociability to MIA offspring that are deficient for IL-17Ra in the S1DZ (Fig. 4e, f, Extended Data Fig. 10d–j). Our data collectively demonstrate that IL-17a mediates the restoring effects on social behaviours by directly acting on IL-17Ra neurons of the S1DZ.

Discussion

Previous data in mice have suggested that increased production of IL-17a in pregnant mothers may present as a risk factor for neurodevelopmental disorders in offspring.15,16,17,18. On the basis of our current findings, we propose that in adult MIA offspring the same cytokine is beneficial and ameliorates sociability phenotypes during episodes of inflammation. Treatment with LPS led to an increase in IL-17a levels in the blood selectively in MIA offspring—but not in other monogenic mutant mice. This suggests that the inflammatory responses may result in beneficial effects only for individuals who have their immune systems primed by prenatal exposure to immune activation, or by other environmental factors. A better understanding of the role of a primed immune system among patients with neurodevelopmental disorders may help to identify those patients whose behavioural symptoms are likely to improve after exposure to fever-associated inflammation. Furthermore, elucidating the mechanisms by which IL-17a can induce two opposing behavioural outcomes depending on when its upregulation occurs (during embryonic brain development or in the adult brain) may provide opportunities to devise therapeutic as well as preventive treatments for the behavioural symptoms associated with ASD.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41586-019-1843-6.

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Mice

All experiments were performed according to the Guide for the Care and Use of Laboratory Animals and were approved by the National Institutes of Health and the Committee on Animal Care at Massachusetts Institute of Technology. C57BL/6 mice were purchased from Taconic institutes of Health and the Committee on Animal Care at Massachusetts and Use of Laboratory Animals and were approved by the National Institute. All experiments were performed according to the Guide for the Care and Use of Laboratory Animals.

Body temperature was sampled in 5-min increments. LPS or vehicle (PBS or MIA) were administered subcutaneously. Mice were given pre-operative slow-release buprenorphine (1.0 mg/kg, intraperitoneal injection) and xylazine (10 mg/kg, intraperitoneal injection). Mice were anaesthetized using a mixture of ketamine (100 mg/kg, intraperitoneal injection) and xylazine (10 mg/kg, intraperitoneal injection). Mice were given pre-operative slow-release buprenorphine (1.0 mg/kg, subcutaneous injection). For manipulating body temperature with LPS or vehicle (PBS or MIA), mice were anaesthetized using a mixture of ketamine (100 mg/kg, intraperitoneal injection) and xylazine (10 mg/kg, intraperitoneal injection). Mice were given pre-operative slow-release buprenorphine (1.0 mg/kg, subcutaneous injection).

Maternal immune activation

Mice were mated overnight with females carrying SFB in their guts. On E12.5, pregnant female mice were weighed and injected with a single dose (20 mg/kg, intraperitoneal injection) of poly(lC) (P9582, Sigma Aldrich) or PBS. Each dam was returned to its cage and left undisturbed until the birth of its litter. All pups remained with the mother until weaning on postnatal day (P)21–28, at which time mice were group-housed at a maximum of 5 per cage with same-sex littermates. Matings between Balb/c (c/c) males and wild-type females were used to make MIA Igμ–Cre mice.

Stereotaxic surgery

Surgery was carried out using aseptic techniques. Mice were anaesthetized using a mixture of ketamine (100 mg/kg, intraperitoneal injection) and xylazine (10 mg/kg, intraperitoneal injection). Mice were given pre-operative slow-release buprenorphine (1.0 mg/kg, subcutaneous injection). For manipulating body temperature with LPS, mice were anaesthetized using a mixture of ketamine (100 mg/kg, intraperitoneal injection) and xylazine (10 mg/kg, intraperitoneal injection). Mice were given pre-operative slow-release buprenorphine (1.0 mg/kg, subcutaneous injection).

Immunohistochemistry

Mice were transcardially perfused with cold paraformaldehyde (PFA) (4% in PBS). Brains were kept in PFA overnight at 4°C before vibratome-sectioning (Leica VT1000S). Brains were cut at 50-μm thickness for FOS quantification. Brains were cut at 100-μm thickness for all other experiments.

Before antibody labelling, sections were incubated in blocking solution (0.4% Triton X-100 and 2% goat serum in PBS) for 30 min. Sections were then incubated in blocking solution containing primary antibodies overnight at room temperature. Primary antibodies used were chicken anti-GFP (1:1,000, Ab5450, Abcam), rabbit anti-FOS (1:500, ABE457, Millipore), rabbit anti-DsRed (1:1,000, 632496, Clontech) and mouse anti-NeuN (1:1,000, MAB377, Millipore). Sections were washed in wash buffer (0.4% Triton X-100 in PBS) three times before secondary antibody labelling. Sections were incubated in blocking solution containing secondary antibodies and DAPI (1:5,000, D1306, Thermo Fisher) for three hours at room temperature. Images of stained slices were acquired using a confocal microscope (LSM710, Carl Zeiss) with a 10×, 20× or 40× objective lens.

