






# Site-specific seeding of Lewy pathology induces distinct pre-motor cellular and dendritic vulnerabilities in the cortex

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
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Circuit-based biomarkers distinguishing the gradual progression of Lewy pathology across synucleinopathies remain unknown. Here, we show that seeding of  $\alpha$ -synuclein preformed fibrils in mouse dorsal striatum and motor cortex leads to distinct prodromal-phase cortical dysfunction across months. Our findings reveal that while both seeding sites had increased cortical pathology and hyperexcitability, distinct differences in electrophysiological and cellular ensemble patterns were crucial in distinguishing pathology spread between the two seeding sites. Notably, while beta-band spike-field-coherence reflected a significant increase beginning in Layer-5 and then spreading to Layer-2/3, the rate of entrainment and the propensity of stochastic beta-burst dynamics was markedly seeding location-specific. This beta dysfunction was accompanied by gradual superficial excitatory ensemble instability following cortical, but not striatal, preformed fibrils injection. We reveal a link between Layer-5 dendritic vulnerabilities and translaminal beta event dysfunction, which could be used to differentiate symptomatically similar synucleinopathies.

The accumulation of misfolded  $\alpha$ -synuclein ( $\alpha$ Syn) in cortical Lewy bodies is a hallmark of Parkinson's disease dementia (PDD) and dementia with Lewy bodies (DLB), together known as Lewy Body Dementias. Here, fibrils comprising misfolded  $\alpha$ Syn propagate trans-neuronally through anatomically interconnected networks, amplifying disease pathology by templating aggregation across molecularly defined cell-types<sup>1–6</sup>. How circuit activity across these vulnerable pathways manifests at the earliest stages of  $\alpha$ Syn insult and evolves as a function of PDD and DLB progression remains unknown. Unraveling such “circuit-level biomarkers” is critical given the symptomatically similar end-stage behavioral deficits observed across the two forms of the disease, for which, at present, no biomarker exists that reliably differentiates between the two.

PDD and DLB are characterized not only by motor deficits but also by cognitive decline, albeit not in the same order<sup>7,8</sup>. In PDD, Lewy pathology is synonymous with striatal dysfunction facilitated by neurodegeneration of dopaminergic neurons in the substantia-nigra pars compacta (SNc)<sup>8</sup>, with cortical dysfunction occurring subsequently<sup>9</sup>. In contrast, in DLB, motor dysfunction occurs within a year of cognitive decline, affecting cortical areas first<sup>10</sup>. These features suggest that cortical vulnerabilities are not necessarily a late-stage effect, as proposed in the Braak et al., hypothesis<sup>11</sup>, but could serve as a critical top-down mediator of motor dysfunction, a notion gaining traction<sup>9,12</sup>. Despite this, it remains unclear how cell-to-cell transmission of misfolded  $\alpha$ Syn occurs under PDD and DLB pathogenesis and impacts activity at the cellular, multicellular, and circuit levels<sup>9</sup>. As the motor

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cortex is vulnerable across both PDD and DLB<sup>7,8</sup>, the influence of  $\alpha$ Syn pathology on recurrent translaminal connections is likely to be significant<sup>13</sup>, wherein the activity of even a single neuron can result in a non-linear change within the network<sup>4,14</sup>. This framework agrees well with the work of Nemani et al., who showed that modest overexpression of  $\alpha$ Syn, albeit in the striatum, does not lead to stark toxicity-related changes. Instead, it impairs glutamate and dopamine release, which could lead to network dysfunction and eventual cognitive change—a feature that could be widespread during the prodromal-phase in the cortex<sup>15,16</sup>. Yet, we know little about the relationship between LB formation and the evolution of functional circuit changes, and whether such shifts are intrinsic to the neurons or a product of functionally connected ensembles of recurrent neural populations. To dissect such changes, one must measure and correlate dynamic activity patterns, including spatial and temporal properties, to pathology spread<sup>4,17</sup>. Such changes could be assayed through cellular-level ensemble dynamics, layer-specific spiking dynamics, and spike-field coherences, which we combinatorially highlight in this study.

Functional neural populations, termed ensembles<sup>18–21</sup>, represent stable multicellular activity maps that predict perception, action, and behavior better than the sum of their constituent neurons<sup>19,20</sup>. In this regard, a recent study using two-photon calcium imaging reported cortical dysfunction across superficial layers under striatal  $\alpha$ Syn pathology load<sup>17</sup>. The study highlighted cortical hyperactivity and possible excitation-inhibition imbalance 9 months post- $\alpha$ Syn-preformed-fibril ( $\alpha$ Syn PFFs) administration but did not capture the evolution of translaminal circuit changes or ensemble-level pattern differences. A similar study by the same authors showed that striatal  $\alpha$ Syn PFF injection results in the loss of spines on pyramidal Layer 5 (L5) neurons in the anatomically connected somatosensory cortex (S1)<sup>22</sup>. Together these studies suggest a cortical pathway for Lewy pathology spread via L5 projection neurons to Layer-2/3 (L2/3) neurons<sup>23,24</sup>, highlighting the need for mapping recurrent translaminal circuit changes. Such changes in deep-layer neurons may contribute to a local network amplification of pathophysiology that leads to impairments at the ensemble level in superficial layers—a feature that potentially reflects the effects of pathology on integration across ascending and descending pathways<sup>24,25</sup>, highlighting selective synaptic vulnerabilities within sensorimotor cortices<sup>25–28</sup>. Thus, key differences in bidirectional L5-L2/3 coupling as well as eventual L2/3 ensemble dynamics could lead to unique biomarkers distinguishing different forms of synucleinopathies.

