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SYNGAP1 Heterozygosity Disrupts Sensory Processing by Reducing Touch-Related Activity within Somatosensory Cortex Circuits

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Abstract

In addition to cognitive impairments, neurodevelopmental disorders (NDDs) often result in sensory processing deficits. However, the biological mechanisms that underlie impaired sensory processing associated with NDDs are generally understudied and poorly understood. We found that *SYNGAP1* haploinsufficiency in humans, which causes a sporadic neurodevelopmental

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Custom code was written in MATLAB to analyze intrinsic imaging experiments. Minor code modifications to the FluoroSNNAP15.04.08 plugin ⁵⁹ were used to analyze somatic GCaMP6 signals. These scripts are available upon reasonable request.

disorder defined by cognitive impairment, autistic features, and epilepsy, also leads to deficits in tactile-related sensory processing. *In vivo* neurophysiological analysis in *Syngap1* mouse models revealed that upper-lamina neurons in somatosensory cortex (SSC) weakly encode information related to touch. This was caused by reduced synaptic connectivity and impaired intrinsic excitability within upper-lamina SSC neurons. These results were unexpected given that *Syngap1* heterozygosity is known to cause circuit hyperexcitability in brain areas more directly linked to cognitive functions. Thus, *Syngap1* heterozygosity causes a range of circuit-specific pathologies, including reduced activity within cortical neurons required for touch processing, which may contribute to sensory phenotypes observed in patients.

Keywords

Intellectual Disability; Epilepsy; Autism; Circuitry; Sensory processing; Synapse; Syngap1

INTRODUCTION

Neurodevelopmental disorders (NDD) often result in poor cognitive functioning and impaired behavioral adaptations. While the precise neurobiological and neurophysiological mechanisms that underlie these impairments remain unknown, it is generally understood that they are a consequence of impaired connectivity and processing within neuronal networks¹⁻⁴. However, it remains unclear what brain areas are disrupted and how neural processing within affected areas contributes to cognitive and behavioral impairments. Therefore, it is critical to identify dysfunctional brain areas that contribute to NDD-relevant phenotypes and to then define the circuit-based mechanisms that contribute to them.

Sensory processing impairments are nearly ubiquitous in NDDs, with a growing appreciation for their central role in behavioral and cognitive deficits associated with these⁵, and related⁶, disorders. Mechanistic studies in NDD animal models have traditionally focused on the circuits and synapses located in brain regions associated with cognitive functions⁴. These studies have demonstrated that NDD risk genes impair synapse properties, such as plasticity and excitatory-inhibitory ratios, which are believed to underlie impaired cognitive functions and common comorbidities, such as epilepsy. However, less attention has been paid to the mechanisms contributing to sensory processing deficits, particularly at the level of primary cortical circuits, which integrate ascending sensory information with topdown modulatory signals from higher cortical areas⁷. Indeed, higher forms of cognition require information regarding the external environment. Primary sensory cortical areas decode stimulus features⁸ and facilitate the construction of complex internal representations of the external world⁹. In this context, disrupted lower-level sensory processing in NDDs could contribute to generalized cognitive and behavioral impairments reported in patients. At present, it remains unclear how impairments in sensory processing contribute to complex cognitive and behavioral phenotypes common to NDD patients. This stems from a relative lack of mechanistic, systems-based studies investigating how highly-penetrant NDD risk factors directly impact the function of circuits that process sensory information.

A tractable entry point for such mechanistic studies is to perform in-depth biological investigations of highly-penetrant rare variants known to cause well-understood childhood NDDs^{10, 11}. The *SYNGAP1* gene is frequently mutated in intellectual disability (ID) associated with classically undefined global developmental delay¹²⁻¹⁵ and is a major risk factor for autism spectrum disorders^{16, 17}. *De novo* nonsense variants in *SYNGAP1* resulting in haploinsufficiency lead to a genetically-defined form of ID (MRD5; OMIM#603384) that may explain up to 1% of these cases^{18, 19}. MRD5 patients suffer from cognitive impairments, such as absent or poor language acquisition, and very low non-verbal IQ^{19, 20}. *SYNGAP1* is also a risk factor for epileptic encephalopathies^{21, 22} and most MRD5 patients have comorbid epilepsy¹⁸⁻²⁰. Currently, the impact of SYNGAP1/*Syngap1* pathogenicity on sensory functions is unknown, but is necessary for a deeper understanding of the complex phenotypes observed in this genetically-defined NDD.

We report altered sensory functions in a *SYNGAP1* patient population, including behaviors related to abnormal tactile processing. Studies in a series of construct-valid mouse models of *SYNGAP1* haploinsufficiency revealed severe impairments in somatosensory cortex (SSC) circuit structure and function. *Syngap1* heterozygosity resulted in reduced connectivity and hypoexcitability within upper-lamina SSC glutamatergic neurons, resulting in reduced touch-related activity within these circuits. These results were surprising, as they were distinct from mechanisms described in associational brain areas more directly linked to cognitive function, such as increased synaptic excitability of circuits in the hippocampus²³ and prefrontal cortex²⁴. Thus, pathogenicity of *Syngap1* causes a range of circuit-specific pathologies. We propose a generalizable scheme where interactions among region-specific circuit pathologies due to causally-linked NDD risk variants drive complex cognitive and behavioral phenotypes observed in patients.

RESULTS

SYNGAP1 heterozygosity in humans leads to touch-related sensory processing defects

To explore how sensory function is impacted by SYNGAP1 pathogenesis, we mined data from a SYNGAP1 patient registry that is a component of an ongoing retrospective MRD5 Natural History Study (NHS). Registries are essential tools to discover common phenotypes in patients with rare genetic disorders²⁵. This registry contains a searchable database with anonymized medical records, including genetic reports and detailed medical histories for SYNGAP1 patients (https://syngap1registry.iamrare.org). Forty-eight unique registry entries contained a comprehensive medical history and completed questionnaire focusing on sensory function. Forty-five of them exhibited features consistent with sensory processing impairments (Supplementary Table 1). Twenty of these entries were supported by detailed narrative accounts describing abnormal responses to tactile stimuli, which included blunted responses to painful touch-related stimuli and/or tactile-seeking or tactile-aversive behaviors (Supplementary Table 2). Genetic reports were available for 17 of the 20. Fourteen of the seventeen reports identified the SYNGAP1 variant as clearly pathogenic (i.e. caused SYNGAP1 heterozygosity) and the cause of their developmental disorder (Supplementary Table 2). The other three reports included variants of undetermined, but potential, clinical significance that require further biological validation, such as predicted splice alterations or

missense mutations. Data from these detailed patient entries indicate that disrupting expression or function of *SYNGAP1* in humans can lead to tactile-related sensory impairments in addition to cognitive impairment and seizure.

Syngap1 heterozygosity in mice causes touch-related deficits in cortical circuit activation

We next utilized mouse models to understand the biological mechanisms of sensory processing deficits associated with human SYNGAP1 heterozygosity. The barrel field of the SSC processes touch-related sensory information generated by movements and angular deflections of whiskers⁸. Computations here promote touch-related sensory perception and object localization, and facilitate the creation of spatial maps of the environment^{8,9}; all processes that facilitate higher-order cognitive functions. First, we mapped cortical receptive fields of whiskers in anesthetized Syngap1 mice by measuring intrinsic optical signals (IOS) generated by whisker deflections (Fig. 1a-b). The amplitude of cortical IOS elicited from C2 and β whisker deflections, as measured by the peak IOS response (Fig. 1c-e) and the area of the absolute value of the thresholded signal (Fig. 1g), was significantly reduced in both whisker-related receptive fields tested in Syngap1 heterozygous mice (Hets). This result was surprising given that Syngap1 pathogenicity has routinely been linked to circuit hyperexcitability ^{18, 21, 23, 24, 26, 27}. However, both the distance between the two fields (Fig. 1f) and their size, as measured by the relative area of the IOS signal (Fig. 1h), did not differ from WT. Reduced cortical activation in Syngap1 mouse barrel SSC was confirmed by widefield, low-resolution imaging of barrel cortex in Thy1-GCaMP6s mice (GP4.3 line²⁸) crossed to Syngap1 mice under anesthesia. In these studies, Syngap1 Het mice also exhibited reduced amplitudes of SSC whisker-evoked responses compared to controls (Supplementary Fig. 1a-1). Moreover, alterations in whisker-evoked signals in SSC were stimulus-dependent, with larger effect sizes occurring in response to stronger stimuli. These findings indicate that receptive field topology is generally unaltered in Syngap1 mice, but that cortical activation driven by whisker input is impaired by *Syngap1* heterozygosity.

To better understand altered SSC functional activation in Syngap1 mice, we measured spikelike supra-threshold somatic calcium events in Layer 2/3 SSC neurons. These SSC neurons were chosen because they are readily accessible by two-photon imaging and known to integrate bottom-up sensory signals originating in the periphery with information arriving from higher cortical areas⁷. Somatic calcium events were measured through *in vivo* twophoton imaging of GCaMP6 dynamics in awake, head-fixed Syngap1 mice crossed to the GP4.3 line (Fig. 2a-c). These studies were performed in awake animals because anesthesia disrupts neuronal activity and can have complex effects in mouse models of NDD risk genes²⁹. We mapped the cortical receptive field of a single whisker and then loaded this whisker into a small plastic holder. Mice spontaneously whisked during the imaging trials, which resulted in the whisker contacting the sides of the holder (Supplementary Video 1). Thus, recorded activity during these trials was comprised of ongoing, spontaneous activity of unknown origins and activity generated by whisker movements and/or touch. Neurons in L2/3 SSC generally appeared less active in Hets compared to WTs (Fig. 2b). Further analysis revealed that while the size of the responsive population was not affected in Syngap1 Hets (Fig. 2d), neurons that were active during the imaging session had significantly smaller (Fig. 2e) and less numerous (Fig. 2f) events compared to WT neurons.

After these imaging sessions, we injected the whisker pad with onabotulinumtoxinA (Botox) to paralyze whisker movements (Supplementary Video 2) and reimaged L2/3 cellular activity in the same animal population one week later. Botox had no effect on the size of the responsive population (Fig. 2d). Botox also had no effect on event amplitudes within genotypes and the difference between genotypes remained even in the presence of whisker paralysis (Fig. 2e). However, Botox had a clear impact on the number of detected events in this experiment. For example, in WT animals, whisker paralysis shifted the cumulative fraction of neuronal activity counts to the left (Fig. 2f). We reasoned that the shift in the "event number" curves after Botox injection reflects the contribution of whisker movements and/or touch to activity within this population. In contrast, Botox did not shift the activity of Het neurons (Fig. 2f). Neurons from Botox(+) Hets appeared to have activity counts that were identical to neurons in Botox(-) Hets, indicating that whisker movements and/or touch results in less activity compared to WTs. Indeed, the Het activity curves were superimposed onto that of Botox-treated WT neurons. While these shifts in spike-like activity may seem subtle, it is known that spiking within a small population of cortical neurons is sufficient to drive a behavioral response 30 . Thus, it is reasonable that a small, yet highly significant change in the number of detected spike-like events generated in Hets, especially spike-like activity linked to touch, is likely to be behaviorally meaningful to these animals.