In situ hybridization

Mice were transcardially perfused with cold PBS. Brains were extracted and embedded in optimal cutting temperature (OCT) compound on dry ice. Sections were cut at 20-μm thickness on a cryostat. In situ hybridizations were performed using the RNAscope 2.5 HD Assay-Red kit (322350, Advanced Cell Diagnostics) using a probe targeting the Il17ra transcript (Mm-Il17ra-O1, 566131, Advanced Cell Diagnostics). The probe was designed to target region 444–882 of the Il17ra transcript (NM_008359.2). Modifications to the kit protocol to improve adherence of tissue to the slide included an extension of the fixation time to 30 min and the addition of a humidified bake step at 40°C immediately before probe hybridization. Sections were counterstained with DAPI. Images were acquired using a confocal microscope (LSM710, Carl Zeiss) with a 10× or 20× objective lens. Il17ra and DAPI expression was quantified using QuPath. Cells were divided into the following categories based on level of Il17ra expression: low = 1–3 puncta, medium = 4–9 puncta, high = 10–15 puncta, highest = >15 puncta.

In situ hybridization followed by immunohistochemistry

For experiments assaying the overlap of Il17ra and NeuN expression, immunohistochemistry for NeuN was performed following a modified in situ hybridization protocol. The RNAscope 2.5 HD Assay-Red kit in situ probe was modified in the following ways: sections were baked at 60°C, followed by a 10-min fixation step with 4% PFA at room temperature. Sections were then stored in 70% ethanol at 4°C overnight. Sections were permeabilized in 8% SDS for 10 min. Sections were washed twice with PBS between each step. After SDS treatment, the RNAscope 2.5 HD Assay-Red kit protocol was followed from the probe hybridization step. Following the completion of the in situ protocol, sections were incubated in blocking buffer containing anti-NeuN antibody overnight at 4°C. Sections were then incubated in blocking buffer containing DAPI and secondary antibody for two hours at room temperature.

Behavioural analysis

Male mice were tested during the light cycle in a room with lighting maintained at 230 lux. Mice were transferred to the testing area at least one hour before the initiation of experiments. Tracking of mouse behaviour was done using the EthoVision XT (Noldus) tracking system.

Three-chamber social approach assay

Adult male mice were assayed for sociability using a three-chamber social approach assay. The arena
was constructed of white acrylic (50 cm × 35 cm × 30 cm). Wire cups (Spectrum Diversified) were placed in the back left and right corner of the arena beneath water-filled 1-l bottles (Nalgene). On day 0, mice were habituated to the arena for 10 min. Immediately after habituation, mice were singly housed. On day 1 (pre-test), mice were placed in the centre of the arena and allowed to freely explore. Following 10 min, mice were confined to the centre of the arena. An inanimate object (rubber stopper) or a male conspecific were placed beneath the wire cups. Placement of the inanimate object and social target were alternated. Mice were then allowed to freely explore the arena for 10 min. Interaction time was defined as time spent in the areas circumscribing the wire cups (<2 cm). Sociability was defined as interaction time with the social target divided by total interaction time and expressed as a percentage. For experiments involving LPS injections, mice were injected with either saline (vehicle) or LPS (50 μg/kg, intraperitoneal injection, L2630, Sigma) on day 2 (test), four hours before testing. For the experiment assaying sociability 72 h after injection of LPS, mice used for 4-h LPS sociability experiments were tested for sociability again at 72 h.

Three-chamber social approach assay with DREADD manipulation. Adult male Vgat–Cre MIA offspring were bilaterally injected with virus encoding the inhibitory DREADD receptor fused to mCherry into the vLPO. After more than three weeks of recovery, mice were assayed on the three-chamber social approach assay outlined above. Baseline sociability was assayed on day 1 (pre-test). On day 2 and day 3, mice were injected with either vehicle or CNO (1.5 mg/kg, intraperitoneal injection, BML-NS105, Enzo Life Sciences), two hours before initiation of behaviour. Injection order was counterbalanced. Following behavioural experiments, post-mortem histology was used to confirm mCherry expression within the vLPO. For experiments assaying the effect of injection of CNO and LPS in Vgat–Cre PBS and MIA offspring that have not undergone surgery, baseline sociability was assessed on day 1. On day 2 and 3, mice received counterbalanced injections of CNO or vehicle. On day 4, mice were injected with LPS.