From an electrophysiological perspective, robust task- or stimulus-driven beta dynamics are observed in physiological brain states and are linked to motor preparation<sup>29</sup> and action<sup>29</sup>, with more recent studies expanding their involvement to encompass a broader range of cognitive functions, such as working memory<sup>30–32</sup>, long-term encoding<sup>33,34</sup>, and decision-making<sup>35,36</sup>. In the cortex, translaminal network synchronization shifts between superficial and deep cortical layers, with slower beta oscillations driven by deep-layer neurons and superficial layers dominated by high-frequency gamma oscillations<sup>30,32</sup>, suggesting that the encoding and maintenance of cortical state is dependent on precisely timed nested oscillatory dynamics<sup>31</sup>. In PD and related dementias, there is strong correlative evidence linking elevated subcortical beta-band activity at rest and beta-power changes in response to stimulation<sup>37,38</sup>. Beta expression is thus primed to strongly predict healthy and pathological motor performance<sup>39–41</sup>. Recently, such beta activity changes were also observed in the olfactory bulb of mice seeded with  $\alpha$ Syn PFFs<sup>42</sup>, suggesting that aberrant beta may not be solely relegated to subcortical motor areas, but might also be highly prevalent in cortical regions impacted by vulnerable pathways<sup>43</sup>. This aspect could be critical, especially in delineating between PD and DLB progression. However, a question that remains unanswered is whether beta and its

pathologically driven amplification mechanisms<sup>16,44</sup> could serve as reliable biomarkers of disease progression.

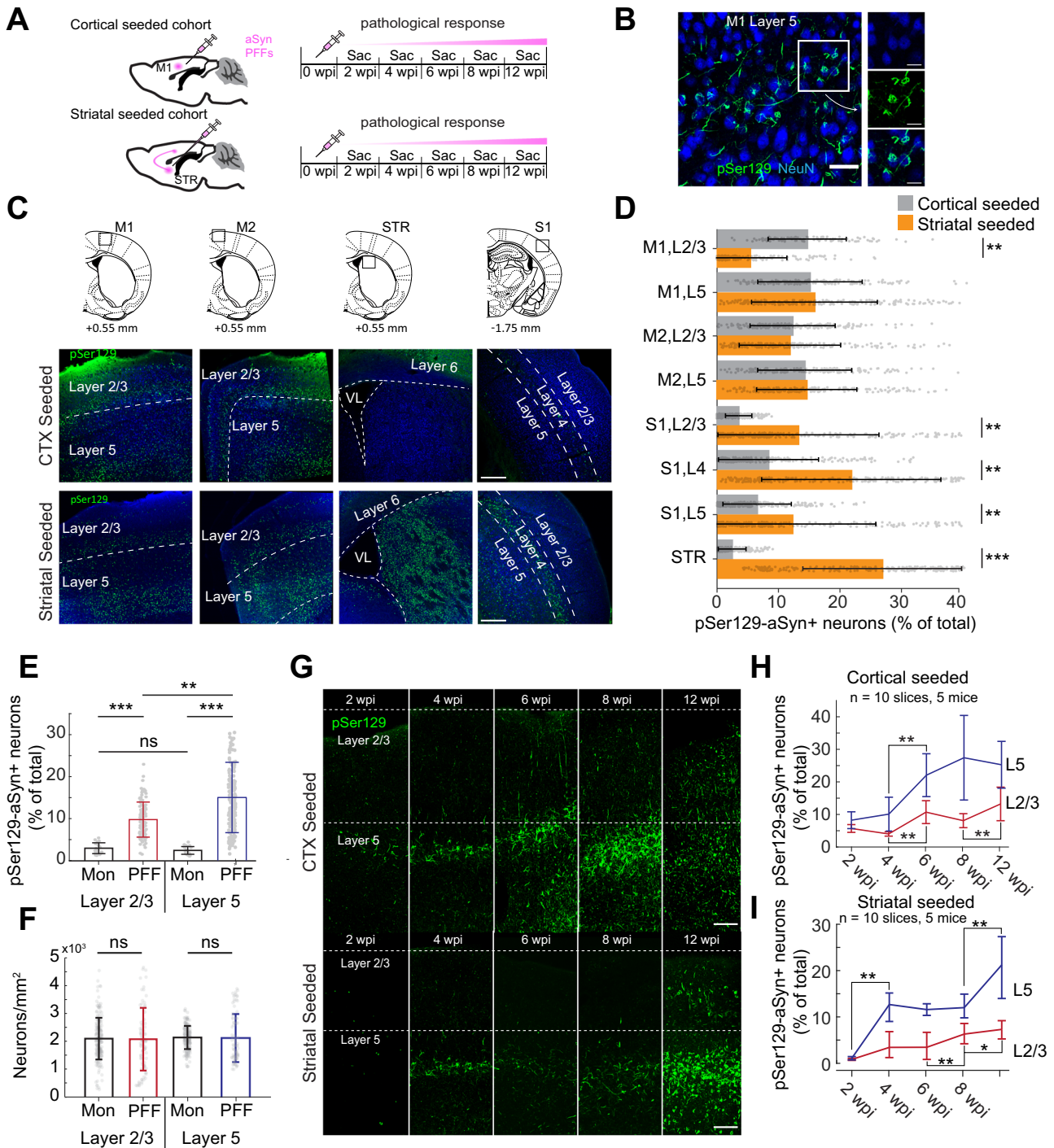
Evidence from computational modeling and experimentation suggests that the after-hyperpolarization and M-type potassium currents of layer 5 (L5) pyramidal neurons are sufficient to drive the firing of these neurons at beta frequencies<sup>45</sup>. Computational evidence, backed by human EEG and MEG recordings<sup>46</sup> suggests that, in the motor cortex of both mice and humans, transient beta events<sup>41,47,48</sup> waves that last ~50 ms—emerge from the integration of bottom-up and top-down inputs along the distal and proximal dendrites of cortical pyramidal neurons. This is significant because these bursts could reflect the hyperexcitability of multiple neurons associated with  $\alpha$ Syn pathology<sup>49</sup> in a pathway-specific manner. Unraveling the cortical biophysical mechanism of these aberrant beta bursts, as well as their specific spatiotemporal interactions in healthy and diseased cortical networks, will help elucidate key biomarkers differentiating across similar syndromes that share characteristic Lewy deposits<sup>8</sup>.

