Contribution of interneuron subtype-specific GABAergic signalling to emergent sensory processing in somatosensory whisker barrel cortex in mouse.

Liad J. Baruchin¹, Michael M. Kohl¹² & Simon J.B Butt¹*

¹. Department of Physiology, Anatomy & Genetics, Sherrington Building, Parks Road, 10 Oxford, OX1 3PT, UK
². Institute of Neuroscience and Psychology, University of Glasgow, Glasgow, G12 8QQ, UK

* Corresponding Author: simon.butt@dpag.ox.ac.uk
Abstract

Mammalian neocortex is important for conscious processing of sensory information. Fundamental to this function is balanced glutamatergic and GABAergic signalling. Yet little is known about how this interaction arises in the developing forebrain despite increasing insight into early GABAergic interneuron (IN) circuits. To further study this, we assessed the contribution of specific INs to the development of sensory processing in the mouse whisker barrel cortex. Specifically we explored the role of INs in speed coding and sensory adaptation. In wild-type animals, both speed processing and adaptation were present as early as the layer 4 critical period of plasticity, and showed refinement over the period leading to active whisking onset. We then conditionally silenced action-potential-dependent GABA release in either somatostatin (SST) or vasoactive intestinal peptide (VIP) INs. These genetic manipulations influenced both spontaneous and sensory-evoked activity in an age and layer-dependent manner. Silencing SST+ INs reduced early spontaneous activity and abolished facilitation in sensory adaptation observed in control pups. In contrast, VIP+ IN silencing had an effect towards the onset of active whisking. Silencing either IN subtype had no effect on speed coding. Our results reveal how these IN subtypes differentially contribute to early sensory processing over the first few postnatal weeks.
Introduction

The mammalian neocortex is a higher order area of the central nervous system responsible for processing of sensory information and initiation of voluntary behaviour. Essential to this role are local circuits comprised of glutamatergic pyramidal cells and locally-projecting GABAergic interneurons (INs). These two populations integrate incoming sensory information – relayed via the thalamus – to generate percepts, which subsequently elicit an appropriate behavioural response through efferent pyramidal cells. Much of our understanding of the processes underpinning such computations – at the cellular and circuit level, has been derived from fundamental research in animal models. One such model is the mouse somatosensory barrel field (S1BF): the area of the neocortex responsible for processing incoming tactile sensory information arising from the whiskers (Petersen, 2007). Investigations performed in mature rodents have revealed that neurons in the columnar and layered structure of S1BF can derive various stimulus properties from incoming signals, such as location, speed, texture and relative novelty (Guić-Robles, Valdivieso, & Guajardo, 1989; Musall, Haiss, Weber, & von der Behrens, 2017; R. S. Petersen, Panzeri, & Diamond, 2002). A body of evidence has identified that GABAergic signalling (Rudy et al. 2011; Yu et al. 2019; Muñoz et al. 2017)) is required for such sensory processing in the mature neocortex (Ayzenshtat, Karnani, Jackson, & Yuste, 2016; Kolasinski et al., 2017; Natan et al., 2015; Wood, Blackwell, & Geffen, 2017). However, the contribution of GABAergic INs to nascent processing in the developing brain is still unknown.

In the developing neocortex there is an additional challenge, namely to balance emergent sensory processing and formative behavioural output with the need to integrate and establish circuit function. This challenge is met across primary sensory areas – including S1BF (Erzurumlu & Gaspar, 2012) – by gradual shifts in synaptic connectivity and plasticity, during which time there are changes in the nature of cortical activity. This includes oscillations not present in the adult neocortex such as the intermittent spontaneous spindle bursts (SB) (Rustom Khazipov et al., 2004; Marat Minlebaev, Ben-Ari, & Khazipov, 2007). To date, our understanding of which neuronal subtypes that contribute to these formative activity patterns and emergent perception is limited (Hangaru-Opatz et al., 2021). What is clear is that GABAergic interneuron diversity has a role to play in constraining the influence of early sensory input and sculpting early circuits (Butt, Stacey, Teramoto, & Vagnoni, 2017; Modol et al., 2020). Of the three main classes of interneuron (Rudy et al., 2011), parvalbumin (PV+)-
expressing INs have been shown to play an important role in the closure of periods of plasticity (Hensch, 2005; McRae, Rocco, Kelly, Brumberg, & Matthews, 2007; Nowicka, Soulsby, Skangiel-Kramska, & Glazewski, 2009) and the onset of fast adult-like signalling (Doischer et al., 2008). In contrast, recent evidence has identified that one of the other prominent IN classes – defined by expression of the peptide somatostatin (SST+), contributes to early neurodevelopment events including synaptogenesis, sensory innervation and neuronal maturation (Marques-Smith et al., 2016; Oh, Lutzu, Castillo, & Kwon, 2016; Tuncdemir et al., 2016). The third main class are defined by expression of the ionotropic serotonin receptor, 5-HT3AR (Rudy et al., 2011). These are born late in embryonic development (S. Butt et al., 2005; Miyoshi et al., 2015) and as such are thought to contribute to circuit refinement towards the onset of active sensory perception (Hanganu-Opatz et al., 2021). That said, one major subtype of 5-HT3AR IN - the genetically-tractable vasoactive intestinal peptide-positive (VIP+) INs, have recently been shown to influence early circuits via their interaction with pyramidal cells and SST+ INs (Batista-Brito et al., 2017; Marques-Smith et al., 2016; Tuncdemir et al., 2016; Vagnoni et al., 2020). Based on this understanding, we hypothesised that both SST+ and VIP+ INs contribute to emergent sensory processing through postnatal life; a role that we can test using conditional silencing of neurotransmitter release via deletion of the SNARE complex protein Snap25 (Marques-Smith et al., 2016; Washbourne et al., 2002).