Three-chamber social approach assay with administration of IL-17a into the S1DZ. Adult male mice were implanted with a cannula into S1DZ bilaterally and allowed to recover for more than two weeks before the behavioural experiments. On day 1 (pre-test), mice were assayed for sociability. On day 2 (test), mice were anaesthetized briefly using isoflurane and either vehicle or IL-17a (50 ng per side in 1 μl at a rate of 180 nl/min, 7956-ML/CF; R&D) was administered bilaterally into the S1DZ through 250-μm projecting injector tips (PlasticsOne). Four hours after administration of vehicle or IL-17a, mice were assayed for sociability. Cannula placements were verified using histology.

Three-chamber social approach assay with S1DZ optogenetic inhibition. EYFP or NpHR were virally targeted to the S1DZ. After two weeks of recovery, mice were allowed to freely explore the arena for 10 min. The following day, the mice were given 3 min of no stimulation (‘off’ session) and 3 min of laser stimulation (‘on’ session) (594 nm, 6 mW).

Three-chamber social approach assay with IL-17a blocking antibody. For central cytokine blockade experiments, adult male mice were implanted with a cannula into the lateral ventricle and allowed to recover for more than two weeks before the behavioural experiments. On day 1 (pre-test), mice were tested for baseline sociability. On day 2 (test), mice were injected with IL-17a blocking antibody (clone 50104; R&D) or isotype-control antibody (IgG2a, clone 54447; R&D). Antibodies were dissolved in saline and administered intracerebroventricularly at 1 mg/kg in 500 nl at a rate of 180 nl/min through 750-μm projecting injector tips (PlasticsOne). Blocking antibody was administered 30 min before administration of LPS. Four hours following administration of LPS, mice were assayed for sociability.

Three-chamber social approach assay in mice deficient for IL-17Ra in the S1DZ. IL-17Ra-knockout was mediated by viral delivery of Cre recombinase, expressed under the control of the human synapsin (hSyn) promoter, into the S1DZ of Il-17ra<sup>−/−</sup> MIA offspring. After at least three weeks of recovery, mice were assayed for sociability. On the next day sociability was assessed four hours following administration of LPS.

Marble burying assay. On day 1, mice were tested for their baseline marble burying phenotype. On day 2, four hours before beginning the marble burying assay, mice were treated with either LPS or vehicle. The marble burying assay was carried out as previously described. Mice were placed into testing arenas (arena size 40 cm × 20 cm × 30 cm; bedding depth 3 cm) each containing 20 glass marbles (laid out in 4 rows of 5 marbles equidistant from one another). At the end of the 15-min exploration period mice were carefully removed from the testing cages and the number of marbles buried was recorded. The marble burying index was arbitrarily defined as the following: 1 for marbles covered >50% with bedding, 0.5 for marbles covered <50% with bedding or 0 for anything less.

Reciprocal social interaction assay. Four hours before testing, mice were injected with vehicle or LPS. Two unfamiliar mice of the same treatment and background were placed in a fresh mouse cage and allowed to freely interact for 10 min. Videos were acquired using IC Capture (The Imaging Source) at a 640 × 480 aspect ratio and 25 frames per second. Social interaction (close following, push–crawl, nose–nose sniffing and nose–anus sniffing) was scored by an observer blind to treatment and background.

Quantification of FOS<sup>+</sup> cells in the brain after administration of LPS

Adult male mice were killed five hours after injection of LPS. FOS<sup>+</sup> cells were quantified using the Cell Counter plugin in Fiji<sup>32</sup>. All cells were counted within a single coronal section of each respective brain region, as defined by the Paxinos and Franklin Mouse Atlas<sup>33</sup>. Regions quantified include: medial prefrontal cortex (mPFC) (comprising prelimbic cortex (PrL) and infralimbic cortex (IL)), AP +1.98; S1DZ, AP −0.46; primary somatosensory cortex barrel field (S1BF), AP −0.46; primary motor cortex (MI), AP −0.46; secondary motor cortex (M2), AP −0.46; secondary auditory cortex dorsal part (AuD), AP −1.94; CeA, AP −1.94; and primary visual cortex (V1), AP −3.64. For experiments testing the IL-17a dependence of LPS-induced changes in FOS expression, mice were injected intracerebroventricularly with control antibodies or blocking antibody against IL-17a 30 min before intraperitoneal injection of vehicle or LPS. Surgical and injection methods were identical to behavioural experiments.