The lack of Botox effects on neuronal activity in Het mice was supported by a cluster analysis of activity counts from neurons imaged in all four conditions. In each condition, active neurons were clustered into low-, medium- and high-activity populations³¹ based on the number of spikes during the recording session (Supplementary Fig. 2a). From this analysis, there was an overall effect on how the neurons clustered across the four groups (Supplementary Fig. 2b) and there was a difference in each of the three activity clusters when comparing the four experimental groups (Supplementary Fig. 2c). However, post hoc analyses comparing the four groups to each other revealed that all three activity clusters from the Botox(-) WTs were different from the corresponding clusters from all other groups (Supplementary Fig. 2d). The difference arose from more neurons present in the high- and medium-active clusters, and fewer in the low-active cluster, in the Botox(-) WT compared to the other three groups, further supporting the idea that Botox reduced activity within WT, but not Het, SSC circuits. Reduced activity generated from free whisking and/or touch in Syngap1 mice could be caused by impaired whisking behaviors in these animals, such as a decrease in total time whisking. Unexpectedly, Syngap1 Hets spent more time freely whisking compared to WTs (Fig. 2g). Taken together, these findings indicate activity driven by free whisker movement and/or whisker curvature driven by touch is poorly encoded by L2/3 SSC neurons in Syngap1 mice.

To further investigate the possibility that *Syngap1* heterozygosity impacts SSC cellular activity generated by whisker curvature, we measured cellular response properties in SSC L2/3 neurons evoked by piezo-driven, passive whisker deflections in awake, head-fixed *Syngap1/GP4.3* mice. We observed an effect of genotype on several measures of stimulus-evoked neuronal activity (Fig. 3a-h), each consistent with reduced activation of L2/3 SSC neurons. A mild stimulus (5 pulses, 5 Hz) resulted in a smaller responsive population in Hets (Fig. 3c), but not the size of individual responses or the probability that a cell responds to the stimulus (Fig. 3d, e). However, a stronger stimulus often used in single whisker-stimulation

detection paradigms, such as 60 pulses at 40 Hz, resulted in a smaller active population (Fig. 3f), a reduced amplitude of sensory-evoked events (Fig. 3g), and reduced response probability of responsive cells (Fig. 3h) in Hets. Active whisking could theoretically disrupt the quality of investigator-controlled passive deflections in a genotype-specific manner through degradation of precise whisker control. To control for this possibility, we performed stimulus evoked trials in the same cohort of animals following injection of Botox into the whisker pad. After whisker paralysis, we continued to observe reduced cellular sensory responsiveness arising from passive whisker deflections in Hets (Fig. 3i-n). For both stimuli, we observed a reduction in the response probably of the whisker-responsive population in Hets (Fig. 3k, n). However, other cellular response phenotypes were less pronounced after whisker paralysis, including the absence of change in the event amplitude of the responsive population (Fig. 3j, m) and the size of the responsive population (Fig. 3i, 1). Together, these data demonstrate that whisker bending through passive deflections results in reduced activity within SSC of *Syngap1* mice.

We next asked if cortex-specific mechanisms contribute to reduced SSC activity observed in Syngap1 Hets by restricting Syngap1 heterozygosity to forebrain glutamatergic neurons. Syngap1 conditional KO mice²³ were crossed to a series of Cre drivers that express the recombinase in different sub-classes of forebrain neurons, and then injected these animals with an AAV vector that drives GCaMP6s expression. Syngap1 cKO mice were first crossed to an Emx1-Cre driver³², where Cre expression is restricted to forebrain glutamatergic neurons and glia, and performed in vivo two-photon imaging of calcium dynamics in L2/3 SSC neurons (Fig. 4a-b). We have previously confirmed that this cross results in reduced SynGAP expression within forebrain excitatory neurons²⁴. Two-photon calcium activity measurements were obtained from anesthetized mice to prevent spontaneous whisking, allowing precise control of stimulus conditions. Restricting Syngap1 pathogenicity to this cellular population did not change the size of the active cell population (Fig. 4c), but did disrupt the amplitude and neuronal response probability of sensory-evoked responses (Fig. 4d, e). These data were largely consistent with findings in awake, conventional nonwhisking awake Syngap1 animals (i.e. Botox-injected animals shown in Fig. 3i-k), though the effects here were stronger than in this prior experiment. The stronger phenotype in this experiment could be due to the method (virally injected GCaMP6s vs. GP4.3 transgene expression), brain state (anesthesia vs. awake), the cell types expressing Syngap1 heterozygosity (EMX1+-restricted population vs. no restrictions), or some combination of these factors. Never-the-less, reduced sensory responsiveness of L2/3 SSC neurons in Syngap1 mice is a reproducible phenotype and implicates Syngap1-mediated pathology directly within forebrain neurons.

Syngap1 cKOs were next crossed to Cre driver lines that induce recombination in nonoverlapping EMX1-positive subpopulations and imaged under the same conditions as the EMX1 experiment. Restricting Syngap1 heterozygosity to upper-lamina cortical neurons in Cux2-CreERT2 mice, an extensively validated Cre driver line selective for upper-lamina neurons in neocortex ³³ (Supplementary Fig. 3a-b), did not impact the size of the responsive population or neuronal response probability (Fig. 4f, h). However, it caused a weak, but significant, reduction in amplitude of evoked responses (Fig. 4g). Next, we restricted *Syngap1* heterozygosity to L5 glutamatergic neurons using the validated Rbp4-Cre diver

line³³ (Supplementary Fig. 3c-d), and imaged calcium responses in L2/3 barrel cortex neurons. Disrupting *Syngap1* only in these neurons did not alter any of the standard measures of neuron responsiveness to sensory stimulation (Fig. 4i-k). While we have measured reduced whisker-evoked activity within Layer 2/3 neurons in both conventional and EMX1-*Syngap1* Het mice, the primary cellular origins of how *Syngap1* heterozygosity leads to effects on Layer 2/3 neurons is complex and not necessarily autonomous to only these neurons. The inability to fully recapitulate the sensory-responsive phenotypes of EMX1-Cre models in Cux2-Cre or RBP4-Cre models indicates that *Syngap1* pathogenicity directly alters the function of multiple EMX1-positive populations that converge to drive the SSC cellular responsiveness phenotypes observed in conventional *Syngap1* Hets.

Reduced sensory responsiveness could be caused by decreased excitation or increased inhibition onto, or within, upper-lamina L2/3 SSC neurons. To test the latter possibility, we crossed *Syngap1* mice with Gad2-T2A-NLS-MCherry mice⁷ and injected them with an AAV vector that drives GCaMP6s expression in inhibitory and excitatory neurons (Fig. 5a-b). Under anesthesia, mCherry-negative (excitatory) neurons exhibited a substantial reduction in neuronal activation in response to the passive whisker stimulation (Fig. 5c-e). The reduced activation of neurons was most apparent in the whisker-responsive population (Fig. 5e). Amongst this population of excitatory neurons, *Syngap1* disruption appeared to preferentially affect the most active cells. Similarly, there was a reduction in whisker-evoked neuronal activation from mCherry-positive (inhibitory) neurons (Fig. 5f-h). A reduced probability of firing in inhibitory neurons in response to whisker stimulation suggests that *Syngap1* heterozygosity does not result in an overactive population of GABA-releasing neurons.

Deficits in touch-related cortical circuit activation in Syngap1 mice are associated with reduced synaptic and intrinsic excitability of upper-lamina SSC neurons

This result prompted us to investigate impaired excitation as a possible cause of reduced sensory-related activity within upper-lamina SSC networks. We performed *in vivo* whole-cell patch-clamp recordings in L2/3 neurons of barrel cortex to determine how *Syngap1* heterozygosity affects whisker-evoked synaptic potentials in anesthetized mice. Synaptic depolarization in response to passive whisker stimulation was reduced in Hets compared to WTs (Fig. 6a-c), a finding consistent with reduced whisker-driven cellular activity within upper-layer SSC circuits. We did not observe any alterations in spontaneous Up/Down state properties or changes in resting membrane potential in neurons from Hets (Supplementary Table 3).

We further hypothesized that structural impairments in L2/3/4 neurons may contribute to reduced whisker-evoked feed-forward excitation in upper-lamina SSC circuits. L4 stellate cells in SSC receive the bulk of sensory-related information arriving from subcortical areas^{8, 34}. Anatomical assessment of digitally reconstructed L4 neurons showed they were smaller in *Syngap1* Hets compared to WTs (Fig. 7a), with dendritic arbors having reduced overall complexity and length. Dendrites from L4 neurons also had reduced spine density (Fig. 7b). Similar anatomical disruptions were found within dendrites and spines of L2/3 pyramidal cells from Hets (Fig. 7c-d), which receive a dense ascending projection from L4

stellate cells that relay sensory-related information to superficial neurons. These findings suggested that there were fewer excitatory synapses in upper-lamina SSC neurons.

Patch-clamp recordings from L2/3 SSC neurons prepared from acute brain slices supported this idea. mEPSC frequency was dramatically reduced in neurons from Hets (Fig. 7e, 7g). Interestingly, mEPSC amplitude was slightly increased in these same neurons (Fig. 7f), possibly reflecting homeostatic compensation arising from too few excitatory synaptic inputs³⁵. To directly measure feed-forward excitation within upper-lamina SSC circuits, we recorded synaptic currents in SSC L2/3 neurons evoked by electrical stimulation of L4 (Fig. 7h). Evoked synaptic currents in L2/3 neurons were reduced in amplitude in Hets compared to WTs (Fig. 7i, j). Finally, it is known that homeostatic compensation of both synaptic and intrinsic neuronal properties contribute to producing stable firing rates. We therefore wondered how intrinsic excitability mechanisms might be engaged to potentially counteract changes in reduced synaptic input onto, and overall activity within, upper-lamina L2/3 SSC neurons. Surprisingly, intrinsic excitability in L2/3 neurons was decreased in *Syngap1* Hets compared to controls (Fig. 7k-m), which may also contribute to reduced activity observed in these neurons.