To assess the role that SST+ and VIP+ INs play in early cortical sensory computations, we recorded spontaneous activity and sensory-evoked responses from S1BF in vivo through the layer (L)4 critical period of plasticity (CP) up to, and including, the onset of active whisking (AW). We found that conditionally silencing SST+ IN signalling led to a reduction in spontaneous SBs during the CP in line with delayed thalamic innervation (Marques-Smith et al., 2016), whereas silencing VIP+ INs had no effect at this early age. At the onset of AW, silencing either IN subtype resulted in increased spike activity across the entire cortical column. In terms of sensory integration, we favoured multi-whisker as opposed to a single-whisker stimulation as this best captures the natural stimulus at early ages (Carvell & Simons, 1990; Kleinfeld, Ahissar, & Diamond, 2006). Beyond assessment of the simple sensory-evoked responses we also focused on speed coding and adaptation. These two processes, that have been previously studied around the onset of active sensation (van der Bourg et al., 2016), underlie more complex perceptual processing in the
mature cortex (Allitt, Alwis, & Rajan, 2017; Arabzadeh, Petersen, & Diamond, 2003; Maravall, Petersen, Fairhall, Arabzadeh, & Diamond, 2007; Ollerenshaw, Zheng, Millard, Wang, & Stanley, 2014); perceptual processing that most likely requires temporal and spatial recruitment of diverse interneuron subtypes. We found that in wild-type animals speed was encoded in a consistent manner from the earliest time point tested. However, adaptation in the sensory response varied in profile over development. Silencing of GABAergic signalling in our two IN subtypes did not affect speed processing per se, but did result in altered sensory-evoked responses in an age and layer specific manner. This confirms differing roles for SST+ and VIP+ INs in emergent sensory processing in the developing somatosensory cortex.
Methods

Mouse Lines

Animal experiments were approved by the University of Oxford local ethical review committee and conducted in accordance with Home Office project (30/3052; P861F9BB7) licenses under the UK Animals (Scientific Procedures) 1986 Act. The following mouse lines, maintained on a mixed (C57B15/J || CD1) backgrounds were used: a conditional floxed-Snap25 [Snap25<tm3mcw>] line and the VIP-ires-Cre [Vip<tm1(cre)Zjh>] and SST-ires-Cre [Sst<tm2.1(cre)Zjh>]. SST-ires-CreHOMO;Snap25C/+ or VIP-ires-CreHOMO;Snap25C/+ were crossed with Snap25C/C mice to generate offspring with either functional (Snap25C+) or silenced (Snap25C/C) SST+ or VIP+ INs respectively. All experiments were performed blind to the genotype, which was ascertained by PCR following completion of the data analysis.

Surgical and recording procedures

Animals were anaesthetised with urethane (U2500; Sigma Ltd., UK) with a dose of 0.5-1g/kg. Depth of anaesthesia was verified by absence of reflexes and the animals’ breathing and heart rate were constantly monitored throughout the recording procedure thereafter. The animal was fixed to a stereotaxic frame (51600; Stoelting; UK) with a mouse adaptor (51615; Stoelting). Contralateral whiskers were fitted into a cannula attached to a piezo electric unit (Thor labs; PB4VB2W), connected to a piezoelectric amplifier (E-650 Piezo Amplifier; PI; Germany). The skull was then exposed and in animals younger than P10 was strengthened by applying a thin layer of cyanoacrylic glue (Loctite). Barrel cortex coordinates were identified using a neonatal brain atlas (Paxinos, Halliday, Watson, & Mustafa, 2020), and a small craniotomy was made with a surgical drill (Volvere i7, NSK Gx35EM-B OBJ30013 and NSK VR-RB OBJ10007). A silicon probe (Neuronexus A1x32-Poly2-5mm-50s-177-A32) covered in Dil solution (1,1'-dioctadecyl-3,3',3'-tetramethyl indocarbocyanine; Invitrogen; UK) was implanted by lowering it slowly into the brain. After a minimum of 30 minutes post-implantation, a baseline period of 20 minutes was recorded, after which the experimental protocols were conducted.

Experimental procedures

Whisking speed manipulation. In each trial, a single whisker deflection was delivered at varying velocities with rectified sine waves with a width equivalents of 5, 10 20, 40
and 80 Hz. The interval between deflections was 30 seconds. The different speeds were given in consecutive blocks of 20 trials, with a different order for each mouse.

**Paired-pulse procedure.** In each trial two consecutive whisker deflections of 80 Hz/288 deg/ms were delivered with a varying inter-stimulus interval of 100, 250, 500, 1000 and 1500 ms. The inter-trial interval was 30 second, and each ISI was delivered in consecutive blocks of 20 trials, with different order for each mouse.

**Current-Source Density analysis and Layer Localisation**

The current-source density (CSD) maps were derived using a previously published method (Nicholson & Freeman, 1975) with a correction for the topmost and bottommost electrodes suggested by Vaknin (Vaknin, DiScenna, & Teyler, 1988). The estimated CSD, \( C \) at depth \( z \) is described as:

\[
C(z) \frac{\phi(z + h) - 2\phi(z) - \phi(z - h)}{h^2}
\]

Where \( \phi \) is the potential at a specific depth and \( h \) is the vertical spacing between the electrodes. Each pair of contacts was averaged when using the procedure to increase the signal to noise ratio. To reduce spatial noise further, we applied the three-point hamming filter (Rappelsberger, Pockberger, & Petsche, 1981):

\[
\phi_{filt}(z) = 0.23\phi(z + h) + 0.54\phi(z) + 0.23\phi(z - h)
\]

The shortest latency, large amplitude sink was classified as the granular layer, the contacts above as the supragranular layers, and those below it as the infragranular layers (M. Minlebaev, Colonnese, Tsintsadze, Sirota, & Khazipov, 2011; Nicholson & Freeman, 1975). In each layer, only contacts that showed consistent activity and a 50 Hz noise below 2 standard deviations (SDs) of the power spectrum curve were chosen for further analysis, while the same number of chosen contacts was used for analysis for each of the layers.
Data Analysis

Data analysis was performed post hoc in Matlab (Matlab 2019b).

Baseline activity: in order to identify spindle burst (SB) activity, we first filtered the signal between 5 and 35 Hz and then isolated periods where the signal exceeded 2 SD of the average signal. Events with a duration of less than 100 ms and/or less than 3 cycles were discarded. The frequency of the SBs was calculated as the sum of troughs divided by the duration of the event. The baseline firing rate was taken by calculating using a running average of 500 ms and then averaging the windows together to get the gross-average.