Enzyme-linked immunosorbent assay

After four hours of administration of vehicle or LPS, mice were anaesthetized by intraperitoneal injection of Fatal-Plus (100 mg/kg). All blood samples were centrifuged at 10,000 g for 10 min at 4 °C. All samples were stored at −80 °C until further analysis. Cytokine concentrations in plasma were measured using an ELISA kit (IFNy; 430804, TNF; 430904, IL-6; 431304 and IL-17a; 432504, Biolegend), following the manufacturer’s instructions.

PCR for assaying IL-17Ra knockout

Il17ra<sup>−/−</sup> male mice were bilaterally injected with virus encoding nuclear Cre fused to EGFP, or control virus encoding only EYFP, into the S1DZ. After more than three weeks, injection sites were dissected from the SI. Single cells were dissociated from brain tissue using a modified version<sup>34</sup> of the Papain Dissociation Kit protocol (LK003153, Worthington) and sorted on a BD FACS Aria (BD Biosciences) based on EGFP and EYFP expression. RNA was extracted from sorted cells using
a Quick-RNA micro-prep kit (Zymo). Twenty nanograms of RNA was converted into cDNA using oligo(dT) (Promega First Strand CDNA Synthesis Kit, NEB). Il17ra and Gapdh mRNA expression was assessed using PCR. One microlitre of cDNA was diluted in a 20-μl reaction volume. Il17ra and Gapdh and mRNA expression was assessed using the following primers: Il17ra 5′-AGATGCCACATCCTGTACC-3′ and 5′-CACAGTACAGCGTGTCGTA-3′; Gapdh 5′-GACCTCAACGCCCTCCACTCTCCG-3′ and 5′-TGGGTCCTCAGTTCTTATCTCCT-3′. Cycling conditions for Il17ra: 95°C × 5 min (1 cycle), 95°C × 20 s, 60°C × 30 s, 72°C × 30 s (32 cycles), 72°C × 5 min (1 cycle) and 4°C hold. Cycling conditions for Gapdh: 95°C × 5 min (1 cycle), 95°C × 20 s, 60°C × 30 s, 72°C × 30 s (28 cycles), 72°C × 5 min (1 cycle) and 4°C hold. Band intensity from gel images was quantified using ImageJ.

In vivo electrophysiology
Electrophysiological experiments were conducted in head-fixed mice trained to walk on a rotating running wheel. Before training, mice were implanted with custom crowns permitting head-fixing above the wheel and allowed to recover for one week before training. Mice were trained to walk on the wheel for three 10-min sessions daily for at least one week. Following training, mice were implanted with a multi-electrode array targeting the S1DZ and allowed to recover before testing. To assess the IL-17a dependence of LPS-induced changes in neural activity, mice were injected intraperitoneally with control antibodies or blocking antibody against IL-17a (1 mg per mouse), 30 min before intraperitoneal injection of vehicle or LPS. Baseline neural activity was measured while mice were running on the wheel immediately before the first injection. After the first injection, mice were returned to their home cage. Post-injection (test) neural activity during wheel running was measured 4 h after injection of vehicle or LPS.

Multi-electrode array construction and implantation. Custom multi-electrode array scaffolds (drive bodies) were designed using 3D CAD software (SolidWorks) and printed in Accura 55 plastic (American Precision Prototyping), as previously described33,34. Before implantation, each array scaffold was loaded with 8–24 independently movable microdrives carrying 12.5-μm nichrome (California Fine Wire Company) tetrodes. Electrodes were pinned to custom-designed 32- or 128-channel electrode interface boards (Sunstone Circuits) along with a common reference wire (A-M systems).

Electrophysiological recordings and spike sorting. Signals were acquired using a Neurolynx multiplexing digital recording system (Neuralynx) through a combination of 32- and 64-channel digital multiplexing headstages plugged into the electrode interface board of the implant. Signals from each electrode were amplified, filtered between 0.1 Hz and 9 kHz and digitized at 30 kHz. Initial spike sorting was performed using MountainSort, followed by manual quality control using MClust toolbox (http://redishlab.neuroscience.umn.edu/mclust/MClust.html).

Analysis of firing rate. At time 0 in each recording session, mice began walking on the wheel rotating at 7.5 cm s⁻¹. Data collected between 1 and 2 s were included in the analysis. The firing rate was calculated across 1-s time windows during which mice were walking stably on the rotating wheel, averaged across 25–30 trials per condition. The first second after the onset of wheel rotation was omitted, to avoid firing rate changes due to acceleration. Firing rate was sampled with a 1-ms bin width passed through a box car filter (100 ms). The resulting peri-stimulus time histograms were then smoothed with a 50-ms Gaussian. To assess the effect of treatment on neural activity, changes in firing rates during wheel running 4 h after injection were normalized to the pre-injection baseline. Cells showing a firing rate below 0.1 Hz were excluded from analysis.