Touch-related behaviors are impaired in Syngap1 mice

Impairments in SSC circuit function in *Syngap1* mice may be predictive of whiskerdependent behavioral deficits. To test this, we first explored the ability of constitutive *Syngap1* mice to detect novel objects exclusively through touch. Importantly, *Syngap1* mice are known to have normal novel object recognition (NOR) memory³⁶. Therefore, we could determine how potential deficits in touch perception may influence novelty detection of similar objects that differ only by subtle change in their surface texture (Fig. 8a). Individual objects with differing textures were equally salient to test subjects (Fig. 8b). During the learning phase, WTs and Hets explored identical objects for a proportionally similar amount of time, though Hets spent more total time exploring objects overall (Fig. 8c-d), which may be related to increased locomotion in this line³⁷. During testing with inclusion of the novel object, WTs distinguished between distinct textures, as evidenced by increased exploration of the novel relative to familiar object (Fig. 8c, e). In contrast, Hets failed to show a bias toward the novel object (Fig. 8d, e). Thus, Hets cannot discriminate between similar objects that differ only by surface texture.

We next used a discrimination task that requires the perception of a stimulus similar to what was used for evoking neuronal activity in our functional studies (Figs. 1-6). We selected a Go/NoGo paradigm, where water-deprived mice learn to perceptually report passive deflections of their whiskers by licking a water-dispensing port during a short "answer" period^{38, 39} (Fig. 8f). This type of learning paradigm is attractive for probing touch-related behaviors because it requires an intact ability to perceive a whisker stimulation³⁹ and activation of whisker-responsive cortical circuits to sufficiently drive learning⁴⁰. Task performance is quantified by the probability of correct choices during two trial types, "Go" (presence of whisker stimulus cue) and "NoGo" (absence of whisker cue) (Fig. 8g). WTs learned to discriminate between trial types after sufficient training (Fig. 8h; Supplementary Fig. 4a-b) with trial discrimination dependent on the strength of whisker stimulation (Fig. 8i-

j), confirming task performance is driven by a stimulus precept computed by the function of whisker-sensitive circuits. Age-matched cohorts of Syngap1 WTs and Hets were then trained, with 86% of WT mice and 0% of Het mice reaching the training goals (Fig. 8k). WTs learned to lick during "Go" trials and withhold licks during "NoGo" trials (Fig. 8h; Supplementary Fig. 4a-b). In contrast, Hets did not improve accuracy in either trial type, likely explaining their static task performance over time (Fig. 81; Supplementary Fig. 4c-d). Importantly, we observed a significant difference between genotypes in the fraction of correct answers over the entire training interval (Fig. 8m) and in the trial discrimination index (Fig. 8n), an objective measure of overall task performance at the completion of training. There was no difference between genotypes in relative weight gain/loss over the training interval (Supplementary Fig. 4e) and Hets exhibited normal licking, indicating familiarity with use of the port (Supplementary Fig. 4f-g). There was also no difference between genotypes in the response times for either trial type (Supplementary Fig. 4h-i) or total trials during training (Supplementary Fig. 4j-1). We also found no evidence of impulsive responding in Hets (Supplementary Fig. 4m-o). Together, these data indicate that poor task performance in Hets was not related to an obvious lack of motivation, training participation, or generalized impairment in the instrumental response.

DISCUSSION

The principle finding of this study is that disease-linked *SYNGAP1/Syngap1* variants lead to impaired sensory processing. Sensory abnormalities are common in NDD patient populations^{41, 42}. Moreover, there is a correlation between the severity of sensory disruptions and behavioral phenotypes in ASD populations⁴³. Human imaging studies from NDD⁵ and schizophrenia patients⁶ have identified altered function of primary sensory areas, leading to the idea that altered sensory processing contributes directly to the complex phenotypes observed in patients. However, the neurophysiological mechanisms that lead to NDD-associated sensory impairments are poorly understood. By mining a *SYNGAP1* patient registry within a retrospective Natural History Study, we confirmed that clinically-significant sensory alterations exist in patients with *SYNGAP1* haploinsufficiency, including abnormal responses to painful tactile stimuli and unusual touch-related behaviors (Supplementary Table 2). Moreover, by utilizing *Syngap1* mouse models, we uncovered unexpected circuit-level mechanisms consistent with impaired touch-related cortical sensory processing. Thus, reverse translation of NDDs caused by single rare variants, such as *SYNGAP1*, hold promise for better understanding these pathobiological mechanisms.

Our findings demonstrate that *Syngap1* heterozygosity reduces activity within upper-lamina SSC circuits of awake *Syngap1* mice. Reduced measures of sensory-evoked activity *in vivo* persisted in awake animals with paralyzed whiskers and in animals under anesthesia. These results were unexpected because many prior reports describe increased excitability and synaptic function in neurons from *Syngap1* heterozygous mutant animals^{23, 24, 27, 44-47}. Indeed, *Syngap1* heterozygosity is believed to generally enhance neuronal excitation through shifting the synaptic E-I ratio in several types of neurons in areas of the brain linked directly to cognitive processing^{23, 44, 48, 49}. *Syngap1* is a risk factor for severe epilepsy²¹, and prior studies in this same *Syngap1* mouse line used in this study have described seizure and circuit hyperexcitability in the hippocampus and prefrontal cortex^{23, 24}. *Syngap1* pathogenicity also

accelerates the maturation of excitatory synapses during development, including several inputs within the hippocampus²³ and the thalamo-cortical synapse in L4 somatosensory cortex²⁷. Moreover, *Syngap1* pathogenicity reduces the level of GABAergic connectivity onto SSC pyramidal neurons²⁶. This loss of inhibitory control contributes to changes in oscillatory cortical rhythms in Syngap1 mice and is also consistent with increased excitability of neural circuits. The most parsimonious explanation for the differences between these prior reports and the current study is that *Syngap1* heterozygosity has celland region-specific effects. For instance, L2/3/4 SSC neurons in adult constitutive Syngap1 mice had smaller dendritic fields with reduced spine density, which contributed to reduced feed-forward excitation during sensory stimulation. However, deep-layer SSC neurons from adult Syngap1 mutants were previously shown to have normally-sized arbors and spine densities⁵⁰. Therefore, the overall impact of *Syngap1* heterozygosity on brain function is defined by a range of circuit-specific impairments that disrupt neuronal excitability and function in complex ways. Our findings, therefore, give insight into the complexity of how a single gene causally-linked to severe ID and epilepsy can disrupt the structure and function of neurons and circuits linked to cognitive processes. We propose a generalizable scheme where interactions among region-specific circuit pathologies caused by causally-linked NDD risk variants drive complex cognitive and behavioral phenotypes observed in patients. The challenge will be to understand how disparate circuit pathologies interact to disrupt behavior, impair cognition, and promote seizure. Furthermore, it will be important to understand to what extent other highly-penetrant ID risk genes cause similarly complex region- and/or circuit-specific pathologies.

Our results support the conclusion that measures of reduced neuronal activity within upperlamina of SSC circuits in Syngap1 mice were caused by a combination of lower intrinsic excitability and reduced synaptic connectivity onto L2/3 neurons in this area. This interpretation is supported by several key pieces of data. First, measures of reduced activity of SSC L2/3 neurons can be attributed to effects of Syngap1 within this cellular population. Restricting Syngap1 heterozygosity to forebrain glutamatergic neurons and some glia [i.e. EMX1(+) populations] was sufficient to impair SSC neuronal activity evoked from passive whisker deflections. This indicates that a primary cause of impaired sensory-evoked activity within Syngap1 mutants arises, at least in part, from altered function of cortical circuits, rather than from disruptions in sensory coding occurring in sub-cortical areas, such as in the mechanoreceptor neurons or thalamic relays. Second, evoked L2/3 SSC neuronal activation was depressed in both GABAergic and glutamatergic cellular populations, indicating that reduced activity was not a consequence of increased inhibition. Third, in vivo patch-clamp recording demonstrated reduced whisker-evoked membrane depolarization in L2/3 neurons from Het mice, which is consistent with our primary observation of reduced cellular activity of neurons in this area. In measurements from acute slices, these neurons also had reduced intrinsic excitability, which may contribute to reduced membrane depolarization observed in vivo. Fourth, we observed that both L4 and L2/3 neurons from Het mice had smaller dendritic fields. We also observed that these shorted dendritic fields contained fewer dendric spines. This anatomical defect translated into reduced functional synaptic connectivity, as we observed reduced mEPSC frequency in Het neurons even though there was a slight increase in mEPSC amplitude. Increased amplitude mEPSC in L2/3 neurons from Hets is

consistent with our prior observation of larger dendritic spines in these cells⁵⁰, which may be a homeostatic adaptation to neurons that have reduced activity³⁵. We also observed direct evidence of reduced feed-forward excitation within upper-lamina of SSC. There was a substantial decrease in synaptic input from L4-to-L2/3 in thalamocortical slices prepared form *Syngap1* mice. A strength of this study is that large impairments in synaptic function within whisker-responsive circuits translated into a significant reduction in touch-related proxy measures of somatic spiking (i.e. super-threshold somatic GCAMP6 events) within these same circuits. Taken together, these data are consistent with a model where L2/3 neurons in *Syngap1* mice poorly encode incoming sensory information due to reduced synaptic excitation within upper-lamina of SSC. Given that L2/3 neurons integrate bottom up sensory codes with top-down modulatory information ⁷, it is possible that the cortexspecific circuit pathologies uncovered in this study disrupt sensory processing related to learning and/or behavioral adaptations. Future studies will be necessary to causally-link activity deficits within cortical circuits that respond to sensory input in *Syngap1* mice to relevant behavioral phenotypes, such as poor learning.

ONLINE METHODS

Collection and analysis of data from the Retrospective SYNGAP1 Natural History Study Registry

The SYNGAPI Patient Registry (https://syngap1registry.iamrare.org) is funded through the National Organization of Rare Disorders. This study was approved through the Hummingbird Institutional Review Board and meets all relevant ethical regulations for protections for human subjects. It is actively managed by a board of trustees comprised of a team of seven stakeholders, including parents with affected children, clinician-scientists that care for MRD5 patients, and neurobiologists that study the gene. The SYNGAP1 (MRD5) Natural History Study Registry is a retrospective longitudinal web-based observational natural history study. Parents or guardians provided informed consent prior to depositing medical history data into the registry. Participants with SYNGAP1 (MRD5) will be followed throughout the course of their lives with either the participant or authorized respondents contributing data at varying intervals throughout the course of the study. Initially, when a new patient is registered, data is collected on demographics, quality of life, medical history including genetic reports, disease phenotypes, event episodic data, retrospective data, participant review of systems, medication and diagnostic data. Each registrant is given a unique identifier to facilitate anonymization of patient data. Initial data collection is done through a series of questionnaires, including a survey of sensory and sensory-related issues. The structure of the database and all questionnaires were reviewed and approved by the members of the Board of Trustees.