Spike Sorting: spike sorting was performed using Kilosort2 (Pachitariu, Steinmetz, Kadir, Carandini, & Kenneth D., 2016; github.com/cortex-lab/KiloSort). After the automatic classification of spikes into units, manual verification was performed using phy (github.com/kwikteam/phy). These spikes were then combined into a multi-unit signal for each layer.

Sensory-evoked response (SER): for both MUA and LFP, the evoked sensory response was derived by averaging across trials, and then finding the peak of the resulting deflection. This was defined as the response amplitude (mV or Hz accordingly), and the time from the onset of whisker deflection to this point was defined as the peak latency (ms). This procedure was repeated for each of the identified layers. To correct for differences in baseline, a baseline subtraction was performed using a baseline of 100 ms before the whisker deflection.

Paired-pulse ratio (PPR): the PPR was calculated by dividing the peak amplitude of the second response by the peak amplitude of the first response.

Histology

At the end of the experiments, following administration of terminal anaesthesia, the brains were dissected and immersed in 4% paraformaldehyde (PFA; Alfa Aesar) in phosphate-buffered solution (PBS, Sigma) for 2 days. The brain was washed three times in a PBS solution, and cut into 80µm thick coronal slices using a vibratome (Leica VT1000S). To assist barrel localization, slices were counterstained with 4',6-diamidine-2' phenylindole 502 dihydrochloride (DAPI, D3571, Molecular Probes; dilution 1:1000) for 5 minutes followed by 2 minutes wash in PBS. Slices were mounted on a slide and imaged using either widefield fluorescent or confocal microscopy (LSM710; Zeiss) to verify the location of the electrode.
Statistical Analysis

Statistical analysis was performed using Prism (Graphpad). Normality of the data was tested using the Shapiro-Wilk test. Differences in populations with normal distributions were tested using Student’s t test or one-way ANOVA. In cases where normality assumptions were violated, Mann-Whitney (M-W), Kruskal-Wallis (K-W), and Wilcoxon tests were used. Bonferroni correction (BfC) and Dunn test (Dunn) were applied for multiple comparisons as appropriate. Alpha levels of $p \leq 0.05$ were considered significant. Data are presented as the mean ± standard error of the mean (SEM).
Results

The Development of Sensory-Evoked Responses in mouse S1BF

We performed multi-electrode *in vivo* electrophysiology to record both local field potential (LFP) and multi-unit activity (MUA) across the depth of S1BF in response to multi-whisker stimulation across early development. We divided our analysis into three developmental time windows: postnatal day (P)5-P8 (equivalent to the layer 4 critical period for plasticity; CP), the next few days prior to active perception (P9-P11; pre-active whisking; pre-AW) and a time window covering the onset of active perception (P12-P16; active whisking; AW); windows that reflect our existing knowledge of underlying circuitry from *in vitro* studies (Anastasiades & Butt, 2012; Anastasiades et al., 2016; Vagnoni et al., 2020). Our initial analysis focused on granular layer 4 (G) as this layer showed robust LFP and MUA across all three developmental time points (Figure 1A,B).
Figure 1. Sensory evoked response (SER) amplitude increases through development. A. CSD and MUA plots across the depth of the cortex after a single multi-whisker deflection (indicated by vertical white dashed line), during CP (P5-P8), pre-AW (P9-P13) and AW (P14-P21); SG, supragranular; G, granular; IG, infragranular layers. B. Granular layer LFP responses in the aforementioned time periods after a 5Hz or 80 Hz whisker deflection (onset indicated by vertical grey bar). C. Averages of the LFP response latencies for the different deflection speeds across the developmental time periods. There was a significant effect for both age (F (2, 188) = 22.19, p < 0.01) and speed (F (4, 188) = 5.801, P <0.01). D. Averages of the LFP response amplitudes for the different deflection speeds across the developmental time periods. There was a significant effect for age (F (2, 188) = 75.92, p < 0.01) and speed (F (4, 188) = 7.135, p<0.01). E. Typical granular layer MUA response in the aforementioned time periods after a 5Hz or 80 Hz whisker deflection. F. Average MUA response latencies across the developmental time periods. There was a significant change with speed (F(4,159) = 25.27, p<0.01) and with age (F(2,159) =63.59, p<0.01). G. Averages of the LFP response peak firing rates across the developmental time periods. There was a significant change with speed (F(4,159) = 17.51, p<0.01) and with age (F(2,159) =11.16, p<0.01 ). LFP: CP: N=11, pre-AW: N = 9, AW: N=21, MUA: CP: N = 8, pre-AW: N = 6, AW: N = 21. Brackets: p<0.05 difference in an ANOVA multiple comparison.

We found that the amplitude and latency of the LFP evoked response scaled with deflection speed starting as early as CP (Figure 1B-D), indicative of differentially encoding of whisking speed prior to active perception (AW). The decrease in latency (Figure 1B,C) and increase in amplitude (Figure 1B,D; p<0.01) of the LFP across developmental time windows suggests that an increase in thalamic synaptic input – most prominent between pre-AW and AW time windows – could be a major influence on the observed sensory responses (Malina, Mohar, Rappaport, & Lampl, 2016). To further examine any cortical contribution to early encoding of speed we further examined MUA responses to different deflection speeds (Figure 1E). Similar to the LFP, the response latency and amplitude were sensitive to speed within developmental time windows (Figure 1E-G). However, the frequency of the sensory response reached a maximum during pre-AW (p<0.01). The LFP in granular layer is a sum of both thalamic synaptic input and intra-cortical activity (Malina et al., 2016), this suggests that maturing intra-cortical inhibition (Daw, Ashby, & Isaac, 2007) counterbalances the increasing thalamic signalling.
Figure 2. Adaptation pattern changes through development. A. CSD and MUA plots showing the sensory response across the depth of the cortex after two deflections with a 0.25s ISI, during CP, pre-AW, and AW. B. Typical granular layer LFP responses during the different time-periods for two deflections with ISIs of 0.1s, 0.50s, and 1.50s. C. Average PPR of the LFP response in the granular layer during CP, pre-AW, AW for ISIs of 0.10s, 0.25s, 0.50s, 1.00s, and 1.50s. There was a significant Age X ISI interaction (F (8, 307) = 3.272, p<0.001; CP: N = 22, pre-AW: N = 25, AW: N = 20). D. Typical granular layer MUA responses during the different time-periods for 2 deflections with ISIs of 0.1s, 0.50s, and 1.50s. E. Average PPR of the MUA response in the granular layer during CP, pre-AW, AW for ISIs of 0.10s, 0.25s, 0.50s, 1.00s, and 1.50s. There was a significant age X ISI interaction (F (8, 198) = 3.876, p<0.001; CP: N = 12, pre-AW: N = 16, AW: N = 18). Green bars show deflection times. * p<0.05 in a simple multiple comparison after an ANOVA.