In this study, we injected  $\alpha$ Syn PFFs in the dorsal striatum and deep-layer motor cortex to induce phosphorylated, aggregated inclusions resembling human LBs<sup>2,50</sup>. We assayed translaminal patterns of motor cortical activity and behavior across a 3- and 12 month pathological timeline—respectively—and revealed how  $\alpha$ Syn PFFs propagating through two distinct pathways generate separable biomarkers within recurrent circuits. First, as a function of seeding location (cortical vs. striatal) and over 12 weeks of Lewy pathology spread, we show that there is diverging phase entrainment between L2/3 and L5 relegated only to beta, and not gamma or theta frequencies. Second, we reveal that cortical seeding destabilizes neuronal ensembles, a feature absent in the brains of mice subjected to striatal seeding. We discovered that such destabilization is delicately tied to shifting patterns in top-down integration, reflected in the form of moment-to-moment bursts of beta transients ( $\beta$ -events). Mechanistically, we show that this increase in  $\beta$ -event power is tightly modulated by apical L5 dendritic  $\text{Ca}^{2+}$  currents, a critical marker of top-down modulation, and that upregulation of N-Methyl-D-Aspartate (NMDA) phosphorylation could drive aberrancies in  $\beta$ -events in both pathological models. Importantly, we describe how such patterns of cortical circuit dysfunction manifested before behavioral deficits, which were observed 1 year after injections under both seeding strategies. Our observations strongly suggest that translaminal spike-field patterns coupled with functional neural ensemble dynamics in the superficial cortex serve as powerful biomarkers and can be used “combinatorially” to differentiate between prodromal-phase synucleinopathies.

## Results

### Characterizing the early spread of Lewy pathology across vulnerable pathways

To explore the extent of pathogenic spread of Lewy bodies and its contribution to circuit dysfunction and prodromal-phase pre-motor deficits, we injected recombinant  $\alpha$ Syn-PFFs (Supplementary Fig. S1a, S1b) in L5 of the mouse primary motor cortex (M1) or dorsal striatum (Fig. 1A, left), and conducted a pathological assessment over a period of 12 weeks post-injection (wpi) (Fig. 1A, right). We compared the pathological burden of  $\alpha$ Syn from both strategies to monomer synuclein injections in both regions in a separate cohort of mice. The injection of  $\alpha$ Syn PFFs in either the cortex or striatum triggered the formation of intracellular Lewy body- and Lewy neurite-like aggregates within M1 (Fig. 1B, C). Interestingly, we also observed the emergence of  $\alpha$ Syn pathology in various regions projecting to the motor cortex and striatum such as the sensorimotor cortex after 12 wpi (Fig. 1C, Supplementary Fig. S1c). Following M1 seeding, we noted apparently higher pathological expression in layer 5 of primary and secondary motor cortex (percentage of pSer129- $\alpha$ Syn+ neurons =  $15.03 \pm 8.38\%$  and  $14.13 \pm 7.54\%$ , respectively) and layer 4 of primary somatosensory cortex ( $8.33 \pm 8.02\%$ ) in comparison to their more superficial layer 2/3



**Fig. 1 | Recombinant preformed fibrils induce gradual progression of  $\alpha$ Syn pathology across layers of the motor cortex.** **A** Schematic of  $\alpha$ Syn PFF injection into the primary motor cortex or dorsal striatum in separate cohorts of mice (left). Pathological assessment of evolving  $\alpha$ Syn aggregate spread over a 12 week timeline (right). **B** Representative image showing pSer129- $\alpha$ Syn+ Lewy body- and Lewy neurite-like aggregates in Layer 5 of the primary motor cortex in a single cortically seeded mouse. Scale bar = 20  $\mu$ m. **C** Pathological pSer129- $\alpha$ Syn+ neurons in select brain regions 12 weeks post-injection in both the cortex and striatum (VL, lateral ventricle). Scale bar: 200  $\mu$ m. **D** Quantification of pSer129- $\alpha$ Syn+ neurons (% of total NeuN+ cells) in gross anatomical regions ( $n = 288, 392, 588, 370$  neurons sampled across 4 slices across 4 mice (2 mice for cortical seeding and 2 mice for striatal seeding) for each region of M1, M2, S1, and STR respectively, one-way ANOVA with Bonferroni comparison test, \*\* $p < 10^{-2}$ , \*\*\* $p < 10^{-4}$  data shown as mean  $\pm$  sd). **E** Comparison of pSer129- $\alpha$ Syn+ neuron counts in PFF- versus monomer-injected mice 12 weeks post injection in the cortex ( $n = 4$  slices, 340 neurons sampled, 4

mice, two-sided Mann-Whitney  $U$ -test \*\* $p < 10^{-7}$ , \*\*\* $p < 10^{-10}$ , data shown as mean  $\pm$  sd). **F** Density of NeuN-expressing neurons compared across L2/3 and L5 layers in PFF- and monomer-injected mice 12 weeks post injection in the cortex ( $n = 4$  slices, 244 regions sampled, two-sided Mann-Whitney  $U$ -test  $p = 0.5$ ,  $p = 0.11$  for L2/3 and L5 respectively, data shown as mean  $\pm$  sd). **G** Progressive accumulation of pSer129- $\alpha$ Syn+ puncta in primary motor cortex and dorsal striatum from 2 to 12 weeks post injection. Scale bar: 100  $\mu$ m. **H** Comparison of pSer129+ neuron counts across weeks in the M1 region of mice injected with  $\alpha$ Syn PFFs in primary motor cortex ( $n = 987$  sampled neurons, 10 slices across 5 mice, 2 slices per week data shown as mean  $\pm$  sem). **I** Comparison of pSer129+ neuron counts across weeks in the M1 region of mice injected with  $\alpha$ Syn PFFs in dorsal striatum ( $n = 1020$  sampled neurons, 10 slices across 5 mice, 2 slices per week data shown as mean  $\pm$  sem). In (**H**, **I**) red and blue traces correspond to the % of pSer129+ neurons in L2/3 and L5, respectively for each seeding strategy; one-way ANOVA with Bonferroni comparison test, \* $p < 10^{-2}$ , \*\* $p < 10^{-5}$ , data shown as mean  $\pm$  sem.