To acquire information of possible sensory alterations in the *SYNGAP1* patient population, the registry database was queried for all entries that answered the sensory questionnaire. The questionnaires for each anonymized entry were then exported to a spreadsheet for analysis (Supplementary Table 1). Entries that noted obvious impairments in tactile alterations, typically though narrative descriptions in columns E, F and/or K of Supplementary Table 1, were placed in Supplementary Table 2. The Registry was then revisited to determine if

genetic reports were available for these entries. Genetic reports were available for 17 of these 20 entries. The presence/absence of an anonymized genetic report for each entry was noted in Column A and for entries with a report, the type of variant was listed in Column B and pathogenicity in Column C. Importantly, decisions on which entries to include in Supplementary Table 2 were made with no prior knowledge of each patient's genotype. Entries containing narratives were unedited, including spelling errors, except when necessary to protect patient information (i.e. redact a name).

Mice

All animal procedures were conducted in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and all protocols were approved by The Scripps Institutional Animal Care and Use Committee. The design and maintenance of the conventional and conditional Syngap1 lines have been described previously^{23, 51}. Thy1-GCaMP6s4.3 (#024275), Gad2-NLS-mCherry (#023140), and the TdTomato Ai9 (#007905) reporter lines were purchased from Jackson Laboratories. Rbp4-Cre (037128-UCD) and Cux2-CreERT2 (032779-MU) were purchased from MMRC. Both males and females were used in all experiments indiscriminately, except for the Go/NoGo task, where only males were used. All animals were older than 6 weeks of age at the beginning of experiments. Data collection occurred from mice >8 weeks of age. Mice were housed 4-5 per cage on a 12hour normal light-dark cycle, except for Go/NoGo experiments, where mice were housed on a reverse light-dark cycle. For experiments requiring chronic cranial window and head-post implantation, mice were singly housed following surgery, with added environmental enrichment consisting of cardboard huts or plastic running wheels (Bio-Serv, Flemington, NJ) for the remainder of the study. Animals expressing the Cux2-CreERT2 allele were injected with tamoxifen (once) at PND2 as previously described⁵⁰. Data collection was semi-randomized. Experimentalists were blind to genotype at the time of data acquisition and analysis. Generation of multiple transgenic mouse lines was labor, time and resource intensive. Additionally, most experiments required 1-3 months to complete, even with small sample sizes. This prevented us from picking WT and Het animals completely at random. Therefore, to obtain comparable sample sizes between genotypes, animal cohorts were generated by allocating equal (if possible) number of age-matched Syngap1 WT and Het littermates from separate litters, usually more than two. Then, animals were assigned a number to hide identity of genotype and/or group assignment. For imaging and behavior tasks, animals were recorded once per day in a randomized order while blinded to genotype. This process enabled balanced populations across experimental groups while minimizing potential biases. Only animals (or equivalent biological specimens) that died (or became non-responsive) during the course of the study or data collection procedures were excluded from analysis. The Life Sciences Reporting Summary contains additional details on data exclusions for specific experiments.

Intrinsic Optical Signal (IOS) Imaging

Animals were anesthetized with 1.6 g/kg urethane (Sigma-Aldrich, St. Louis, MO), followed by implantation of a custom head plate. The skull was scraped gently with a scalpel but otherwise left intact. The skull was thinned removing most of the spongy bone. Following gluing of the head plate, the skull was sealed with 1.5 % low melting point agarose dissolved

in lactate ringer's solution under a glass coverslip. Imaging was performed under a 4X objective on an upright microscope frame (BW51X; Olympus, Tokyo, Japan). The skull was illuminated with a 630 nm LED light mounted on the 4X objective⁵². The images were acquired with a Zeiss Axiocam camera (Carl Zeiss Microscopy Inc, Thornwood, NY) controlled by µManager software (Open Imaging, Inc.). Acquisition rate was approximately 10Hz. Whiskers were deflected using a piezoelectric bending actuator controlled by a linear voltage amplifier (Piezo Systems Inc, Woburn, MA). A single sinusoidal wave with a 5 ms rise and a 5 ms decay times were generated using Clampex software (Molecular Devices, Sunnyville, CA). Bending of the piezo was calibrated using a laser based displacement device (LD1610-0.5 Micro - Epsilon, Raleigh, NC). A single whisker deflection was approximately 200 μ m at 2 mm away from the whisker pad (~6 ⁰ or 1200 ⁰/s). Each IOS imaging trial consisted of a 2 s baseline imaging period followed by 20 deflections at 10 Hz. 50-70 trials were performed for each whisker (now called an imaging session) and averaged using IO and VSD Signal Processor plugin in ImageJ ⁵². Images taken between 1 s to 3 s after the start of the stimulus, were averaged, and defined as the response. IOS images were obtained by calculating (response-baseline)/baseline value for each pixel using custom scripts written in MATLAB (MathWorks, Natick, MA) according to established procedures ^{53, 54}. The animals that died during an intrinsic imaging experiment (due to anesthesia) or animals on which no visually reliable IOS map could be obtained (due to blood vessel contamination etc.) were excluded from the analysis (less than 5% of the cases). Image analysis was performed with the investigator blind to animal genotype. Briefly, images were first filtered with a Gaussian filter. Afterwards a baseline and a response region was manually selected in the final IOS image to minimize contamination by blood vessels^{53, 55}. Response size was determined as the minimum value of the response region subtracted by median of the baseline region. Image thresholding was performed in the response region in order to determine the area of activation. Thresholds based on the absolute response size were specified in the figures. Relative thresholding values were set at 50-80 % of the response size for each image.

GCaMP Widefield Imaging (GWI):

Thy-1 Gcamp6s4.3 mice were implanted with a chronic cranial window according to established procedures^{56, 57}. Briefly animals were anesthetized with avertin (Tribromoethanol, Sigma-Aldrich, St. Louis, MO) and intra-peritoneally injected with dexamethasone (4 mg/kg), rimadyl (carprofen 10mg/kg) and enroflox (enrofloxacin 5mg/kg). A 3 mm cranial window was made over the barrel cortex (A/P –2.0 mm, D/V +3.5 mm). The cranial window was sealed using two 3 mm glass coverslips glued onto a 5 mm glass coverslip. Animals were supplied with rimadyl for one week in drinking water for pain management. Following 2-3 weeks recovery from the surgery, animals were anesthetized with 1.6 g/kg urethane. Widefield fluorescent imaging of GCaMP6 was performed through the cranial window using 490 nm illumination under a standard 4X objective. Signals were acquired using a standard eGFP epifluorescence filter set. Whisker deflection and acquisition parameters were the same as with the IOS imaging, although we varied frequency and the number of deflections as specified in the figures. F/F images were obtained by calculating (response-baseline)/baseline for each pixel similar to intrinsic imaging. A fixed rectangle of 150 µm centered at the functionally defined barrel center

(90 % relative thresholding) was used for creating individual and averaged F/F traces. Bias was minimized in these analyses. Image analyses were performed blind to the animal genotypes and performed by automated MATLAB scripts (minimal investigator-driven selection artifacts).

In vivo GCaMP imaging in barrel cortex:

For awake Thy-1 Gcamp6s4.3 experiments, both male and female mice at least 6 weeks of age were fitted with a chronic cranial window and implanted with a titanium head-post according to established procedures with minor alterations³⁹. Briefly, animals were anesthetized with isoflurane (5 % induction, 1.5-2 % maintenance) and IP injected with a cocktail of dexamethasone (4 mg/kg), rimadyl (carprofen 10 mg/kg) and enroflox (enrofloxacin 5 mg/kg). Animals were mounted on a stereotaxic frame (David Kopf Instruments, Tujunga, CA) and body temperature was maintained with a thermal regulator (Harvard Apparatus, Holliston, MA). The scalp was shaved and sterilized with alternating swabs of betadine and 70 % alcohol. A small skin flap was removed, periosteum gently cleared, and the skull was scraped with a scalpel. A small, circular craniotomy was made over the left barrel cortex (3 mm diameter; center relative to bregma: lateral 3.5 mm; posterior 1.8 mm) using a dental drill, and the dura was left intact. The cranial window was sealed by gluing, two 3 mm glass coverslips glued onto a 5 mm glass coverslip, directly to the bone (VetBond, 3M). The titanium head-post was implanted by adhering it directly to the bone using VetBond, and then dental cement (Metabond, Parkell, Edgewood, NY). Animals recovered on a warm blanket before being placed back in their home cage. Rimadyl was injected (5 mg/kg) for three consecutive days after surgery for pain management. Following 1 week recovery from surgery, IOS imaging was performed through the cranial window, as described above, using light (0.5-1 %) isoflurane anesthesia to locate principle whisker areas (typically C2 whisker). The following week, animals were slowly habituated to headfixation with increasing time spent under head-fixation, up to one hour, which was continuously monitored via IR videography (Basler, acA640-120um). Mice were head-fixed in a custom-built stainless-steel body tube and mounting brackets. Noise associated with the resonant scanner was recorded and continuously played through speakers (Avisoft-UltraSoundGate, Avisoft Bioacoustics) within the microscope enclosure during habituation and imaging. Once mice were comfortable with head-fixation (typically four days), awake in vivo GCaMP imaging was performed. Mice were then injected with 0.5 MU of onabotulinumtoxinA (Botox, Allergan; prepared in PBS) into the right whisker pad under isoflurane anesthesia using a Hamilton syringe (2.5 µL 62 RN model, beveled tip), and allowed to recover for 2 days before repeating awake in vivo GCaMP imaging. Whisker paralysis, as demonstrated in Supplemental Movie 2, lasted ~5 days.

For Gad2-T2A-NLS-MCherry X *Syngap1* (i.e. Fig. 5) and Cre-driver experiments (i.e. Fig. 4), both male and female mice, at least 6 weeks of age were fitted with a chronic cranial window and implanted with a titanium head-post according to procedures detailed above. GCaMP6s was expressed following transduction with a rAAV (AAV1.Syn.GCaMP6s.WPRE.SV40, University of Pennsylvania Vector Core). During cranial window surgeries, iontophoresis via pulled-glass capillary micropipettes (Harvard Apparatus, Holliston, MA, 1.2 mm O.D., 0.69 mm I.D., inner tip diameter 20 um) was used

for infection at 4-6 sites within the craniotomy. Iontophoresis was performed with 5 uA at 7s 'on' and 7 s 'off' cycles for 5 min total per injection site at a depth of ~200 µm. Following 2-3 weeks recovery from surgery, IOS imaging was performed as described above, using light (0.5-1 %) isoflurane anesthesia, to locate principle whisker areas that overlapped with GCaMP6s expression (typically C2 whisker). *In vivo* GCaMP imaging was performed ~4 weeks following surgery under light (1-1.5 % isoflurane) anesthesia.