To study the development of adaptation we used a paired-pulse paradigm – two identical stimuli presented in sequence with varying inter-stimulus intervals (ISIs),...
similar to previous research (van der Bourg et al., 2019, Zehendner et al., 2013), however, over a larger range of developmental time points (CP to AW). We recorded LFP and MUA (Figure 2A) in response to a paired-pulse paradigm of varying ISI (between 0.1 and 1.5s), focusing our analysis on granular layer 4 (Figure 2B-E). Prior to AW, we observed failure of the second response at the shortest ISI test (0.1s) in 12 of 22 animals during CP and 13 out of 25 during pre-AW. No such failure was observed during AW. In the older animals, analysis of LFP (Figure 2B) showed a pattern of depression that becomes weaker with longer ISIs. However, during the CP time window, the paired-pulse ratio (PPR) profile takes a distinctive ‘reverse-U’ shape in respect to ISI (Figure 2D), whereby the PPR peaked at 0.5s and fell away at shorter (0.1s: p<0.01 compared to AW) and longer ISIs (1.5s; p<0.01 compared to AW). The same was true during pre-AW, although to a lesser degree (p<0.01 for 0.1s and 1.5s compared to AW). Analysis of MUA (Figure 2C,E) revealed a similar pattern in PPR over varying ISI, with the exception of 0.5s ISI during the CP (Figure 2E). Stimulating whiskers at 0.5s ISI within CP resulted in a larger MUA in response to the second stimulation (p<0.01 compared to pre-AW and AW) and the only positive PPR observed in L4 MUA across early development.

Impact of silenced interneuron signalling on spontaneous cortical activity in neonatal S1BF

GABAergic INs play an important role in cortical circuit formation and maturation (Ben-Ari, Khalilov, Represa, & Gozlan, 2004; S. J. Butt et al., 2017; Modol et al., 2020). However there is little understanding of how GABAergic IN diversity contributes to early activity on the millisecond time scale despite the observation that application of GABA antagonists can disrupt whisker-evoked activity (M. Minlebaev et al., 2011). To address the role for GABAergic IN signalling, we used a conditional Snap25 knockout line (Snap25\textsuperscript{C/C}), to produce mutant animals in which we conditionally abolished action potential-dependent release of GABA (‘silenced’) in SST\textsuperscript{+} (SST\textsuperscript{Cre};Snap25\textsuperscript{C/C}; termed SST\textsuperscript{cs}) or VIP\textsuperscript{+} (VIP\textsuperscript{Cre};Snap25\textsuperscript{C/C}; VIP\textsuperscript{cs}) INs (Marques-Smith et al., 2016; Washbourne et al., 2002), and compared them to wild-type (WT) animals. We first investigated the impact of conditional silencing of SST\textsuperscript{+} or VIP\textsuperscript{+} INs on spontaneous spindle bursts (SB), synchronous neural activity, that is observed in early development, equivalent to our CP window (Rustom Khazipov et al., 2004; Marat Minlebaev et al., 2007). Across all three genetic backgrounds we observed oscillatory
network events in the absence of sensory input (Figure 3A), and quantified their oscillation frequency and duration (Figure 3B). SBs in WT pups had similar rate of occurrence, duration and frequency (Figure 3C-E) to previous studies in mouse (Khazipov & Milh, 2017; Seelke & Blumberg, 2010). However, we observed a decrease in SB occurrence in SST\textsuperscript{cs} animals when compared to both WTs (p<0.01) and VIP\textsuperscript{cs} (p<0.01) animals (Figure 3C). We observed no difference in the duration (Figure 3D) and intra-spindle frequency (Figure 3E) of SB events across all 3 backgrounds. These findings are consistent with the role of thalamus on SB generation (Khazipov et al., 2004). The reduced SB occurrence in SST\textsuperscript{cs} pups can explained in part by weaker thalamic innervation of L4 spiny stellate neurons reported in SST\textsuperscript{cs} animals \textit{in vitro} (Marques-Smith et al., 2016).
We next examined the impact of silencing these INs subtypes on spontaneous cortical action potential discharge (MUA) both during CP and the later time windows (Figure 4A,B). We found that cortical firing is reduced in SSTcs animals during CP compared to WT (p<0.05), while VIPcs animals had comparable cortical firing to WT (Figure 4C). During the pre-AW time window, we observed no difference in the spontaneous firing rate across all three background (Figure 4D). At the onset of AW there was a considerable increase in spontaneous firing rate (Figure 4B,E). This was more pronounced in both SSTcs (p<0.01) and VIPcs (p<0.01) animals which had a higher firing rate than WT (Figure 4B,E). This observation is consistent with SST+ and VIP+ INs regulating spontaneous cortical activity at this later age. The reduced spiking
during CP in SST\textsuperscript{cs} pups most likely reflects the reduction in SBs, since cortical firing is mostly limited to these events during this period (Rustom Khazipov et al., 2004). This further supports a role for SST+ INs in early network formation and function (Marques-Smith et al., 2016; Tuncdemir et al., 2016), one that later switches to an adult-like suppression of cortical activity (Urban-Ciecko, Fanselow, & Barth, 2015).