(L2/3) counterparts ( $12.54 \pm 6.26\%$ ,  $12.21 \pm 6.83\%$ ,  $3.69 \pm 2.11\%$ ; L2/3 of M1, M2, and S1 respectively). This was accompanied by a clear lack of pSer129- $\alpha$ Syn pathology in the dorsal striatum ( $2.48 \pm 2.22\%$ ) (Fig. 1C, D). In contrast, dorsal striatal seeding led to a strong pathological spread in the striatum (percentage of pSer129- $\alpha$ Syn+ neurons =  $26.58 \pm 12.74\%$ ), followed by a comparable pathological burden in L5 of both primary and secondary motor cortex (percentage of pSer129- $\alpha$ Syn+ neurons =  $15.74 \pm 10.09\%$ ,  $14.63 \pm 8.04\%$ ). Moreover, L4 and L2/3 of S1 showed a larger burden compared to cortically seeded ( $p < 0.001$ ), with a comparable spread in M1 L5 and M2 L2/3. (Fig. 1C, D). Notably, although the percentage of neurons that stained positive for pSer129- $\alpha$ Syn was markedly greater in the L2/3 and L5 regions of mice injected in the cortex with  $\alpha$ Syn PFFs versus  $\alpha$ Syn monomer (monomer control), we observed no difference in overall somatic density across layers 12 weeks after injections between the two groups of animals. (Fig. 1E, F) (L2/3 monomer:  $2093 \pm 750$ , L2/3 PFF:  $2072 \pm 1128$ , L5 monomer:  $2134 \pm 417$ , L5 PFF:  $2141 \pm 864$  neurons per  $\text{mm}^2$ ;  $p = 0.35$ ,  $0.22$  for layer 2/3 and layer 5 respectively,  $n = 4$  mice), suggesting a lack of early-stage cellular atrophy and death consistent with other reports<sup>5,17,51</sup>.

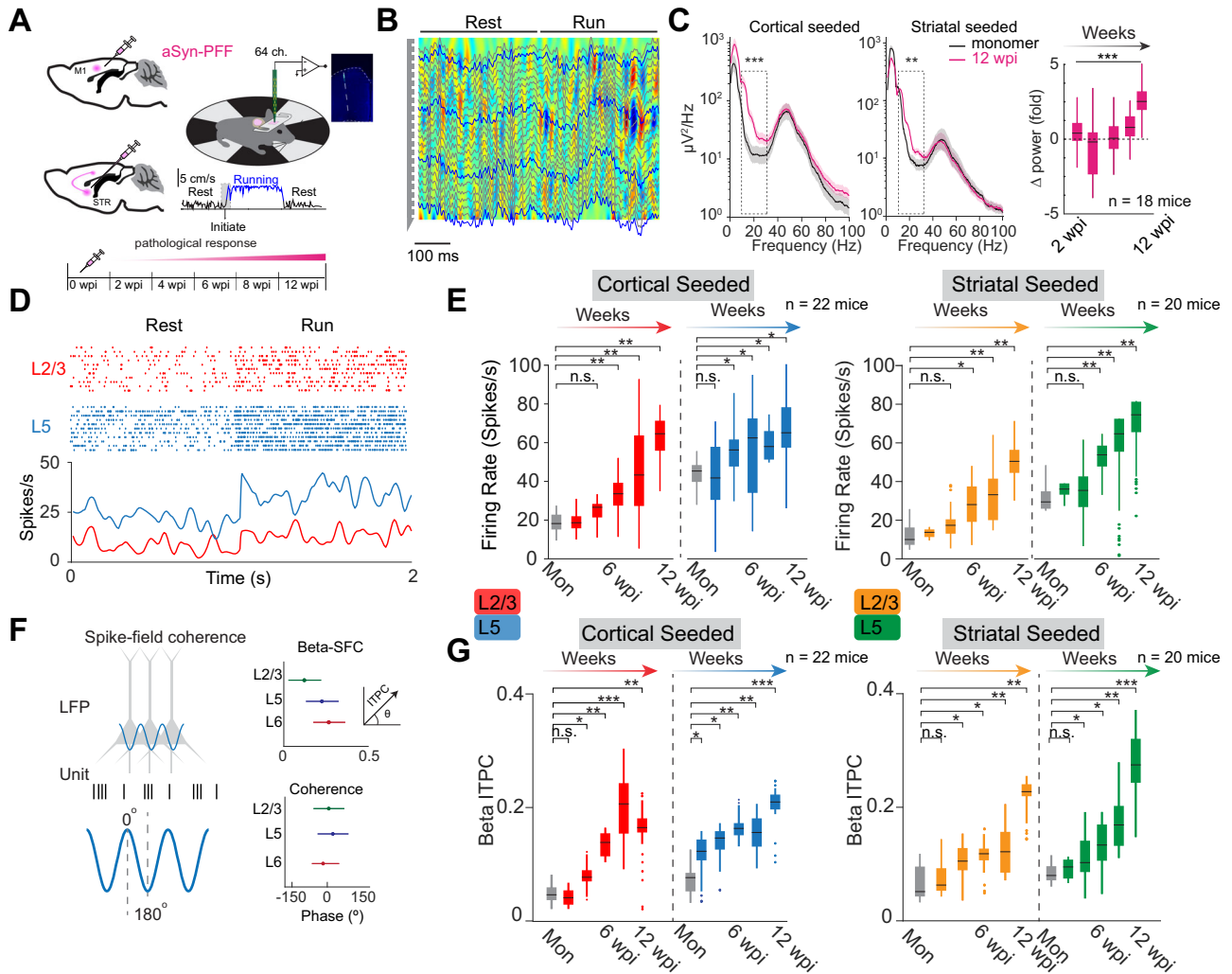
To characterize Lewy body spread across cortical layers over the 3 month period, we performed immunohistochemistry every 2 weeks of sections prepared from the primary motor cortex of mice injected with  $\alpha$ Syn PFFs in the cortex or striatum (see Methods). We observed a gradual increase in the burden of pS129- $\alpha$ Syn aggregates across all layers in M1 of animals injected in either location, with the highest density focused in L5 and subsequently spreading to L2/3 (Fig. 1G, see Supplementary Fig. S1 for additional slices with counterstaining). Notably, we observed that significant Lewy pathology (consisting of obvious, somatic Lewy-like inclusions) did not appear in the M1 L2/3 region of cortically seeded mice until 6 wpi (percentage of pSer129- $\alpha$ Syn+ neurons =  $10.7 \pm 3.51\%$ ;  $p < 0.001$ ) (Fig. 1H). Moreover, the appearance of pS129- $\alpha$ Syn aggregates was delayed (and the aggregate burden was reduced) in the M1 L2/3 region of striatally seeded mice compared to cortically seeded animals (percentage of pSer129- $\alpha$ Syn+ neurons =  $6.49 \pm 2.13\%$  versus  $13.22 \pm 5.15\%$ ,  $p < 0.01$ ) at 12 wpi (Fig. 1I). These findings suggest that the gradual spread in  $\alpha$ Syn pathology differed slightly in cortically versus striatally seeded mice, but still consisted of a strong retrograde transynaptic progression of Lewy-like inclusions in M1, ending in similar pathological staging across layers. The continued propagation of  $\alpha$ Syn aggregates across different brain regions also led to a heavy pathological load throughout the brain 12 months post-injection (Supplementary Fig. S1d). In addition, we observed little to no observable behavior changes (Supplementary Fig. S2a–d) involving locomotion in the case of striatally and cortically seeded mice within the first 3 months of  $\alpha$ Syn-PFF seeding, whereas clear behavioral deficits were exhibited by animals in both groups 12 months post-injection (Supplementary Fig. S2e–i) (24 cortically injected and 22 striatally injected mice).