Statistics and reproducibility
Statistical analyses were performed using GraphPad Prism. Sample size was chosen on the basis of similar previous studies, and not on statistical methods to predetermine sample size46. Within each iteration of an experiment, mice were randomly assigned to groups with approximately balanced sample size. Behavioural results from mice with inaccurate targeting of viral infection or cannula implantations were excluded. Experimenters were blind to subject treatment during data collection and analysis.

Reporting summary
Further information on research design is available in the Nature Research Reporting Summary linked to this paper.

Data availability
Source Data for Figs. 1–4 and Extended Data Figs. 1–4, 6–10, containing raw data for all experiments, are provided with the paper. All other data are available from the corresponding author on reasonable request.

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Author contributions
M.D.R., Y.S.Y., J.R.H. and G.B.C. wrote the manuscript with input from the co-authors.

Competing interests
The authors declare no competing interests.

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Correspondence and requests for materials should be addressed to J.R.H. or G.B.C.

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**Extended Data Fig. 1** Cntnap2, Fmr1 and Shank3 mutant mice show variable sociability performance. a, Sociability performance. MIA, n = 13; wild-type, n = 22; Cntnap2 mutant, n = 71; Fmr1 mutant, n = 165; Shank3 mutant, n = 50; from 30 independent experiments. b, Time spent investigating social (S) versus inanimate (I) objects for mice described in a. c, d, Total interaction time (c) and distance travelled (d) during the three-chambered sociability experiments described in a. All n values refer to the number of mice used. Statistics calculated by one-way ANOVA with Dunnett’s post-hoc test (a, c, d) or two-way ANOVA with Dunnett’s post-hoc test (b). Graphs are mean ± s.e.m.
Extended Data Fig. 2 | Further behavioural analyses for sociability performance after treatment with LPS in PBS and MIA offspring, and monogenic mutant mice.  

**a**–**d**, Time spent investigating social (S) versus inanimate (I) objects (**a**), total interaction time (**b**), time spent in social (S), centre (C) or inanimate (I) chamber (**c**), and distance travelled (**d**) for sociability experiments in Fig. 1c. PBS offspring + vehicle, n = 10; PBS offspring + LPS, n = 9; MIA offspring + vehicle, n = 10; MIA offspring + LPS, n = 12; wild type (WT) + vehicle, n = 8; wild type + LPS, n = 11; Cntnap2 mutant + vehicle, n = 11; Cntnap2 mutant + LPS, n = 11; Fmr1 mutant + vehicle, n = 11; Fmr1 mutant + LPS, n = 13; Shank3 mutant + vehicle, n = 8; Shank3 mutant + LPS, n = 10; from 3 independent experiments. All n values refer to the number of mice used. Statistics calculated by two-way ANOVA with Sidak’s (a) or Dunnett’s (c) post-hoc test, or two-way repeated-measures ANOVA with Sidak’s post-hoc test (**b**, **d**). Graphs are mean ± s.e.m.
Extended Data Fig. 3 | LPS-induced rescue of MIA behavioural phenotypes is transient, effective in aged mice and extends beyond three-chambered sociability. a–e, Sociability measured 72 h after injection with vehicle (Veh) or LPS, in PBS and MIA offspring from Fig. 1c. Data expressed as per cent sociability (a), time spent investigating social (S) versus inanimate (I) objects (b), total interaction time (c), time spent in social (S), centre (C) or inanimate (I) chamber (d), and distance travelled (e) during three-chambered sociability experiments. PBS offspring + vehicle, n = 7; PBS offspring + LPS, n = 7; MIA offspring + vehicle, n = 8; MIA offspring + LPS, n = 6; from 2 independent experiments. f–j, Sociability measured before and 4 h after injection of vehicle or LPS in aged MIA mice (9–12 months old). Data are expressed as per cent sociability (f), time spent investigating social (S) versus inanimate (I) objects (g), total interaction time (h), time spent in social (S), centre (C) or inanimate (I) chamber (i), and distance travelled (j) during three-chambered sociability experiments. MIA offspring + vehicle, n = 6; MIA offspring + LPS, n = 7; from 2 independent experiments. k, Reciprocal social interactions measured after treatment with vehicle or LPS in PBS or MIA offspring. PBS offspring + vehicle, n = 9; PBS offspring + LPS, n = 9; MIA offspring + vehicle, n = 11; MIA offspring + LPS, n = 11; from 4 independent experiments. l, Marble burying index (percentage of buried marbles) measured before and 4 h after treatment with vehicle or LPS in PBS or MIA offspring. PBS offspring + vehicle, n = 12; PBS offspring + LPS, n = 12; MIA offspring + vehicle, n = 12; MIA offspring + LPS, n = 11; from 5 independent experiments. All n values refer to the number of mice used. Statistics calculated by two-way ANOVA with Sidak’s (a–c, e, g), Dunnett’s (d, i) or Tukey’s (k) post-hoc tests, or two-way repeated-measures ANOVA with Sidak’s post-hoc test (f, h, j, l). Graphs are mean ± s.e.m.
Extended Data Fig. 4 | Acute increase in body temperature is insufficient to promote sociability.