**Figure 4. SST+ and VIP+ INs control spontaneous action potential discharge in an age dependent manner.** A. Typical spontaneous spiking activity in WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals during CP. B. Typical spiking activity of these animals during AW. C. The average firing rate in WT, SST\textsuperscript{cs}, and VIP\textsuperscript{cs} animals during CP. There was a significant different between genotypes (F(2,21) = 3.956, p<0.05; WT: N = 12, SST\textsuperscript{cs}: N=7, VIP\textsuperscript{cs}: N = 5). D. The average firing rate in these genotypes during pre-AW. There was no significant effect for genotype (F(2,21) = 0.048, p=0.952. WT: N = 13, SST\textsuperscript{cs}: N = 5, VIP\textsuperscript{cs}: N = 6) E. The average firing rate in these genotypes during AW. There was a significant effect for genotype (F(2,31) = 6.850, p<0.01. WT: N = 22, SST\textsuperscript{cs}: N = 6, VIP\textsuperscript{cs}: N = 6). * p<0.05 in a multiple comparison after an ANOVA.

**Contrasting Dynamics of SST+ and VIP+ in Sensory Responses**

Our observation that INs differentially modulate spontaneous activity through development led us to ask how these interneurons might contribute to emergent sensory processing. We first examined how IN silencing affects the response to a single multi-whisker stimulation presented at different speeds. As before (Figure 1), we focused on our analysis on the LFP in the granular layer (Figure 5A-C). Silencing SST+ or VIP+ IN populations had no effect on the ability to encode speed across early development: across all three backgrounds and development time windows, we consistently observed a decrease in peak latency (Figure S1A-C) and an increase in amplitude (Figure 5A-C) for the L4 sensory responses according to speed. However, we observed an effect of IN silencing on the amplitude of the neural response to a single stimulus. During CP, both SST\textsuperscript{cs} (p<0.05) and VIP\textsuperscript{cs} (p<0.01) animals exhibited
a reduced amplitude of the sensory response compared to controls (Figure 5A). During pre-AW, the response amplitude observed in SST\textsuperscript{cs} animals was comparable to WT, whereas VIP\textsuperscript{cs} animals exhibited a larger response than both WTs and SST\textsuperscript{cs} (Figure 5B; p<0.01 in both cases). This changed during AW: VIP\textsuperscript{cs} animals were indistinguishable from WT, whereas SST\textsuperscript{cs} animals exhibited delayed latency (Figure S1C; p<0.01 compared to WTs and VIP\textsuperscript{cs}) and reduced amplitude of the sensory response compared to both WTs and VIP\textsuperscript{cs} animals (Figure 5C; p<0.01 compared to WT and VIP\textsuperscript{cs}). Analysis of MUA (Figure 5D-F) revealed a similar pattern to that observed in the LFP with one significant exception: During AW, MUA in L4 of SST\textsuperscript{cs} was indistinguishable from that recorded in WT and VIP\textsuperscript{cs} animals (Figure 5F), consistent with delayed maturation of thalamic input into L4 S1BF in these animals (Marques-Smith et al., 2016)
Figure 5. Time-dependent effect of SST+ and VIP+ IN silencing on SER.

A. Top: LFP response after an 80Hz deflection in the granular layer of SSTcs and VIPcs animals during CP. Bottom: the average peak amplitude of WT (n = 11), SSTcs (n=7) and VIPcs (n=6) animals during CP. There was a significant effect for speed (F(4,48) = 6.743, p<0.001) and genotype (F(2,48) = 3.665, p < 0.05).

B. Top: LFP response in the granular layer of SSTcs and VIPcs animals during pre-AW. Bottom: the average peak amplitude of WT (n=9), SSTcs (n=5) and VIPcs (n=6) animals during pre-AW. There was a significant effects for speed (F(4,85) = 6.373, p<0.001) and genotype(F(2,85) = 11.59, p<0.001).

C. Top: LFP response in the granular layer of SSTcs and VIPcs animals during AW. Bottom: average peak amplitude of WT (n=21), SSTcs (n=6) and VIPcs (n=6) animals during AW. There was a significant effects for speed (F(4,152) = 7.205, p<0.001) but not for genotype (F(2,152) = 9.600, p<0.001).

D. Top: MUA response after an 80Hz deflection in the granular layer of SSTcs and VIPcs animals during CP. Bottom: average peak firing-rate of WT (n=8), SSTcs (n=5) and VIPcs (n=5) animals during CP. There was an effect for speed (F(4,70) = 4.58,
p<0.001), and genotype (F(2,70) = 3.834, p <0.05).

E. Top: MUA response in the granular layer of SSTcs and VIPcs animals during pre-AW. Bottom: average peak firing-rate of WT (n=6), SSTcs (n=4) and VIPcs (n=6) animals during pre-AW. There was no effect for speed (4,60) = 1.650, p = 0.0.173), but there was an effect for genotype (F(2,60) = 7.208, p<0.01).

F. Top: MUA response in the granular layer of SSTcs and VIPcs animals during AW. Bottom: average peak firing-rate of WT (n=21), SSTcs (n=6) and VIPcs (n=6) animals during AW. There was an effect for speed (F(4,147) = 16.81, p<0.001), but not for genotype (F(2,147) = 2.404, p = 0.094. Brackets show a p<0.05 multiple comparison after an ANOVA.

A role for SST+ but not VIP+ INs in paired-pulse facilitation.

We examined the paired-pulse response in the LFP (Figure 6A,B) and found that in both SSTcs and VIPcs there was a linear relationship between PPR and ISI. PPR increased in SSTcs across the developmental windows tested (Figure 6C; p<0.01 between all age groups) whereas in VIPcs animals there was a decrease in PPR during pre-AW not observed in WT and SSTcs animals (Figure 6D; p<0.01 compared to both CP and AW). We then compared the PPR between the three backgrounds across the different time-points. This revealed that SSTcs animals had a lower PPR than both VIPcs (p<0.01) and WT animals (p<0.01) during CP (Figure S2A,B). But, during pre-AW, VIPcs had a lower PPR over the shorter ISIs compared to controls (Figure S2B; p<0.01 for 0.1s and 0.25s ISI). All three backgrounds showed a similar PPR profile during AW (Figure S2A,B). Our subsequent analysis of MUA responses (Figure 6E, F) revealed that silencing SST+ INs abolished the positive PPR for 0.50 ISI (Figure 6E, G) observed in controls (Figure 2E, Figure S2C). Silencing VIP+ INs resulted in greater adaptation following paired stimuli at short ISIs up until AW (Figure S2D), but otherwise the profile was similar to control animals (Figure 6H). These data suggest that GABAergic signalling via SST+ INs play a role in paired-pulse facilitation of MUA observed during CP, whereas an absence of signalling from VIP+ INs impairs the response to stimuli presented at short ISIs, prior to AW.
A LFP
ISI: 0.50 s