### Electrophysiology reveals layer-specific vulnerabilities as a result of $\alpha$ Syn pathology

To investigate the effects of differing  $\alpha$ Syn pathology on recurrent motor circuitry during active behavior<sup>52</sup>, we performed high-density electrophysiological recordings using silicon probes (see Methods) across layers of M1 in awake  $\alpha$ Syn PFF-seeded and monomer seeded mice. Locomotory movements are the behavioral correlates of active motor preparation and execution<sup>23</sup>. Thus, mice were habituated on a running wheel 1 week before experiments and engaged in bouts of sustained running during recordings (Fig. 2A) (see Methods). We generated a velocity-triggered window to quantify periods of active locomotion. Using spectral decomposition, we analyzed the local field potential (LFP) during this period and identified frequency bands of theta (8–12 Hz), beta (12–30 Hz), and gamma (30–90 Hz) activity

(Fig. 2B). We observed a significant increase in LFP power across these frequency bands in mice injected in the cortex or striatum with  $\alpha$ Syn PFFs versus age-matched  $\alpha$ Syn monomer (monomer control) at 12 weeks post-PFF injection (wpi) for both treatment groups (Fig. 2C, left, middle,  $n = 5$  mice, Cortical Monomer  $n = 2$ , Cortical PFF  $n = 3$ , Striatal Monomer  $n = 2$ , Striatal PFF  $n = 3$ ). Specifically, beta LFP power significantly increased by 12 wpi, highlighting a distinct change in cortical beta-band activity induced by  $\alpha$ Syn pathology (Fig. 2C, right). During this 3-month period, no significant motor dysfunction was observed (Supplementary Fig. S2a–d). However, changes in exploratory behavior, rearing, and motor control were observed 6 months and 1 year after injection (Supplementary Fig. S2e–i). Investigating LFP alone reflects network-wide changes induced by  $\alpha$ Syn pathology and does not capture the precise temporal contribution of underlying circuits. We thus measured putative excitatory single units across cortical layers and found that these showed rate-dependent modulation during locomotion (Fig. 2D). Analysis of putative excitatory single units revealed a gradual increase in mean firing rates in the brains of cortically and striatally seeded animals (Fig. 2E). However, while initially the spike rates in L5 were higher in comparison to L2/3, the rate of increase in L2/3 was rapid and steep towards the 12 week mark, suggesting an entrainment and amplification by L5. Specifically, cortical seeding drove a substantial increase in the mean firing rates of L2/3 neurons compared to L5 neurons during active locomotion (Two-way unbalanced ANOVA with Bonferroni comparison test,  $p < 0.01$ ;  $N = 958$  units (43 well-defined units per animal per timepoint);  $n = 22$  Mice CTX; see figure legend for exact mice across weeks) (Fig. 2E, left). In striatally seeded animals, we observed a similar increase in firing rates over the 12 week period, but in this case, L2/3 seemed to linearly track the trend in L5 (Fig. 2E, right). Importantly, in comparison to cortical seeding, a slightly lower response of mean firing rates across both L2/3 and L5 neurons (Two-way unbalanced ANOVA with Bonferroni comparison test,  $p < 0.01$ ;  $N = 709$  units (35 well-defined units per animal per timepoint);  $n = 20$  striatally seeded mice in total; see figure legend for exact mice across weeks) was observed. Notably, under all monomer conditions, where monomeric mice at 2 and 12 weeks after injection were pooled, firing rates of L2/3 pyramidal neurons were less than L5 (average firing rate during locomotion =  $19.3 \pm 5.1$  L2/3 spikes/s;  $39.4 \pm 6.3$  L5 spikes/s;  $p < 0.01$ ).

Firing patterns of neurons are heavily influenced by phase synchronization to the local-field potential (LFP). Therefore, we employed spike-field coherence<sup>53,54</sup> (Fig. 2F, Supplementary Fig. S3a) to couple LFP and single-unit activity across cortical layers (see Methods). We focused on beta LFP and calculated the spike-field coherence and phase across cortical depths (Fig. 2F). To quantify the preference of single units to beta LFP, we looked at the beta inter-trial phase coherence (ITPC)<sup>53,54</sup>. Under monomer conditions pooled at 2 and 12 weeks, L5 neurons exhibited stronger beta ITPC than L2/3 neurons (ITPC during locomotion =  $0.12 \pm 0.03$  for L5 and  $0.06 \pm 0.02$  for L2/3;  $p < 0.05$ ), in line with the theory that beta predominantly targets deep cortical layers<sup>45,55</sup> (Fig. 2F, Supplementary Fig. S3b). Strikingly, in mice with cortically seeded  $\alpha$ Syn pathology, we observed a shift in beta ITPC across weeks; superficial L2/3 ITPC was lower than L5 but then rapidly changed alongside L5 beginning at the 6-week time point, suggesting a gradual entrainment (Fig. 2G, left). However, in striatally seeded mice, increases in beta ITPC were more apparent in L2/3 at earlier timepoints relative to monomeric injections. This was followed by a larger increase of L5 ITPC in comparison to L2/3 12 weeks after seeding (Two-way unbalance ANOVA with Bonferroni comparison test,  $*p < 0.01$ ,  $**p < 0.001$ ,  $***p < 0.0001$ ,  $n = 22$  Mice CTX,  $n = 20$  Mice STR) (Fig. 2G, right). Such gradual changes in ITPC were primarily relegated to the beta and not gamma nor theta-bands for both seeding locations (Supplementary Fig. S3c–f). Importantly, such changes in spike rates or beta-ITPC were not observed when we compared monomeric cortical and striatal injected mice to un-injected mice under the same behavior