**a**, Body temperature profile after injection of vehicle or LPS in MIA offspring. Vehicle, $n = 10$; LPS, $n = 10$; from 4 independent experiments. The initial spike in body temperature is due to handling stress.

**b–e**, Data are expressed as time spent investigating social (S) versus inanimate (I) objects (**b**), total interaction time (**c**), time spent in social (S), centre (C) or inanimate (I) chamber (**d**), and distance travelled (**e**) during three-chambered sociability experiments described in Fig. 1g. $n = 9$ for all groups, from 2 independent experiments. All $n$ values refer to the number of mice used.

**f–j**, Sociability performance in Vgat-Cre PBS and MIA offspring after treatment with vehicle, CNO or LPS. Data are expressed as percent sociability (**f**), time spent investigating social (S) versus inanimate (I) objects (**g**), total interaction time (**h**), time spent in social (S), centre (C) or inanimate (I) chamber (**i**), and distance travelled (**j**) during three-chambered sociability experiments. PBS offspring, $n = 11$; MIA offspring, $n = 7$; from 2 independent experiments. All $n$ values refer to the number of mice used. Statistics calculated by two-way repeated-measures ANOVA with Bonferroni’s (**a**) or Dunnett’s (**f, h, j**) post-hoc tests, two-way ANOVA with Sidak’s (**b, g**) or Dunnett’s (**d, i**) post-hoc tests, or one-way repeated-measures ANOVA with Tukey’s post-hoc test (**c, e**). Graphs are mean ± s.e.m.
Extended Data Fig. 5 | Histological identification of the S1DZ. a, Coronal section of the cortex counterstained with DAPI to highlight the abrupt reduction in cell density in layer 4, between the S1DZ and the S1BF, at AP −0.46 mm. \( n = 5 \), from 1 independent experiment. D, dorsal; V, ventral. 

\[ b \] Coronal section of the cortex imaged with differential interference contrast, further highlighting the reduced layer 4 in the S1DZ at AP −0.46 mm. \( n = 3 \), from 2 independent experiments. All \( n \) values refer to the number of mice used. White arrows indicate borders of S1DZ. Scale bars, 300 µm (a); 300 µm (b).
Extended Data Fig. 6 | See next page for caption.
Extended Data Fig. 6 | Treatment of MIA offspring with LPS does not have a distinguishable effect on FOS expression in the other cortical regions analysed. a, Full cortical depth of S1DZ FOS staining as shown in Fig. 2a, for PBS and MIA offspring after administration of vehicle or LPS. PBS offspring + vehicle, n = 8; PBS offspring + LPS, n = 9; MIA offspring + vehicle, n = 13; MIA offspring + LPS, n = 11; from 3 independent experiments. Scale bar, 200 μm. b, c, Representative images (b) and quantification (c) of FOS (green) and NeuN (red) colabelled cells within the S1DZ of PBS and MIA offspring. PBS offspring, n = 4; MIA offspring, n = 3; from 1 independent experiment. Scale bar, 50 μm. d, e, Representative images (d) and quantification (e) of FOS expression in a series of cortical regions and after injection of vehicle or LPS in PBS or MIA offspring. Sections are stained for FOS (green) and DAPI (blue). Scale bars, 200 μm. For S1BF, M2, M1 and AuD: PBS offspring + vehicle, n = 8; PBS offspring + LPS, n = 9; MIA offspring + vehicle, n = 13; MIA offspring + LPS, n = 11. For mPFC: PBS offspring + vehicle, n = 8; PBS offspring + LPS, n = 9; MIA offspring + vehicle, n = 12; MIA offspring + LPS, n = 11. For V1: PBS offspring + vehicle, n = 7; PBS offspring + LPS, n = 8; MIA offspring + vehicle, n = 12; MIA offspring + LPS, n = 9; from 4 independent experiments. All n values refer to the number of mice used. Statistics calculated by unpaired two-tailed t-test (c) or two-way ANOVA with Tukey’s post-hoc test (e). Graphs are mean ± s.e.m.
Extended Data Fig. 7 | Further behavioural analyses of S1DZ optical-inhibition-mediated rescue of sociability in monogenic mutant mice.