B

C

D

E MUA
ISI: 0.50 s

F

G

H

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Figure 6. Age-dependent effect of SST+ and VIP+ IN silencing on adaptation. A. LFP adaptation response of a SSTcs animal to two consecutive deflections with a 0.50s ISI during CP and AW. B. LFP adaptation response of VIPcs animal to two consecutive deflections with a 0.50s ISI during CP and AW. C. Average PPR of the LFP response of SSTcs animals for different ISIs. There was a significant change with ISI (F(4,95) = 18.31, p<0.01) and genotype (F(2,95) = 31.20, p<0.01), but no interaction (F(8,95) = 1.67, p=0.11); CP: N = 9, PAW: N = 6, AW: N = 7. D. Average PPR of the LFP response of VIPcs animals for different ISIs. There was a significant effect for both ISI (F(4,80) = 20.67, p<0.01) and age (F(2,80) = 10.10, p<0.01) but no interaction (F(8,80) = 1.66, p = 0.12); CP: N = 7, PAW: N = 6, AW: N = 6. E. MUA adaptation response of SSTcs animal to two consecutive deflections with a 0.50s ISI during CP and AW. F. MUA adaptation response of VIPcs animal to two consecutive deflections with a 0.50s ISI during CP and AW. G. Average PPR of the MUA response of SSTcs animals for different ISIs. There was a significant change with ISI (F(4,95) = 21.59, p<0.01) and genotype (F(2,95) = 27.40, p<0.01), but no interaction (F(8,95) = 1.36, p =0.23); CP: N = 8, PAW: N = 4, AW: N =6. H. Average PPR of the MUA response of VIPcs animals for different ISIs. There was an interaction between age and ISI (F(8,50) = 4.10, p < 0.01); CP: N = 4, PAW: N = 6, AW: N = 6. Brackets signify p<0.05 in a simple comparison post ANOVA. * signify p<0.05 in a simple multiple comparison.

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GABAergic INs, notably SST+ and VIP+ subtypes, are not evenly distributed across the layers of neocortex (X. Xu, Roby, & Callaway, 2010). Moreover, in vitro data has identified the presence of transient translaminar networks mediated by both subtypes during early postnatal life (Marques-Smith et al., 2016; Vagnoni et al., 2020). During the CP time window, MUA is largely confined to granular L4. As such we focused our investigation across supragranular (SG) and infragranular (IG) layers to the pre-AW and AW time windows. During pre-AW, the peak firing rate in SG layers was not affected by silencing of either IN subtype (Figure 7A). However we did observed a decrease in IG layer MUA in both VIPcs and SSTcs compared to WT animals (p<0.01 in both cases) (Figure 7C). The latency of the sensory response was slower across both SG and IG layers in both SSTcs and VIPcs animals (Figure S3A,S3C; p<0.05) compared to wild-type. During AW, both SG and IG response amplitudes of the SSTcs animals had increased MUA compared to both VIPcs and WT animals (Figure 7B; p<0.05), consistent with an inhibitory role for this IN subtype (Naka et al., 2019). In contrast, the IG response of VIPcs animals was lower than WT animals (p<0.05), consistent with a dis-inhibitory role for this IN subtype in IG layers (Pfeffer, Xue, He, Huang, & Scanziani, 2013; Pi et al., 2013). Latencies were similar regardless of genotype (Figure S3B,D) with the exception of the response latency of SSTcs animals at the slowest deflection speed (Figure S3B; p<0.05) in the SG layers. The paired-pulse response was consistently altered across layers with responses in both SSTcs...
and VIP\textsuperscript{cs} animals, having lower PPRs, signifying stronger adaptation, than WTs, especially at shorter ISIs (Figure S4A,B; p<0.05). During the AW time window the PPR of the responses were comparable to controls in line with observations from granular L4 (Figure S4C,D).

**Figure 7.** Layer-specific effects of SST\textsuperscript{+} and VIP\textsuperscript{+} IN silencing on SER. A. Top. SG evoked response traces for the three genotypes. Bottom. The average pre-AW MUA response for a single whisker deflection in the SG layers of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals. There was an effect for speed($F(4,58)=2.44$, p<0.05) but not for genotype($F(2,58)=0.18$, p= 0.84; WT: N =5, SST\textsuperscript{cs}: N =4, VIP\textsuperscript{cs}: N= 6) B. The average AW MUA response for a single whisker deflection in the SG layers of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals. There was an effect for both speed ($F(4,142)=7.42$, p<0.01) and genotype ($F(2,142)= 5.60$, p<0.01; WT :N=20, SST\textsuperscript{cs}: N = 6 ;VIP\textsuperscript{cs} :N=6) C. The average pre-AW MUA response for a single whisker deflection in the IG layers of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals. There was an effect for both
speed(F(4,68) = 13.84, p<0.01) and genotype(F(2,68) = 25.24, p<0.01; WT: N =6, SST\textsuperscript{cs}: N =5, VIP\textsuperscript{cs}: N= 6) D. The average AW MUA response for a single whisker deflection in the IG layers of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals. There was an effect for both speed (F(4,142) =7.87 ,p<0.01) and genotype(F(2,142) =7.40 , p<0.01; WT: N = 21, SST\textsuperscript{cs}: N =6, VIP\textsuperscript{cs}: N= 6 ). Brackets signify p<0.05 in a post-hoc multiple comparison.

Discussion

The contribution of GABAergic interneuron subtypes to early sensory-evoked activity on the millisecond timescale is poorly understood. In this study we have used genetic silencing of GABA release in interneurons to determine the role that SST+ and VIP+ INs play in the acquisition of normal sensory function in S1BF during the first few weeks of development. Analysis of our in vivo data reveals a different role for these two subtypes in mouse: SST+ INs contribute to thalamocortical maturation and plasticity in line with previous in vitro circuit analysis, whereas VIP+ INs regulate spiking activity through both inhibitory and dis-inhibitory mechanisms towards the onset of active whisking.