Imaging was performed with a VivoScope two-photon microscope equipped with a resonant scanner (Scientifica, UK). The light source was a Mai Tai HP 100 femtosecond-pulse laser (Spectra-Physics) running at 940 nm for GCaMP and 1040 nm for Mcherry imaging. The objective was a 16× water immersion lens with 0.8 NA (Nikon). Images were acquired using ScanImage 5 (vidriotechnologies.com). Functional images (512×512 pixels, 4X zoom, $150 \times 150 \mu$ m) of L2/3 cells (70-250 μ m under the pia) were collected at 10 Hz. 100% laser power was 165 mW at the front aperture of the objective. For all GCaMP imaging experiments, we used 30-60% power depending on the imaging depth. A similar number of imaging sessions at similar depths (~ 7 sessions starting > 70 μ m under the pia in 30 μ m intervals) were acquired for each animal.

Analysis of GCaMP activity in barrel cortex:

Calcium images were corrected for motion artifacts using the moco plugin in ImageJ⁵⁸. All subsequent analyses were performed in MATLAB R2015b using FluoroSNNAP15.04.08 plugin⁵⁹ with the following parameter choices. Regions of interest (ROIs) corresponding to identifiable cell bodies were selected manually. The fluorescence time course was measured by averaging all pixels within the ROI, then corrected for neuropil contamination. The neuropil ROIs were also manually drawn where there are no visible cell bodies and were the same for all cells within an imaging frame. After neuropil correction, the F/F of each ROI was calculated as (F-F0)/F0, where F0 was the mean of the lower 50% of the proceeding 10 s period. For the first 10 s period, minimum value of F0 was used (Patel et al., 2015). A template search-based algorithm was used in order to detect calcium events using built-in templates in FluoroSNNAP15.04.08. A correlation coefficient of 0.8 and F/F amplitude threshold of 15 % was used during template search. Spontaneous activity was recorded over 2 mins. Neurons with at least one detected calcium event during a 2 mins period were classified as spontaneously active neurons. Whisker stimulation induced activity was also recoded over a 2 mins period from the same ROIs. Whisker stimulation consisted of 5 whisker stimulations at 5 Hz or 60 whisker stimulations at 40 Hz, with an intertrain interval of 10 s with the same whisker stimulation parameters used in IOS imaging ($\sim 6^{\circ}$). Therefore, each single stimulus train lasted for 1 or 1.5 s. A total of 12 trains were given during a 2 mins period. Raw response probability was calculated as the total number of calcium events during the 1 or 1.5 s stimulation window divided by the total number of trains (12). In order to adjust for the probability of obtaining calcium events non-specifically during the stimulation detection window, we subtracted expected number of spontaneous calcium events from the raw calcium event count before dividing by 12. Cells with response probabilities greater than 0.083 (responding 1 out 12 stimulation) are considered responders. For Gad2-T2A-NLS-MCherry X Syngap1 experiments, mCherry fluorescence was recorded for 1 minute. Images were flattened and merged onto GCaMP6s images to delineate

inhibitory neurons from excitatory neurons. ROIs for excitatory and inhibitory neurons were analyzed in a similar fashion described above.

In vivo whole cell recordings:

A custom in vivo whole cell patch clamp system was built as described ⁶⁰. Mice were anesthetized with 1.6 g/kg urethane (Sigma-Aldrich, St. Louis, MO), followed by implantation of a custom head plate and a 1 mm craniotomy was made over the barrel cortex. Recordings were performed the same day in current clamp mode with the following internal solution in the electrode (mM): 130 potassium gluconate, 5 KCl, 10 HEPES, 10 sodium phosphocreatine, 0.4 EGTA, 1 Na-GTP and 4 Mg-ATP (pH 7.3, 285-290 mOsm). Electrophysiological signals were amplified with Multiclamp 700B (Molecular Devices, Sunnyville, CA), filtered at 2 KHz, digitized (10 KHz) with NI USB-6363 (National Instruments, Austin, TX) and recorded using NI acquisition system in MATLAB⁶¹. UP/ DOWN states were identified as described²⁹. Briefly, the signal was first filtered with a median filter (100 ms window) in order to eliminate action potentials. Voltage segments lasting at least 100 ms and higher or lower than a third of the standard deviation away from the signal mean were classified as either up or down states respectively. In order to obtain synaptic currents, whiskers were deflected 5 times at 5 Hz with intensity parameters used during imaging experiments. In order to obtain synaptic amplitudes, 50 trials were averaged, excluding the trials with the action potentials. Action potentials were defined as samples where the speed of depolarization exceeded 6 mV/ms and whose values were more than two standard deviations away from the mean.

Dendrite reconstructions: AAV systemic Injections, Tissue Clearing, Sholl Analysis

Dendrite reconstructions were performed in mice (*Syngap1*+/– Hets crossed to Ai9+/+) that were injected with an rAAV9-packaged Cre-expressing virus via the superficial temporal vein (STV) of PND1 mouse pups as described previously⁶². Briefly, pups were sedated by covering them with ice for 3 min. STV is visualized using a handheld transilluminator (WeeSight; Respironics, Murrysville, PA, USA), and a pair of standard reading glasses. Virus solution was prepared by 1:50 dilution of stock solution in Dulbecco's phosphate-buffered saline (PBS), also supplemented with 0.001% pluronic-F68. Virus solution (50 nL) was injected using a 100 nL Nanofil syringe attached with a 34-gauge Nanofil beveled needle (World Precision Instruments, Sarasota, FL). A correct injection was verified by noting blanching of the vein. After the injection, pups were returned to the incubator until active and then returned to their dam.

For dendritic tracing, PND 60 animals were deeply anesthetized with pentobarbital (Nembutal) and transcardially perfused with 4 % PFA/PBS (wt/vol). Extracted brains were post-fixed in 4 % PFA/PBS at 4°C for 10 h and cryoprotected in 20 % sucrose/PBS (wt/vol) at 4°C for 24 h. Brains were cut on a vibratome (500 µm thickness) collecting the somatosensory cortices. Slices were immediately submerged in Scale A2 solution in order to clear the tissue. When the tissue was transparent (at least two weeks), brain slices were mounted in Petri dishes, covered in agarose, and imaged using standard confocal microscopy. Three-dimensional image stacks were collected (x: 2048, y: 2048 pxl; step size 1 µm) using confocal microscopy equipped with water immersion objective lens (ULTRA

25x, numerical aperture 1.05, Olympus). A computer-based tracing system (Neurolucida360; MicroBrightField) was used to generate three-dimensional neuron tracings that were subsequently visualized and analyzed with NeuroExplorer (MicroBrightField). In order to select a neuron, the following criteria were strictly followed: 1) neuron was selected starting toward the middle of the stack (~150 μ m ± 30 μ m) to ensure the accurate reconstruction of an entire dendritic arbor; 2) neuron was distinct from other neurons to allow for identification of branches; 3) neuron was not truncated in some obvious way. For every reconstructed neuron, an estimate of dendritic complexity within layer 2/3 and 4 neurons was obtained with Sholl analysis. A 3D Sholl analysis was then performed in which concentric spheres of increasing radii (20 μ m increments) were layered around the cell body until branches were completely enveloped. The total length of branches, the number of dendritic intersections at each sphere, and the dendritic orders were measured ⁵⁰. Neurons were traced by an experimenter blind to genotype.

Spine Density Analysis

Because the clearing tissue method enhanced the quality of our images, spine density was determined using the same set of images previously acquired for the tracing experiment. As previously described ⁵⁰, ten to fifteen dendritic segments of somatosensory cortex L2/3 and L4 (20-120 μ m in length) at PND 60 mice were collected and considered for analysis. All measurements were performed by an experimenter blind to the experimental conditions. Pictures were visualized and elaborated with Neurolucida 360 software (MicroBrightField).

Acute slice preparation and in vitro electrophysiology

Acute thalamocortical (TC) slices (350 μ m) from 8 week old *Syngap1* WT and Het mice were cut using standard methods as previously described⁶³. Ice-cold cutting solution contained (in mM): 110 choline-Cl, 25 NaHCO₃, 1.25 NaH₂PO₄, 2.5 KCl, 0.5 CaCl₂, 5 MgCl₂, 25 Glucose, 10 Ascorbic acid, 5 Pyruvic acid (pH 7.4, ~300 mOsm). The slices were then warmed to 35°C for 40 minutes in standard artificial cerebrospinal fluid (aCSF), composed of (mM): 125 NaCl, 2.5 KCl, 24 NaHCO₃, 2 CaCl₂, 1.25 NaH₂PO₄, 2 MgSO₄, and 10 D-Glucose, and equilibrated with 95 % O₂ and 5 % CO₂ (pH 7.4, ~300 mOsm). Following this, slices were maintained in bubbled aCSF at room temperature until transferred to a submerged-type recording chamber (Warner Instruments, Hamden, CT). All experiments were performed at 32°C±2 (2-3 mL/min).

Whole-cell patch clamp experiments were conducted from visually identified L2/3 neurons using infrared DIC optics and regular spiking was confirmed in current clamp. Recordings were made using borosilicate glass pipettes (3-6 MΩ; 0.6 mm inner diameter; 1.2 mm outer diameter; Harvard Apparatus). All signals were amplified using Multiclamp 700B (Molecular Devices, Sunnyville, CA), filtered at 2 KHz, digitized (10 KHz), and stored on a personal computer for off-line analysis. Analog to digital conversion was performed using the Digidata 1440A system (Molecular Devices). Data acquisitions and analyses were performed using pClamp 10.2 software packsge (Clampex and Clampfit programs; Molecular Devices).

For current clamp and evoked excitatory postsynaptic current (eEPSC) recordings, the following internal solution was used (in mM): 130 potassium gluconate, 5 KCl, 10 HEPES, 0.25 EGTA, 10 phosphocreatine disodium, 0.5 Na-GTP and 4 Mg-ATP (pH 7.3, 285-290 mOsm). For miniature excitatory postsynaptic current (mEPSC) recordings, the following internal solution was used (in mM): 120 CsCl, 5 NaCl, 1 MgCl₂, 10 HEPES, 10 EGTA, 3 Mg-ATP and 0.3 Na-GTP (pH 7.3, 285-290 mOsm). Cells with access resistance >30 M Ω or were unstable (>20 % change) were discarded from further analysis.

In eEPSC experiments, slices were incubated with 100 µM picrotoxin. L4 was stimulated by placing a concentric bipolar stimulating electrode (FHC, 25 µm inner diameter; 125 µm outer diameter, ME, USA) in the center of a barrel. L2/3 neurons were recorded directly above the stimulus site. The stimulation intensity (0.2 ms, constant-current pulses) was regulated by a stimulus isolation unit (ISO-Flex, A.M.P.I). A minimum stimulus intensity protocol was employed (6 sweeps with 15 s ISI) in current clamp, as eEPSPs appeared to be more stable than eEPSCs recorded in voltage clamp. The stimulus intensity was gradually increased until the emergence of an eEPSP with a 30-50 % failure rate. This stimulus intensity was used in voltage-clamp mode to elicit eEPSCs from 15 sweeps (15 s ISI). eEPSCs were quantified by averaging the peak amplitudes, within a 30 ms post-stimulus window, excluding failures. Only one neuron was recorded from a single barrel column and a maximum of 2 neurons per slice. mEPSCs were recorded in TTX (1 µM), picrotoxin (100 μ M) and APV (100 μ M) at a holding potential of -75 mV. For each recording, the first 500 events were analyzed using Mini Analysis software (Synaptosoft Inc, NJ). Intrinsic properties of neurons were recorded at the resting membrane potential and measured as described²⁴.