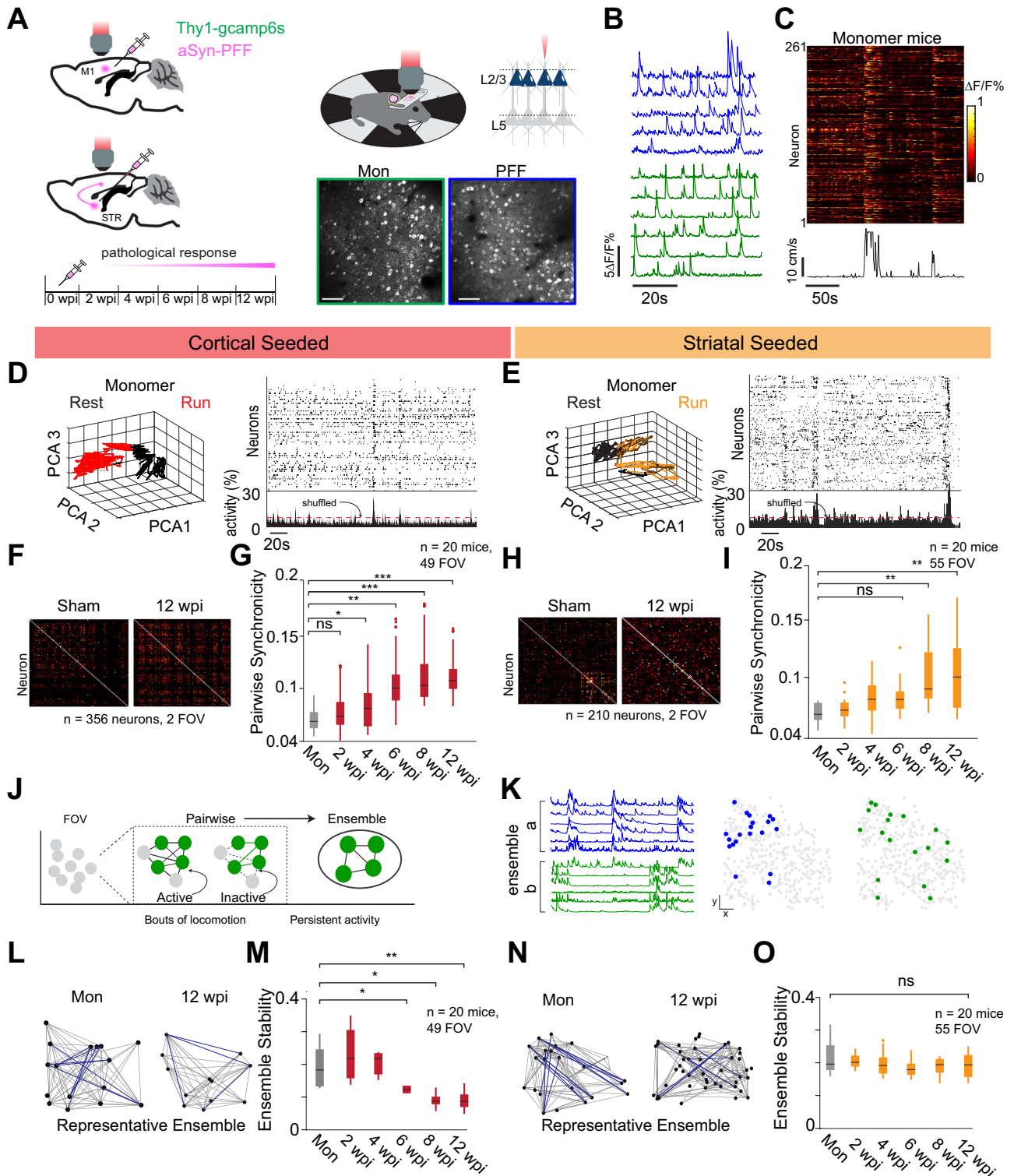


**Fig. 2 | Cortical and striatal seeding drives hyperexcitability and distinct beta coupling in primary motor cortex.** **A** Schematic of experimental setup. **B** Representative CSD and wideband (1–100 Hz) LFP during initiated and running epochs averaged across trials from a single mouse ( $n = 407$  trials). **C** (Left, middle) Power distribution of broadband LFP across cortical and striatal pathological weeks ( $n = 1083$  trials,  $n = 5$  mice, Cortical Monomer  $n = 2$ , Cortical PFF  $n = 3$ , Striatal Monomer  $n = 2$ , Striatal PFF  $n = 3$ ). Note the shaded region corresponding to beta band (13–30 Hz). (One-way Anova with Bonferroni comparison test,  $n = 1040$  trials). (Right) Changes of integrated beta LFP across weeks in cortically seeded mice  $F(4, 19) = 5.35, p = 0.007$  ( $n = 18$  mice, 3, 4, 4, 3, 4 across each two-week time point, i.e. every two weeks starting at week 2 and ending at week 12, respectively). **D** Representative single unit response of L2/3 and L5 pyramidal neurons during rest and running periods from a single monomer mouse ( $n = 11$  trials). **E** Firing rate during active running for each layer across weeks in mice injected with  $\alpha$ Syn PFFs in the cortex (left)  $F_{\text{Mon}^{\text{PFF}}}(5, 20) = 9.83, p = 3.7 \times 10^{-9}$  or striatum (right)  $F_{\text{Mon}^{\text{PFF}}}(5, 20) = 15.12, p = 3.3 \times 10^{-6}$ . (Two-way unbalanced ANOVA with Bonferroni