**a**, Quantification of FOS-expressing cells in the S1DZ of monogenic mutant mice. Wild type, *n* = 6; Cntnap2 mutant, *n* = 21; Fmr1 mutant, *n* = 17; Shank3 mutant, *n* = 15; from 5 independent experiments. **b**, Correlation of FOS expression in the S1DZ with severity of sociability deficits across monogenic mutant mice. Cntnap mutant, *n* = 21; Fmr1 mutant, *n* = 17; Shank3 mutant, *n* = 15; from 4 independent experiments. Black solid lines represent regression line; grey lines indicate 90% confidence intervals. **c**, Individual data for experiments in Fig. 2f. **d–f**, Data are expressed as time spent investigating social (S) versus inanimate (I) objects (d), total interaction time (e), and distance travelled (f) during the three-chambered sociability experiments described in Fig. 2f. Wild type + EYFP, *n* = 7; wild type + NpHR, *n* = 8; Cntnap2 mutant + EYFP, *n* = 11; Cntnap2 mutant + NpHR, *n* = 9; Fmr1 mutant + EYFP, *n* = 8; Fmr1 mutant + NpHR, *n* = 12; Shank3 mutant + EYFP, *n* = 8; Shank3 mutant + NpHR, *n* = 10; from 6 independent experiments. All *n* values refer to the number of mice used. Statistics calculated by one-way ANOVA with Dunnett’s post-hoc test (a), linear regression (b), one-way repeated-measures ANOVA with Dunnet’s post-hoc test (c), two-way ANOVA with Sidak’s post-hoc test (d) or two-way repeated-measures ANOVA with Sidak’s post-hoc test (e, f). Graphs are mean ± s.e.m.
Extended Data Fig. 8 | See next page for caption.
Extended Data Fig. 8 | Il17ra expression in the S1DZ of PBS and MIA offspring and further behavioural analyses of S1DZ IL-17a rescue of sociability in MIA offspring and monogenic mutant mice. a, Representative images of Il17ra expression in the S1DZ of PBS and MIA offspring. Scale bar, 1 mm. b, Quantification of Il17ra expression within the S1DZ of MIA offspring according to cortical layer. n = 6, from 2 independent experiments. c, Quantification of overall Il17ra expression in the S1DZ of PBS and MIA offspring. PBS offspring, n = 8; MIA offspring, n = 6; from 2 independent experiments. d–h, Further behavioural analyses of experiments described in Fig. 3f. Time spent investigating social (S) versus inanimate (I) objects (e), total interaction time (f), time spent in social (S), centre (C) or inanimate (I) chamber (g), and distance travelled (h). PBS offspring + vehicle, n = 11; PBS offspring + IL-17a, n = 12; MIA offspring + vehicle, n = 14; MIA offspring + IL-17a, n = 10; wild type + vehicle, n = 11; wild type + IL-17a, n = 11; Cntnap2 mutant + vehicle, n = 8; Cntnap2 mutant + IL-17a, n = 10; Fmr1 mutant + vehicle, n = 9; Fmr1 mutant + IL-17a, n = 11; from 6 independent experiments. All n values refer to the number of mice used. Statistics calculated by unpaired two-tailed t-test (e), two-way ANOVA with Sidak’s (f) or Dunnett’s (g) post-hoc tests, or two-way repeated-measures ANOVA with Sidak’s post-hoc test (f, h). Graphs are mean ± s.e.m.
Extended Data Fig. 9 | IL-17a is necessary for LPS-induced behavioural rescue and reduction of FOS expression in MIA offspring. a–e. Further behavioural analyses of experiments described in Fig. 4a. Time spent investigating social (S) versus inanimate (I) objects (b), total interaction time (c), time spent in social (S), centre (C) or inanimate (I) chamber (d), and distance travelled (e). PBS offspring + vehicle + isotype, n = 9; PBS offspring + LPS + isotype, n = 11; MIA offspring + vehicle + isotype, n = 9; MIA offspring + LPS + isotype, n = 10; MIA offspring + LPS + anti-IL-17a, n = 10; from 7 independent experiments. f. Quantification of FOS-expressing cells in the S1DZ and CeA after injection of vehicle or LPS in MIA offspring pre-treated intracerebroventricularly with isotype-control antibody or blocking antibody against IL-17a (anti-IL-17a). PBS offspring + vehicle + isotype, n = 14; MIA offspring + vehicle + isotype, n = 9; MIA offspring + LPS + isotype, n = 10; MIA offspring + LPS + anti-IL-17a, n = 10; from 4 independent experiments. All n values refer to the number of mice used. Statistics calculated by two-way ANOVA with Sidak’s (b) or Dunnett’s (d) post-hoc tests, two-way repeated-measures ANOVA with Sidak’s post-hoc tests (c, e) or one-way ANOVA with Dunnett’s post-hoc test (f). Graphs are mean ± s.e.m.
Extended Data Fig. 10 | See next page for caption.
Extended Data Fig. 10 | Further analyses of the necessity of IL-17a for the LPS-induced reduction of firing rate in the S1DZ, and the necessity of S1DZ IL-17Ra expression for the LPS-induced rescue of sociability deficits in MIA offspring. 