Novel Texture Discrimination Task (NoTeDt)

Before testing for NoTeDt, mice were first habituated to an Open Field Arena. Cohorts of mice were handled for several minutes on three separate days prior to commencement of behavioral testing. On the first day of behavioral assay, mice were subjected to the open field habitation phase, during which an animal was allowed to explore a custom-made clear acrylic arena (43 cm x 43 cm x 32h cm) for 10 minutes, two daily sessions, for two days. Opaque white acrylic dividers surrounded each arena to prevent distractions from activities in adjacent boxes. Activity was monitored with two CCTV cameras (Panasonic WV-BP334) feeding into a computer equipped with Ethovision XT 11.5 for data acquisition and analyses. A white noise generator (2325-0144, San Diego Instruments) was set at 70 dB to mask external noises and provide a constant noise level. Fluorescent linear strip lights placed on each of the four walls of the behavioral room adjacent to the ceiling provided a lower lighting (100 lux) environment. Distance travelled was analyzed as an indicator of familiarization to the experimental environment.

NoTeDt was performed in the same arenas used for open field habituation. The cylindricalshaped objects were conceived to promote whiskers-like explorations. Objects were designed by optimizing criteria from previously described and validated texture-based object discrimination tasks^{64, 65}. Briefly, white polyoxymethylene objects were 15 cm tall, fixed to a 5 cm diameter base and the rough portion of the object covered 7.5 cm of the surface starting 2 cm from its base. Two different grades of roughness (T1 and T2) were chosen and tested to determine whether the objects shared similar degree of attractiveness.

After the habituation phase (see open field section for details), mice were engaged in exploring two identical objects (T1) for 10 minutes (learning phase). After 5 minutes, necessary for rearranging the arena (cleaning and replacing one of the objects T1 with the novel object T2), mice were exposed to the testing phase, engaging in the exploration of both old and new objects. The location of the novel object was pseudo-randomized and counterbalanced between groups. Time spent in exploring textures was assessed by manually scoring each trial (offline), during both learning and testing phases. Mice that did not explore the objects during the learning phase, explored only one of the two objects during the testing phase, or had a total investigation time of less than 20 s during the learning phase, were excluded from the study for lack of adequate exploratory activity. Behavioral scoring was performed by an experimenter blind to the experimental conditions.

Whisker-dependent Go/NoGo task

Syngap1 WT and Het littermates (male, housed in a reverse light-dark room) were trained one session per day (~5 days/week) to perform a head-fixed whisker-dependent Go/NoGo task³⁹. The behavioral apparatus was controlled by open-source BControl software (C. Brody, Princeton University) on a RT Linux machine⁶⁶. A custom titanium head-post was implanted onto the skull at six to ten weeks after birth. Briefly, animals were anesthetized with isoflurane (5% induction, 1.5-2% maintenance) and injected IP with rimadyl (carprofen 10 mg/kg) and enroflox (endrofloxacin 5 mg/kg). A small flap of skin was removed over the midline, and the skull scraped with a scalpel before a thin layer of cyanoacrylate glue (Vetbond, 3M) was applied. The head-post was implanted onto this layer of glue and secured with dental cement (Metabond). Animals were then singly housed and monitored for 1 week following surgery, with added environmental enrichment (running wheel). All behavioral sessions occurred during the dark phase. Weekly and throughout the experiment, all whiskers were trimmed to the base except for the C2 whisker on either side. Following recovery from head-post surgery, mice were placed on water restriction (1 ml/day) for 10 days prior to and then throughout training (food ad libitum). Water-restricted mice were monitored for health issues and weighed daily. On day six of water restriction, habituation to head-fixation commenced with increasing time spent under head-fixation, up to one hour. Mice were head-fixed in a custom-built stainless-steel body tube, inside a fully enclosed light and soundproof box, which was continuously monitored via IR videography. On the last day of habituation, video recordings (30 FPS, Basler, acA640-120um) were obtained (45 mins duration) under IR illumination to determine basic whisking properties (bouts and bout durations). Whisking was scored manually, offline (Solomon Coder) by an experimenter blinded to mouse genotypes.

Mice moved through a series of training steps. First, lick-port training (1 session lasting 10 mins) allowed mice to associate water availability via licking from a lick-port, with the C2 whisker inserted into a pipette (with ~ 2 mm at the base exposed) attached to a piezo actuator (Physik Instrumente), however, no passive whisker deflections were introduced. The metal lick-port was positioned directly in front and slightly below the mouth, within reach of

the tongue. Detection of licks was performed electronically⁶⁷ and precise water delivery (8 µl/reward) was controlled with a solenoid valve (The Lee Company, CT). Next, Step 1 training lasted for 2 sessions and represented the first-time mice were exposed to passive whisker deflections. Here, sessions consisted of 90% Go and 10% NoGo trials. For Go trials, the whisker was deflected by the piezo actuator, controlled by a linear voltage amplifier (Physik Instrumente) and a waveform generator (BK Precision), for 1.5 s with a 40 Hz sinusoidal wave (rostral to caudal, 1200 °/s). Bending of the piezo was calibrated using a laser-based displacement device (LD1610-0.5 Micro - Epsilon). For NoGo trials, a "dummy" piezo, placed just above the whisker deflecting piezo was driven with the same stimulation, but was not attached to a whisker. The response window was defined from 0.1-4 s after the start of piezo stimulation. For Go trials, hits consisted of trials where mice licked for a water reward within the response window, and misses were scored as lack of licking during this same period. For NoGo trials, a false alarm (FA) resulted when the animal licked during the response window and a correct rejection occurred when the animal withheld licking. No punishments were given for FAs and no auditory cue was presented. Inter-trial interval remained constant at 4 s, however, mice were required to withhold licking for 1.5 s before the piezo was stimulated for trials to proceed. Mice performed the task until satiated. Step 2 training was similar to Step 1, except sessions consisted of 50% Go and 50% NoGo trials (200-350 trials/session) with no more than 3 consecutive trials of the same type. The response window was shortened to 2 s. Training persisted until mice reached several performance criteria for at least 2 consecutive days. 1) Overall performance reached > 70%correct for all trials, 2) Hits > 70%, 3) FAs < 30%, and 4) discrimination index (d') > 1.1. D' was calculated in Excel (Microsoft) as d' = z(hit) - z(FA), with z scores computed using the function NORMSINV. If mice did not reach criteria by 21 days, training was halted. "Good performers" graduated to Step 3, whereby the stimulus duration was reduced to 0.5 s. Mice continued this training until the same criteria was met as in Step 2. These mice graduated to a reduced stimulation protocol. Here, sessions consisted of the same experimental setup as Step 3, except angular velocity of passive whisker deflections were reduced from 1200 °/s to a range between 300-900 °/s.

Statistics:

Data analyses were conducted in SPSS (IBM Corp, version 20), MATLAB (Mathworks, version 2015b, Natick, MA) and GraphPad Prism 7 (GraphPad Software, CA). D'Agostino-Pearson omnibus normality test was applied to determine data distributions and the appropriate parametric or non-parametric statistical test was performed accordingly. However, for experiments with small sample sizes (N<8 subjects per group), statistical tests tend to lack power to detect deviations from normality, and therefore a more subjective approach was used. For these cases where data appeared to approximate a normal distribution, we assumed normality and used parametric statistical tests (i.e. Fig. 5c, Fig. 8n, but not limited to these panels). For analysis of imaging data, the following tests were used: two-sided student's t-test was used to compare IOS amplitudes, IOS inter-barrel distance, GWI integrated F/F, GWI ratio of 4th/1st pulse to first pulse and the fraction of responding cells (except for Fig. 2d, where <u>2-way RM-ANOVA</u> was used) between genotypes; <u>2-way RM-ANOVA</u> was used to compare area measures for both relative and absolute thresholding methods for IOS and for peak responses and integrated F/F in GWI experiments;

Kolmogorov-Smirnov test was used to compare event amplitudes, event counts/2 minutes, and response probabilities, with Bonferroni-adjusted P values to control for type I errors when multiple pairwise comparisons were made. For cluster analyses of spike counts, we pooled all spikes for each genotype and treatment, then performed a two-step cluster analysis. We specified a fixed number of cluster to three, for low-, medium- and highresponsive neurons and used the Euclidean distance for computing the similarity between clusters. The number of neurons in each cluster were then separated into groups based on genotypes and treatment for further analysis. Differences in the clustered data were analyzed using Chi-square analysis. Pairwise post-hoc comparisons between clusters and groups were performed using Chi-square analysis with Bonferroni-adjusted P values to control for type I errors. For analysis of the neuronal morphology of layer 2/3, and 4 neurons, two-sided student's t-test, assessing the contribution of *total length, # of nodes* and the *spine density* were performed. For electrophysiology data, two-sided student's t-test or Mann-Whitney U test when applicable or Kolmogorov-Smirnov test was used to compare measurements between WT and Het mice. 2-way RM-ANOVA was used to compare response amplitudes of responses to each whisker deflection and the number of spikes in response to current injections. NoTeDt behavioral data are expressed as the median \pm interquartile except data showed in fig 1b and fig.1e which are expressed as the mean \pm SEM. Non-parametric Friedman and Mann-Whitney U-tests were used to analyze WT and Het performance in both learning and testing phases. Post-hoc comparisons were performed by using sign test and the false discovery rate (FDR)-controlling procedure Benjamini/Hochberg, was used to compute the adjusted level of significance. One sample t-tests were performed against a chance value of 50% to determine if the mice were able to discriminate between the novel and the old textures. For Go/NoGo behavioral data, the following tests were used: 2-way RM-ANOVA with Bonferroni's multiple comparisons post-hoc test was used to compare performance (P(lick)) of WT and Het mice, performance with reduced angular velocity of whisker stimulation, normalized weights, number of trials performed, response times, and number of total licks; one-way repeated measures ANOVA with Bonferroni's multiple comparisons post-hoc test was used to compare d' with reduced angular velocity of whisker stimulation; two-sided student's t-test or Mann-Whitney U-tests were used to compare d' between WT and Hets, mean water licks, total reward licks, mean trials/session, total trials performed, mean licks/trial (total), number of total licks and basic whisking parameters; fisher's exact test was used to compare proportion of performers vs. non-performers in WT and Het mice. Unless otherwise stated, data represents mean \pm SEM. No statistical test was used to predetermine sample sizes, however, our sample sizes are similar to those previously reported in the field⁶⁸.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1: Reduced sensory-evoked brain activity in Syngap1 SSC.