To study the early role of INs we conditionally deleted exons 5 and 6 of the t-SNARE protein Snap25 in SST+ and VIP+ neurons, thereby preventing action potential-dependent neurotransmitter release (Marques-Smith et al., 2016; Verhage et al., 2000; Washbourne et al., 2002). We favoured this approach over alternative genetic manipulations – including expression of potassium rectifier channels – as we felt it was more selective in preventing GABAergic signalling through the time window of our analysis from the onset of the critical period of plasticity to active whisking. We focused on two of the major cortical IN subtypes, SST+ and VIP+ INs, which we could reliably target genetically using Cre lines (Taniguchi et al., 2011). The lack of a specific Cre line for fast spiking, PV+ basket cells at early postnatal ages precluded assessment of these INs during the time window analysed.

Given the key role of locally projecting GABAergic INs in balancing excitation and inhibition in adult neocortex, it is unsurprising that silencing INs led to an increase in spontaneous activity at the later ages tested, at the onset of active perception. However, in the case of VIP+ INs this is counterintuitive given the body of data that suggest these cells exert a primarily dis-inhibitory effect on pyramidal cells via the inhibition of SST+ INs (Pfeffer et al., 2013; Pi et al., 2013). That said, this observation echoes recent findings that suggest that VIP+ INs directly inhibit pyramidal cells in a
state-dependent manner (Batista-Brito et al., 2017; Vagnoni et al., 2020). Moreover, it is clear from the effect of silencing INs on spontaneous activity that VIP+ INs develop to control spontaneous pyramidal cell activity around the onset of active whisking. In contrast, our evidence shows that SST+ INs contribute to activity as early as the critical period of plasticity. Silencing action potential-dependent release of GABA from this IN subtype led to a decrease in spontaneous SB and associated spike activity at early ages. Early SB activity is thought to be important for normal sensory development and play a role in the prevention of activity-dependent apoptosis, amongst other formative processes (Khazipov & Milh, 2017). A number of potential mechanisms could explain the effect of SST+ IN silencing on SB generation: first, we broadly targeted SST+ INs. This will affect signalling elsewhere in forebrain, notably the ventral posteromedial nucleus (VPM) of the thalamus. Second, silencing SST+ INs could lead to an increase in local GABAergic signalling through dis-inhibition. Third, silencing SST+ INs could result in delayed maturation of thalamic innervation of neocortex given the role of these cells in early thalamocortical circuits (Yang et al., 2013). The first option is unlikely given that intra-thalamic inhibiton has not fully matured at this age (Zolnik & Connors, 2016) and intra-thalamic SST+ neurons do not synapse onto VPM relay neurons (Clemente-Perez et al., 2017). Our in vivo observations are consistent with our previous study that identified both delayed thalamocortical innervation and compensatory increase in the local GABAergic innervation – most likely from immature basket cells (Daw et al., 2007), following SST+ IN silencing in vitro (Marques-Smith et al., 2016). Moreover our data are consistent with the thalamic origin of SB generation (Marat Minlebaev et al., 2007) but further support the notion that interneurons circuits across infra- and granular layers interpret afferent sensory signalling to constrain and direct circuit development (Marques-Smith et al., 2016).

Encoding velocity is a key requirement for somatosensory detection by the vibrissae (Kleinfeld et al., 2006; Szwed, Bagdasarian, & Ahissar, 2003). We could detect speed coding in the cortex from the earliest postnatal time point recorded. This suggests that while this sensory computation develops independent of cortical maturation, probably as a result of phase coding as early as at the brainstem level of sensory processing (Szwed et al., 2003; Wallach, Bagdasarian, & Ahissar, 2016). This is entirely consistent with other reports that have identified various stimulus properties encoded in the VPM in adult animals, including speed (Bale, Ince, Santagata, &
Petersen, 2015). Further support for upstream processing of speed comes from the lack of an effect for SST+ or VIP+ IN silencing on this computation.

In contrast to speed coding, the profile of sensory response adaptation changed over development. In young animals – during the critical period for plasticity in L4, adaptation took an ‘Inverse-U’ shape with significant depression of the second response at both short and long inter-stimulus intervals, while an interval of 0.5s led to its facilitation. The depression of the second response at short intervals in young animals can likely be explained by low release probability of the immature thalamocortical synapses (Crair & Malenka, 1995; Isaac, Crair, Nicoll, & Malenka, 1997). However, this is less likely to underpin the attenuation observed at longer intervals, which could involve recurrent GABAergic networks. Certainly, it would appear that SST+ INs contribute to the observed facilitation at 0.5s interval as this is abolished in animals in which SST INs are silenced. This can be either directly through facilitation of the TC input onto pyramidal cells through excitatory GABAergic signalling (Ben-Ari, 2002), or indirectly through SST+ IN influence on local GABAergic circuits, as was shown before in the infragranular layers of S1BF (Tuncdemir et al., 2016).

Another possibility is that the altered facilitation in SSTCS animals is a by-product of the attenuated thalamic input in these animals. However, it is less likely, as we do not observe any impact of VIP+ IN silencing on the facilitation despite these animals also exhibiting reduced thalamic input. Of note is the fact that the 0.5s interval corresponds to the frequency of whisker stimulation that evokes the largest haemodynamic response in S1BF (Sintsov, Suchkov, Khazipov, & Minlebaev, 2017), and results in long term potentiation (LTP) during this developmental time window (An, Yang, Sun, Kilb, & Luhmann, 2012). Together these lines of evidence support the notion that this specific interval is physiologically relevant for the young animal, matching slow passive stimuli it receives, mostly from its littermates (Akhmetshina, Nasretdinov, Zakharov, Valeeva, & Khazipov, 2016). Moreover that it requires functional SST+ IN network.