a–c, Further analyses for experiments described in Fig. 4b–d. 

a, Example of a head-fixed mouse on the running wheel used during single-unit recording. 
b, Representative image of a tetrode placement in the S1DZ. Scale bar, 500 μm. 
c, Firing rate for individual cells before and 4 h after injection of vehicle or LPS in PBS and MIA offspring pre-treated with isotype-control antibody or blocking antibody against IL-17a (anti-IL-17a).

d–h, Further analyses for experiments described in Fig. 4e, f. 

d, Time spent investigating social (S) versus inanimate (I) objects (e), total interaction time (f), time spent in social (S), centre (C) or inanimate (I) chambers (g), and distance travelled (h). R17rafl/fl; EYFP, n = 9; R17rafl/fl;EGFP:nCre, n = 10; from 5 independent experiments.

Il-17rafl/fl ;EYFP, n = 9; Il-17rafl/fl ;EGFP:nCre, n = 10; from 5 independent experiments.

i, j, Representative images (i) and corresponding quantification (j) of Il17ra and Gapdh amplicon following PCR using cDNA derived from cells isolated from the cortical region centred on S1DZ of Il-17rafl/fl;EYFP and Il-17rafl/fl;EGFP:nCre mice. n = 3 for both groups; from 1 experiment. All n values refer to the number of mice used. Statistics calculated by two-way ANOVA with Sidak’s post-hoc test (e, g), two-way repeated-measures ANOVA with Sidak’s post-hoc tests, (f, h) or unpaired two-tailed t-test (j). Graphs are mean ± s.e.m.
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- For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted. Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
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- Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

Data collection

Behavioral experiments were recorded using Ethovision XT. IL-17R knockout and c-Fos+ cell quantification were performed using FIJI (ImageJ 1.49m). qPCR data for SFB quantification were collected using Lightcycler R 96 SW1.1. In situ and immunohistochemistry automated quantification was performed using QuPath. Single unit recording data were acquired using a Neuralynx multiplexing digital recording system. Spike sorting for single-unit recording data was performed using MountainSort and the MClust toolbox.

Data analysis

Data were analyzed using Graphpad Prism v8.0.1.

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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | Sample size was chosen based on similar previous studies (8,15) |
|-------------|---------------------------------------------------------------|
| Data exclusions | Behavioral results from mice with inaccurate targeting of viral infection or cannula implantations were excluded. |
| Replication | Key experiments were reiterated with representatives from each group with similar observations across iterations. |
| Randomization | Within each iteration of an experiment, animals were randomly assigned to groups with approximately balanced sample size. |
| Blinding | Experimenters were blind to subject treatment during data collection and analysis. |

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**Materials & experimental systems**

| n/a | Involved in the study |
|-----|-----------------------|
| ☑ Antibodies |
| ☑ Eukaryotic cell lines |
| ☑ Palaeontology |
| ☑ Animals and other organisms |
| ☑ Human research participants |
| ☑ Clinical data |

**Methods**

| n/a | Involved in the study |
|-----|-----------------------|
| ☑ ChIP-seq |
| ☑ Flow cytometry |
| ☑ MRI-based neuroimaging |

**Antibodies**

Antibodies used:

- chicken anti-GFP (1:1000, Ab5450, Abcam),
- rabbit anti-cFos (1:500, ABE457, Millipore),
- rabbit anti-DsRed (1:1000, 632496, Clontech),
- mouse anti-NeuN (1:1000, MAB377, Millipore).

Validation:

These are all commonly used antibodies. Statements regarding validation can be found at the manufactures website.

**Animals and other organisms**

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**Laboratory animals**

WT mice used for experiments were male C57BL/6 mice. Male IL17Ra KO, IL17Ra(fl/fl), Cntnap2, Fmr1, Shank3, and Vgat-Cre mice in the C57BL/6 background were also used for experiments. Sources for all mice used are reported in the methods.

**Wild animals**

Study did not involve wild animals.

**Field-collected samples**

Study did not involve field-collected samples.

**Ethics oversight**

Committee on Animal Care at Massachusetts Institute of Technology provided approval for all experiments.

Note that full information on the approval of the study protocol must also be provided in the manuscript.