comparison test,  $*p < 0.01, **p < 0.001, ***p < 0.0001$ ). **F** Schematic of spike field coherence (left). Beta spike-field coherence and phase calculated across depth ( $n = 2$  cortical monomer-injected mice, 67 trials) (right). **G** Beta ITPC across depths and pathological weeks between monomer and cortical seeded PFF (left)  $F_{\text{L23Mon}^{\text{L23PFF}}}(5, 20) = 19.42, p = 8.66 \times 10^{-9}, F_{\text{L5Mon}^{\text{L5PFF}}}(5, 20) = 63.77, p = 1.42 \times 10^{-9}$  and striatal seeded mice (right),  $F_{\text{L23Mon}^{\text{L23PFF}}}(5, 18) = 17.6, p = 0.0024, F_{\text{L5Mon}^{\text{L5PFF}}}(5, 18) = 8.99, p = 0.0004$ . (Two-way unbalanced ANOVA with Bonferroni comparison test,  $*p < 0.01, **p < 0.001, ***p < 0.0001$ ). Data shown on box plots with 25th, median and 75th percentiles and with whiskers indicating min and max values,  $n = 22$  cortically injected mice in total. Monomer Cortex  $n = 4$  mice (2 mice at wpi and 2 mice at 12 wpi), Cortical PFF  $n = 18$  mice (3, 4, 4, 3, 4 PFF mice across each two-week time point, i.e. every two weeks starting at week 2 and ending at week 12, respectively).  $n = 20$  striatally injected mice in total. Striatal monomer  $n = 4$  mice (2 mice at 2 wpi and 2 mice at 12 wpi) Striatal PFF  $n = 16$  mice (3, 3, 3, 3, 4 PFF animals across each two-week time point, i.e. every two weeks starting at week 2 and ending at week 12, respectively), wpi weeks post-injection.

(Supplementary Fig. S4a–d). Remarkably, the disruption observed under both seeding strategies in overall spike-LFP coherence emerged earlier than changes in broadband LFP power (Fig. 2C, right), indicating that the ITPC is a much more sensitive metric of pathology progression compared to gross LFP changes and can serve as a key marker of  $\alpha$ Syn-induced circuit alterations in the motor cortex<sup>16</sup>. The rate of change in beta-ITPC across weeks shows that the measure carries within it a seeding location-dependence. Given that L5 recurrently modulates L2/3 via monosynaptic projections<sup>25,27</sup>, specifically via Layer 5A, it suggests that the L5 to L2/3 coupling might be particularly vulnerable to  $\alpha$ Syn aggregates and a potential driver of L2/3 dysfunction.

The differences in the beta-ITPC profiles between animals injected with PFFs in the cortex versus striatum could be explained as follows: (i) propagation of pathology must occur over a greater distance to impact L5 when administered in the striatum; and (ii) PFF injection at the two sites would be expected to affect non-identical circuit pathways, due to difference in seeding sites (Fig. 1D, H, and I), across the L5->L2/3 axis<sup>26</sup>, thereby leading to differences in ITPC perturbation. Finally, given that coherency of single units to LFPs is critical for both local and global synchronization of neuronal activity, the circuit alterations observed over the prodromal-phase could have significant implications for information processing and communication between



different brain regions and cortical networks<sup>16,53,56</sup> in pathologically similar forms of disease.

**Superficial network hyperactivity patterns are pathway-specific**  
 Fundamentally, networks of cortical pyramidal neurons in L2/3 exhibit sparse representation of sensory stimuli, allowing the cortex to non-linearly represent sensory information in a distributed and efficient manner<sup>57,58</sup>. Based on the observation that αSyn PFF seeding in the cortex or striatum led to distinct beta-band ITPC profiles that progressively amplified in superficial layers over 12 weeks, we examined

the extent to which this trend impacted cellular activity patterns and functional ensemble dynamics. To address this, we employed two-photon (2P) calcium imaging with GCaMP6s expression under the control of the Thy-1 promoter (see Methods) in separate cohorts of mice (of either sex) injected with αSyn PFFs in the cortex or striatum, respectively (Fig. 3A, left). We imaged population activity of L2/3 pyramidal neurons in the motor cortex across 12 weeks and mapped their spatial and temporal correlations as a function of cortical and striatal seeding<sup>19,20</sup>, a feature not captured in our silicon probe recordings (Fig. 3A, right) (*n* = 9034 L2/3 pyramidal neurons from 20

**Fig. 3 | Population calcium imaging reveals distinct differences in synchronicity and ensemble activity.** **A** Schematic of experimental setup. **B** Denoised calcium transients in GCaMP6s expressing L2/3 neurons of monomer (green traces, bottom) and PFF-injected mice (blue, top). **C** Running evoked fluorescent activity and encoder trace in monomer animal under a representative running trial. **D–I** Pairwise analysis of data obtained under cortical and striatal PFF seeding models. **D, E** Low-dimensional neural state space representing transitions of rest and active running states 12 weeks post seeding in primary motor cortex or striatum of representative cortical monomer mouse (left) ( $n = 33$  trials and  $n = 23$  trials, respectively) and calcium event (right). Each row represents a neuron and each raster point represents a measure of neural activity. Percentage of neurons coactive within a given frame (bottom inset, right plots in **D, E**). Highly coactive regions above chance values are marked by a dashed line (See methods and Supplementary Fig. S5). **F–I** Pairwise analysis of neuronal activity. **F, H**  $\alpha$ Syn PFF injection into motor cortex M1 (**F, G**) and dorsal striatum (**H and I**). Representative pairwise response from mice injected in the cortex (**F**) or striatum (**H**) with monomer control or  $\alpha$ Syn PFFs 12 weeks post injection (‘12 wpi’) ( $n = 566$  neurons, 4 mice). **G, I** Pairwise synchronicity values of evoked  $\text{Ca}^{2+}$  population responses across cortical. **G**  $F_{\text{Mon}^{\text{PFF}}}(5, 19) = 10.81, p = 5.18 \times 10^{-6}$  and striatal (**I**)  $F_{\text{Mon}^{\text{PFF}}}(5, 18) = 13.11, p = 1.043 \times 10^{-7}$ ,  $\alpha$ Syn pathology compared to shuffled data (See methods).  $n = 9034$  neurons from 20 cortically injected mice total. Cortical monomer  $n = 4$  (2 mice at 2 wpi and 2 mice at 12 wpi), Cortical PFF mice  $n = 16$  mice (3, 3, 3, 3, 4 PFF animals