In adult mice, SST+ and VIP+ INs have been shown to have layer-specific functions (Muñoz et al., 2017; Pfeffer et al., 2013; Pi et al., 2013; H. Xu, Jeong, Tremblay, & Rudy, 2013). Though our work focused primarily on granular L4 – given the consistency of the sensory-evoked response in this layer, we did observe layer-specific changes in sensory processing in our genetically modified mice across the developmental time window tested. In adult mice SST+ INs exert an inhibitory effect in the upper layers but are dis-inhibitory in L4 (H. Xu et al., 2013). We observed an
increase in sensory-evoked spiking in supragranular layers after the onset of active whisking, suggesting that the inhibitory effect reported in adults emerges in line with active somato-sensation. VIP+ INs, present mostly in upper layers, have a dis-inhibitory role in the mature cortex (Muñoz et al., 2017; Pfeffer et al., 2013; Pi et al., 2013). In our animals, silencing this IN subtype led to a decrease in response, consistent with dis-inhibition. However, this effect was mostly in the lower, infragranular layers. We did not observe any change in supragranular layer activity, in line with the late integration of VIP+ INs in the local network (Batista-Brito et al., 2017; Vagnoni et al., 2020). Before the onset of active sensation, VIP+ IN silencing led to a transient increase in sensory response in the granular layer. This is consistent with previous findings showing an increase in synapses between these INs and pyramidal cells during this time period (Vagnoni et al., 2020). Overall, the changes we observed are consistent with an inside-out pattern of innervation involving both IN subtypes, whereby the interneurons first integrate in infragranular layers before sequentially innervating supragranular target neurons.

Taken together, our results show that sensory processing develops in line with cortical maturation. We demonstrate that SST+ and VIP+ INs both contribute to early processing of sensory information, with SST+ INs having a distinct role in early regulation of spontaneous activity and facilitation. VIP+ INs play more of a role toward the onset of active perception, regulating incoming sensory information. Our data identify the importance of IN diversity in in vivo cortical processing, across early postnatal development.

Author Contribution
LJB, MK and SJBB conceived experiments which were conducted by LJB, who in addition analysed the data. LJB and SJBB wrote the manuscript. All authors edited the manuscript.

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Supplementary Figures

Figure S1. A. The average peak LFP latency of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals during CP. There was a significant effects for speed (F(4,98) = 25.26, p<0.001) but not for genotype(F(2,98) = 2.14, p=0.123 ). B. The average LFP latency of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals during pre-AW. There was a significant effects for speed (F(4,85) = 5.826, p<0.001) but not for genotype (F(2,85) = 1.148, p=0.322 ). C. The average LF latency of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals during AW. There was a significant effect for speed (F(4,152) = 24.33, p<0.001) and for genotype(F(2,152) = 15.35, p<0.001 ). D. The MUA latency of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals during CP. There was an effect for speed (F(4,70) = 30.04, p<0.01), but not for genotype (F(2,70) = 0.75, p=0.475) E. The average MUA latency of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals during pre-AW. There was a genotype X speed interaction (F(8,60) = 2.374, p< 0.05), due to SST\textsuperscript{cs} being slower at 1Hz deflections (p<0.05). F. The average MUA latency of WT, SST\textsuperscript{cs} and VIP\textsuperscript{cs} animals during AW. There was an effect for speed (F(4,147) = 15.29, p<0.001), but not for genotype (F(2,147) = 2.488, p = 0.086). Brackets signify p<0.05 for a post-hoc multiple comparison. * p<0.05 for a simple post-hoc multiple comparison.
Figure S2. A. The average difference in LFP PPR of SST<sup>cs</sup> animals compared to the average in controls over development. B. The average difference in LFP PPR of VIP<sup>cs</sup> animals compared to the average in controls over development. C. The average difference in MUA PPR of SST<sup>cs</sup> animals compared to the average in controls over development. D. The average difference in MUA PPR of VIP<sup>cs</sup> animals compared to the average in controls over development.
Figure S3. The effect of VIP+ and SST+ IN silencing of response latency. 

A. Average response latencies in SG layers during pre-AW. There was an effect for both speed (F(4,58) = 11.51, p<0.01) and genotype (F(2,58) = 4.71, p<0.01). 

B. Average response latencies in SG layers during AW. There was an interaction between genotype and speed (F(8,142) = 4.44, p<0.01). The interaction stemmed from the 5 Hz deflection where both silenced genotypes had slower latencies (p<0.01). 

C. Average response latencies in IG layers during pre-AW. There was an effect for both speed (F(4,68) = 14.10, p<0.01) and genotype (F(2,68) = 8.17, p<0.01). 

D. Average response latencies in IG layers during AW. There was only an effect for speed (F(4,142) = 13.22, p<0.01), but not for genotype (F(2,142) = 2.31, p=0.10).
Figure S4. A. The average PPR of the MUA response in the SG layer of WT, SST\textsuperscript{cs}, and VIP\textsuperscript{cs} animals during pre-AW. There was an effect for both ISI (F(4,100) = 11.61, p<0.01) and genotype (F(2,100) = 4.336, p<0.05; WT: N = 13, SST\textsuperscript{cs}: N = 4, VIP\textsuperscript{cs}: N = 6). B. The average PPR of the MUA response in the IG layer of WT, SST\textsuperscript{cs}, and VIP\textsuperscript{cs} animals during pre-AW. There was an effect for both ISI (F(4,120) = 11.40, p<0.01) and genotype (F(2,120) = 9.013, p<0.01; WT: N = 18, SST\textsuperscript{cs}: N = 4, VIP\textsuperscript{cs}: N = 6). C. The average PPR of the MUA response in the SG layer of WT, SST\textsuperscript{cs}, and VIP\textsuperscript{cs} animals during AW. There was an effect for ISI (F(4,173) = 8.521, p<0.01), but not for genotype (F(2,173) = 1.673, p=0.191; WT: N = 27, SST\textsuperscript{cs}: N = 5, VIP\textsuperscript{cs}: N = 6). D. The average PPR of the MUA response in the IG layer of WT, SST\textsuperscript{cs}, and VIP\textsuperscript{cs} animals during AW. There was an effect for ISI (F(4,188) = 16.46, p<0.01) but not for genotype (F(2,88 = 0.671, p=0.512; WT: N = 30, SST\textsuperscript{cs}: N = 6, VIP\textsuperscript{cs}: N = 6)
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