(**a-b**) Stimulus paradigm used during IOS imaging. (**c**) Example IOS signals from one animal in each genotype obtained for β and C2 whiskers. (**d-f**) Scatter plots showing reduced IOS amplitudes in β (**d**) and C2 (**e**) whisker fields, but normal inter-barrel distance (**f**) in adult Syngap1 mutants. Unpaired two-sided t-Tests: t(20)=3.76 p=0.0014 for β amplitude (n=11 WT n=11 Het mice), t(16)=3.70 p=0.001 for C2 amplitude (n=9 WT, n=9 Het mice), t(19)=1.49 p=0.15 for inter-barrel distance(n=10 WT n=11 Het mice). (**g**, **h**) Quantification of the area responding to β or C2 whisker stimulation according to absolute (**g**) or relative thresholding methods (**h**). 2-way RM-ANOVA for absolute area F_(1,36)=11.18 p=0.002 for genotype, F_(3,108)=1.74 p=0.163 for genotype*threshold; 2-way RM-ANOVA for relative area F_(1,36)=2.8 p=0.1 for genotype, F_(3,108)=2.92 p=0.037 for genotype*threshold (n=20 WT n=20 Het IOS imaging sessions from different whiskers). For **d-f**, open circles represent animal means, black lines indicate population means and error bars indicate SEMs. Data in this figure were acquired from two independent cohorts of animals that were pooled together.





(a) Experimental setup for awake, *in vivo* two-photon calcium imaging in *Syngap1* crossed with Thy-1 Gcamp6s4.3 mice, with and without Botox injection. (b-c) Representative *in vivo* two-photon microscopy images (left) and F/F traces (right) of spontaneous activity in 9 (1-9) neurons of L2/3 SSC of WT and Het *Syngap1* X Thy-1 Gcamp6s4.3 mice, without
(b) and with (c) Botox. ROI number 10 is the neuropil signal. Asterisks indicate detected calcium events (blue for WT, red for Het). (d-f) Cellular sensory properties from awake WT

and Het mice without and with Botox injection. (d) Scatter plot showing fraction of spontaneously active cells in WT and Het mice (2-way RM-ANOVA, Genotype: F(1, 14) =0.29, p = 0.60, Treatment: F(1,14) = 0.034, p = 0.86, Interaction: F(1, 14) = 0.012, p = 0.92; WT n=8 mice, Het n=8 mice). (e-f) Cumulative probability plots of F/F amplitudes (e; KS Tests: WT/Botox- vs. Het/Botox-, p= 3.38E-7; WT/Botox+ vs. Het/Botox+, p=0.0002; WT/ Botox- vs. WT/Botox+, p= 0.67; Het/Botox- vs. Het/Botox+, p= 0.078; WT: n_{Botox}=1622 neurons, $n_{Botox+} = 1827$ neurons; Het: $n_{Botox-} = 1667$ neurons, $n_{Botox+} = 1600$ neurons) and spike counts (f, KS Tests, WT/Botox- vs. Het/Botox-, p= 1.82E-7; WT/Botox- vs. WT/ Botox+, p= 7.99E-6; WT/Botox- vs. Het/Botox+, p= 2.87E-8; Het/Botox- vs. Het/Botox+, p=0.99; WT/Botox+ vs. Het/Botox+, p=0.24). (g) Whisking behavior in head-fixed Syngap1 mice (Fraction of time whisking: Unpaired t-test t(12)=7.493, p= 7.0E-6; WT n=7 mice, Het n=7 mice. Bout frequency: Unpaired t-test t(12)=6.298, p= 4.0E-5; WT n=7 mice, Het n=7 mice. Mean bout duration: Unpaired t-test t(12)=0.8714, p=0.40; WT n=7 mice, Het n=7 mice). Data were pooled from two independent cohorts of animals and thus obtained from 2169 neurons in 56 imaging planes from 8 WT mice; 1971 neurons in 55 imaging planes from 8 Het mice. In scatter plots, open circles are animal means, black lines indicate population means, and error bars indicate SEMs. ***P<0.001, ****P<0.0001 for cumulative probability plots. All statistical tests were two-sided.

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Figure 3: Reduced whisker responsiveness of SSC neurons in behaving *Syngap1* mice. Representative *in vivo* two-photon microscopy images (**a**) and representative F/F traces (**b**) of 9 (1-9) neurons in WT and Het (Botox-) mice in response to 5 passive whisker deflections at 5 Hz. Gray vertical lines indicate the timing of whisker stimuli. Green asterisks indicate calcium events within the response detection window. Red asterisks show spontaneous calcium events. ROI number 10 is the neuropil signal. (**c-h**) Cellular sensory properties from awake animals in response to 5 pulses at 5 Hz (**c-e**) and 60 pulses at 40 Hz whisker stimulation (**f-h**). (**c**) Scatter plot showing fraction of responding cells in WT and Het mice

(Unpaired t-test t(14)=2.48, p=0.026; WT n=8 mice, Het n=8 mice). (d-e) Cumulative probability and scatter plots (inserts) of F/F amplitudes (d, KS Test, p=0.13; WT: n=406 neurons, Het: n=330 neurons) and response probabilities (e, KS Test, p=0.23) in responding neurons. (f) Scatter plot depicting fraction of responding cells in WT and Het mice (Unpaired t-test t(14)=3.07, p=0.0084; WT n=8 mice, Het n=8 mice). (g, h) Cumulative probability and scatter plots (inserts) of F/F amplitudes (g, KS Test, p=0.00050; WT: n=467 neurons, Het: n=368 neurons) and response probabilities (h, KS Test, p=0.0059) in responding neurons. (i-n) Cellular sensory properties from awake animals following Botox injection in response to 5 pulses at 5 Hz (i-k) and 60 pulses at 40 Hz whisker stimulation (l**n**). (i) Scatter plot depicting fraction of responding cells in WT and Het mice (Unpaired ttest t(14)=0.35, p=0.73; WT n=8 mice, Het n=8 mice). (j, k) Cumulative probability and scatter plots (inserts) of F/F amplitudes (j, KS test, p=0.32; WT: n=413 neurons, Het: n=416 neurons) and response probabilities (k, KS Test, p=0.0061) in responding neurons. (l) Scatter plot showing fraction of responding cells in WT and Het mice (Unpaired t-test t(14)=0.11, p=0.92; WT n=8 mice, Het n=8 mice). (m, n) Cumulative probability and scatter plots (inserts) of F/F amplitudes (m, KS Test, p=0.39; WT: n=481 neurons, Het: n=445 neurons) and response probabilities (n, KS Test, p=0.010) in responding neurons. (c-h) Data obtained from 1921 neurons in 54 imaging planes from 8 WT mice; 2044 neurons in 54 imaging planes from 8 Het mice. (c-n) Data was pooled from two independent cohorts of animals and thus obtained from 2169 neurons in 56 imaging planes from 8 WT mice; 1971 neurons in 55 imaging planes from 8 Het mice. In scatter plots, open circles are animal means, closed circles are individual cells, black lines indicate population means, and error bars indicate SEMs. All statistical tests were two-sided.



Figure 4: Reduced sensory responsiveness of L2/3 SSC neurons in *Syngap1* mice is cortex-specific.

(a) Representative *in vivo* two-photon microscopy images of L2/3 SSC of WT and Het Emx1-Cre X *Syngap1* cKO mice. (b) Representative F/F traces of 9 (1-9) neurons in WT and Het mice in response to 5 passive whisker deflections at 5 Hz (from **a**). Gray vertical lines indicate the timing of whisker stimulus. Green asterisks indicate calcium events within the response detection window. Red asterisks show spontaneous calcium events. ROI number 10 is the neuropil signal. (**c-e**) Cellular sensory properties pooled from two

independent cohorts of Emx1-Cre X Syngap1 cKO mice in response to 5 pulses at 5 Hz whisker stimulations under anesthesia. (c) Scatter plot showing fraction of responding cells in WT and Het mice (Unpaired t-test t(14)=0.85, p=0.41; WT n=8 mice, Het n=8 mice). (d, e) Cumulative probability and scatter plots (inserts) of F/F amplitudes (d, KS Test, p=0.016; WT: n=327 neurons, Het: n=306 neurons) and response probabilities (e, KS Test, p= 8.2E-5) in responding neurons. Data obtained from 1671 neurons in 55 imaging planes from 8 WT mice; 1877 neurons in 58 imaging planes from 8 Het mice. (f-h) Cellular sensory properties pooled from two independent cohorts of Cux2-CreERT2 X Syngap1 cKO mice in response to 5 pulses at 5 Hz whisker stimulations under anesthesia. (f) Scatter plot showing fraction of responding cells in WT and Het mice (Unpaired t-test t(14)=0.56, p=0.58; WT n=8 mice, Het n=8 mice). (g, h) Cumulative probability and scatter plots (inserts) of F/F amplitudes (g, KS Test, p=0.0002; WT: n=435 neurons, Het: n=445 neurons) and response probabilities (h, KS Test, p=0.67) in responding neurons. Data obtained from 2015 neurons in 57 imaging planes from 8 WT mice; 1901 neurons in 56 imaging planes from 8 Het mice. (i-k) Cellular sensory properties pooled from two independent cohorts of Rpb4-Cre X Syngap1 cKO mice in response to 5 pulses at 5 Hz whisker stimulations under anesthesia. (i) Scatter plot showing fraction of responding cells in WT and Het mice (Unpaired t-test t(13)=0.64, p=0.53; WT n=8 mice, Het n=7 mice). Cumulative probability and scatter plots (inserts) of F/F amplitudes (j, KS Test, p=0.74; WT: n=340 neurons, Het: n=317 neurons) and response probabilities (k, KS Test, p=0.26) in responding neurons. Data obtained from 1684 neurons in 56 imaging planes from 8 WT mice; 1411 neurons in 46 imaging planes from 7 Het mice. In scatter plots, open circles are animal means, closed circles are individual cells, black lines indicate population means, and error bars indicate SEMs. All statistical tests were two-sided.

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Figure 5: Reduced sensory responsiveness in both excitatory and inhibitory neuronal populations in L2/3 SSC of *Syngap1* mutants.

(a) Representative *in vivo* two-photon microscopy image of L2/3 SSC of a Gad2-T2A-NLS-MCherryX*Syngap1* WT mouse expressing GCaMP6s (green) and Mcherry (red). White arrows indicate MCherry positive (inhibitory) neurons expressing GCaMP6s. (b) Representative F/F traces of 9 (1-9) excitatory neurons in WT and Het mice in response to 5 passive whisker deflections at 5 Hz. Gray vertical lines indicate the timing of whisker stimuli. Green asterisks indicate calcium events within the response detection window. Red asterisks show spontaneous calcium events. ROI number 10 is the neuropil signal. (c-h) Cellular sensory properties of excitatory (c-e) and inhibitory (f-h) neurons. (c) Scatter plot showing fraction of excitatory neurons responsive to whisker stimulation (Unpaired t-test t(11)=1.891, p=0.0853; WT n=7 mice, Het n=6 mice). (d, e) Cumulative probability and scatter plots (inserts) of F/F amplitudes (d, KS Test, p=0.0011) in responding excitatory neurons. (f) Scatter plot showing fraction of inhibitory neurons responsive to whisker stimulation (Unpaired t-test t(11)=1.164, p=0.2691; WT n=7 mice, Het n=6 mice). (g, h) Cumulative probability and scatter plots (inserts) of F/F amplitudes (inserts) of F/F amplitudes (g, KS Test, p=0.9848;

WT n=49 neurons, Het n=20 neurons) and response probabilities (**h**, KS Test, p=0.0176) in responding inhibitory neurons. Data in this figure was pooled from two independent cohorts of animals and thus obtained from 850 excitatory and 240 inhibitory cells in 48 imaging planes from 7 WT mice; 825 excitatory and 193 inhibitory cells in 45 imaging planes from 6 Het mice. Open circles are animal means, closed circles are individual cells, black lines indicate population means, and error bars indicate SEMs. All statistical tests were two-sided.



Figure 6: *In vivo* patch clamp reveals that L2/3 SSC neurons in *Syngap1* mutants have reduced sensory-evoked synaptic input.

(a) Representative *in vivo* traces for whole-cell patch clamp experiments in response to passive whisker stimulations. (b) Scatter plot showing the overall response peaks (t-test: t(15)=2.59, p=0.021; n=8 for WT n=9 for Het. (c) Individual response amplitudes (2-way RM-ANOVA, F(1,14)=4.82, p=0.045 for genotype effect, F(4,56)=0.72, p=0.58 for genotype and stimulus interaction) in response to whisker stimulation in L2/3 neurons from *in vivo* patch clamp recordings. For **b**, open circles represent animal means, black lines indicate population means and error bars indicate SEMs. For **c** closed circles represent population means and error bars indicate SEMs. Data obtained from two cohorts of *Syngap1* animals. All statistical tests were two-sided.



Figure 7: *Syngap1* heterozygosity degrades synaptic connectivity and reduces intrinsic excitability of upper layer SSC neurons.

(**a**, **c**) Representative 3D reconstruction of L4 (**a**, left) and L2/3 (**c**, left) of SSC excitatory neurons depicting dendritic complexity and scatter plot (right) showing the total length and # of nodes by using Sholl analysis. (**a**) Total length: (WT = 5 mice, Het = 5 mice), unpaired t-Test, t(8)=4.002, p = 0.0030; # of Nodes: unpaired t-Test, t(8)=3.017, p = 0.0166. (**c**) Total length: (WT = 5 mice, Het = 6 mice), unpaired t-Test, t(9)=3.7713, p = 0.0044; # of Nodes: unpaired t-Test, t(9)=3.7090, p = 0.0048, from a single cohort of animals. (**b**, **d**) Examples of

L4 (b, left) and L2/3 (d, left) apical dendrites and scatter plots (right) depicting the density of dendritic spines. (b) Layer 4 - spine density: (WT = 5 mice, Het = 5 mice), unpaired t-Test, t(8)=4.059, p = 0.0036. (d) Layer 2/3 - spine density: (WT = 5 mice, Het = 5 mice), unpaired t-Test, t(8)=7.80, p = 5.2E-5, from a single cohort of animals. (e) Representative traces depicting $L^{2/3}$ excitatory neuron mEPSCs from acute WT and Het TC slices. (f, g) Cumulative probability and scatter plots (inserts) of mEPSC amplitudes (f, Kolmogorov-Smirnov test p = 2.2E-16) and mEPSC IEI (g, Kolmogorov-Smirnov test p = 2.1E-16). (f, g) Data was acquired from a single cohort of animals with n=7986 mEPSC events from 16 neurons in 4 WT and n=6765 mEPSC events from 16 neurons in 4 Het mice. (h) Cartoon depicting experimental setup for investigating feed-forward excitation in L2/3 excitatory neurons from L4. (i) Representative traces depicting L2/3 excitatory neuron eEPSCs from acute WT and Het TC slices. (j) Scatter plot of eEPSC amplitudes in L2/3 following stimulation of L4 (Mann-Whitney test, U=14.00, p=2.5E-5; Data obtained from a single cohort of animals; WT n=14 neurons from 4 mice, Het n=14 neurons from 4 mice). (k) Representative current-clamp traces from L2/3 excitatory neurons from acute WT and Het TC slices and (I) graph depicting a decrease in the number of spikes (2-way RM-ANOVA, F(1,46)=5.51, p=0.023 for genotype effect F(9,414)=5.46, 9.0E-9 for genotype and stimulus interaction, n=22 neurons from 5 WT mice, n=26 neurons from 6 Het mice) in response to current injections. (m) Scatter plot showing increased rheobase (Student's t-test: t(44)=3.50, p = 0.0010) in the same set of neurons as in **l**. Data was obtained from a single cohort of animals. For morphology data, open circles are animal means, closed circles are individual cells, black lines indicate population means, error bars indicate SEMs, colored triangles represent spines (blue = WT, red = Het) and white triangles represent filopodia. For f, g, j, m, open circles are individual cells, black lines indicate population means, and error bars indicate SEMs. For l, circles represent population means and error bars indicate SEMs. All statistical tests were two-sided.



Figure 8: Impaired texture discrimination and whisker-dependent Go/NoGo task performance in *Syngap1* mice.

(a) Cartoons depicting different texture roughness of the objects used in Novel Texture Discrimination Task and relative protocol. (b) Scatter plot showing no preference in exploring T1 or T2. (T1 vs T2: n = 12 mice; unpaired t-Test, t(22) = 0.0016, p = 0.9986. (c-d) Box plots (solid line represents median, box represents interquartile range and whiskers represent maximum and minimum values) depicting time spent exploring identical textured objects during the learning phase and time spent exploring the novel (T2) and the old (T1)

object for WT (c) and Het groups (d) [(WT mice Friedman test: χ^2 (n = 18 mice, df = 3) = 28.87, exact sign. p = 1.4206 E-7; (HET mice, Friedman test: χ^2 (n = 15 mice, df = 3) = 15.24, exact sign. p = 0.0016). Pairwise comparisons: (WT(learning) vs WT(testing), Sign test: n = 18 mice, Z = 4.007 p = 0.00006; HET(learning) vs HET(testing), Sign test: n = 15 mice, Z = 2.065, p = 0.04, ns). Statistical significance was accepted at the p < 0.03125. WT(learning) vs HET(learning), Mann-Whitney U test: Z = -2.567 p = 0.01; WT(testing) vs HET(testing), Mann-Whitney U test: Z = -2.821 p = 0.0048). (e) Scatter plot showing Exploration Index for animals in c and d: WT mice, paired t-Test: t(17) = 4.707, p = 0.0002; HET mice paired t-Test: t(14) = 1.641, p = 0.123; One sample test: WT(learning), t(17) =1.555, p = 0.138; WT(testing), t(17) = 8.579, p = 1.39E-7; HET(learning), t(14) = -2.164, p = 0.048, ns; HET(testing), t(14) = -2.415, p = 0.03, ns. (f) Cartoon representation of Go/ NoGo setup. Water-restricted, head-fixed mice were rewarded with water for licking a lickport in response to a passive whisker (C2) deflection. (g) Detection task trial structure for Step 2 training. Go trials are identical to NoGo trials, except for the passive whisker deflection. Note, NoGo trials include activation of a "dummy piezo" not attached to any whisker to control for noise/vibration associated with piezo activation. (h) Step 2 training learning curve for WT mice showing the probability of licking (P(lick)) on Go (black, hit) or NoGo (blue, FA) trials (n=7 mice; 2-way RM-ANOVA with Bonferroni's multiple comparison, Trial type: $F_{(1,6)}=67.19 \text{ p}=0.0002$; Session: $F_{(15,90)}=0.4827 \text{ p}=0.9437$; Trial type*Session interaction $F_{(15,90)}=5.86 \text{ p}=2.9\text{E-8}$. (i, j) Reductions in angular velocity of whisker deflections impairs the ability of "Good Performing" WT mice to discriminate between trial types (i: n=6 mice; 2-way RM-ANOVA with Bonferroni's multiple comparison, Trial type: $F_{(1.5)}=471.1 \text{ p}=3.9\text{E-6}$; Velocity: $F_{(3.15)}=1.469 \text{ p}=0.263$; Trial type*Velocity interaction: $F_{(3,15)}=30.12 \text{ p}=1.4\text{E-6}$) and results in a reduced discrimination index (j: n=6 mice; RM-ANOVA with Bonferroni's multiple comparison, $F_{(3,15)}=24.52$ p= 4.9E-6). (k) Proportion of mice to learn (performers, P vs. non-performers, Non-P) the task (WT n=7, Het n=7: Fisher's Exact Test: p=0.0047). (I) Step 2 training learning curve for Het mice (n=7 mice; 2-way RM-ANOVA with Bonferroni's multiple comparison, Trial type: $F_{(1.6)}$ =8.44 p=0.027; Session: $F_{(15.90)}$ =2.416 p=0.0054; Trial type*Session interaction $F_{(15,90)}=0.8852$ p=0.5825). (m) Learning curves depicting the fraction of total trials correct in Step 2 training (WT n=7, Het n=7; 2-way RM-ANOVA with Bonferroni's multiple comparison, Genotype: $F_{(1,12)}=10.13$ p=0.0079; Session: $F_{(15,180)}=3.665$ p= 1.4E-5; Genotype*Session interaction: $F_{(15,180)}=4.398 p= 5.2E-7$). (n) Scatter plot showing the discrimination index at the completion of Step 2 training (WT n=7, Het n=7; Unpaired t-test, t(12) = 4.281, p=0.0011). Data for both Novel Texture Discrimination and Go/NoGo Tasks were obtained from two independent cohorts of animals. Open circles are individual animals, closed circles and solid black horizontal lines indicate population means and error bars or shaded area represent the SEMs, except for boxplots in c and d which are described above. **P*<0.05, ***P*<0.01, ****P*<0.001, *****P*<0.0001 for post-hoc multiple comparisons. (h, i, l) Solid black and blue dashed lines indicate performance criteria for hit and FAs, respectively. (m) Solid black line indicates performance criteria for total trials correct. All statistical tests were two-sided.