across each 2-week time point, i.e. every 2 weeks starting at week 2 and ending at week 12, respectively).  $n = 7509$  neurons from 20 striatal mice total, Striatal monomer  $n = 4$  (2 mice at 2 wpi and 2 mice at 12 wpi). Striatal PFF mice  $n = 16$  mice (3, 3, 3, 3, 4 PFF animals across each 2-week time point, i.e. every 2 weeks starting at week 2 and ending at week 12, respectively).  $*p < 0.01, **p < 0.001, ***p < 0.0001$ , Two-way unbalance ANOVA with Bonferroni correction. **J** Schematic of evolved cortical networks based on pairwise activity and the emergence of cortical ensembles that are functionally dependent on time (FOV, field of view). **(K)** Representative traces of the two most stable ensembles across the behaving epoch from a single mouse (left) and their spatial arrangement (right). **L–O**  $\alpha$ -Syn PFF injection into motor cortex M1 (**L, M**) and dorsal striatum (**N, O**). **L, N** Ensemble of neurons after mathematically decomposing data from binarized matrix under cortical and striatal seeding. **M, O** Probability of ensemble recruitment response across  $\alpha$ Syn pathology under cortical  $F_{\text{Mon}^{\text{PFF}}}(5, 19) = 13.62, p = 1.37 \times 10^{-5}$ , and striatal seeding model  $F_{\text{Mon}^{\text{PFF}}}(5, 18) = 0.24, p = 0.94$  ( $*p < 0.01, **p < 0.001, ***p < 0.0001$ , two-way unbalanced ANOVA with Bonferroni correction).  $n = 1950$  ensembles from 20 cortically seeded mice, Cortical monomer  $n = 4$  mice (2 mice at 2 wpi and 2 mice at 12 wpi), Cortical PFF  $n = 16$  mice,  $n = 1871$  ensembles from 20 striatally seeded mice, Striatal monomer  $n = 4$  mice (2 mice at 2 wpi and 2 mice at 12 wpi), Striatal PFF  $n = 16$  mice, data shown on box plots with 25th, median and 75th percentiles and with whiskers indicating min and max values, wpi weeks post-injection.

cortically seeded mice; see figure legend for exact Ns across weeks,  $n = 7509$  L2/3 pyramidal neurons from 20 striatal-seeded mice; see figure legend for exact mice across weeks).

We quantified  $\text{Ca}^{2+}$  dynamics in motor cortex L2/3 (see Methods), across both PFF and monomer-injected mice (monomer experiments performed and pooled at 2- and 12 week timepoints only) (Fig. 3B, C) during spontaneous bouts of running. Because of the relatively large number of neurons imaged, a clear transition of cortical dynamics could be observed during locomotion, which could be projected into a lower-dimensional state-space (Fig. 3D, E). We then extracted pairwise correlations (based on synchronized  $\text{Ca}^{2+}$  activity) triggered by spontaneous bouts of locomotion (see Methods). Using this metric, we observed a gradual yet progressive increase in pairwise correlations for mice injected with  $\alpha$ Syn PFFs in either seeding location (Fig. 3F–I). Notably, pairwise synchronicity extracted from the covariance maps (Fig. 3F, H) obtained for cortically seeded mice (Fig. 3G) reflected a marked difference as early as 4 weeks post-injection in comparison to striatally seeded, where a difference was seen only 8 weeks after PFF administration (Fig. 3I). These differences were significantly above monomer-injected mice for both cortically ( $n = 184$  neurons, 33,850 pairwise connections (per FOV);  $n = 20$  cortically seeded mice in total; see figure legend for exact Ns across weeks;  $p < 0.001$ ; Two-way unbalance ANOVA with Bonferroni comparison test) and striatally seeded mice ( $n = 140$  neurons, 19,600 pairwise connections (per FOV);  $n = 20$  striatally seeded mice in total; see figure legend for exact mice across weeks). While an increase in cortical pairwise activity alone is not surprising, our results highlight possible changes in cellular and circuit mechanisms critical for motor control and learning. Specifically, distinct pathway-specific hyperactivity patterns, observed from changes in spiking activity (Fig. 2E, G), can influence and alter computations requiring sparse L2/3 coding, for example, during motor learning<sup>59,60</sup>, in a manner that varies with pathology propagation across vulnerable pathways. Central to this line of thinking would be the disruption of neuronal ensemble dynamics, which we examine below.

While pairwise hyperactivity trends (Fig. 3G, I) are informative in conveying how any two neurons are coactive, we wondered if there was a starker difference in the activity patterns that could help differentiate pathway-specific progression. It is not clear, from average pairwise synchrony alone, how groups of 3 or more neurons coordinate activity within a dysfunctional network. Indeed, throughout locomoting bouts, a subset population of multiple coactive neurons, termed neuronal ensembles, emerged<sup>19</sup>. Ensemble dynamics reflect

flexible spatial emergences and can signify a mixed representation of sensory features (Fig. 3J)<sup>61</sup>. Importantly, stable reactivation of neuronal ensembles is a critical feature of healthy brains and has been linked to motor learning<sup>21,59,60</sup>. To identify these ensembles, we measured population activity as multi-dimensional vectors corresponding to the degree of similar temporal activity across populations of neurons (see Methods and Supplementary Fig. S5a–d). We searched this multi-dimensional vector (i.e., time) space to mathematically extract various ensemble combinations, which were projected onto a 2D Euclidean space to reveal their spatio-temporal structure (Fig. 3K).

To test whether specific features of ensembles were altered in response to  $\alpha$ Syn pathology, we quantified ensemble stability – the probability that neurons within an ensemble reactivate together (see Methods, Fig. 3L–O). The strength of co-activation of neurons within an ensemble, termed a connection, was critical to define the stability of the ensemble (Fig. 3L, N, Supplementary Fig. S5d). Interestingly, under cortical seeding we discovered a reduction in ensemble stability over the 12 week period in contrast to the progressive increase in pairwise synchronicity observed for cortically seeded mice (Fig. 3M) (Two-way unbalance ANOVA with Bonferroni comparison test,  $p < 0.001$ ;  $n = 40$  ensembles (per FOV);  $N = 49$  FOV;  $n = 20$  cortically seeded mice in total; see figure legend for exact mice across weeks). However, when we applied the same analysis to striatally seeded mice, we observed no such change in ensemble stability (Fig. 3O) (Two-way unbalance ANOVA with Bonferroni comparison test,  $p = 0.94$ ,  $N = 69$  ensembles (per FOV);  $N = 55$  FOV;  $n = 20$  striatally seeded mice in total; see figure legend for exact mice across weeks). A gradual reduction of ensemble stability implies that the probability for the same neurons within an ensemble to reactivate decreases across the pathological timeline. A comparison of pairwise network activity (synchronicity) and ensemble stability reveals a dichotomous relationship in mice with evolving  $\alpha$ Syn pathology that is only found in cortically (but not striatally) seeded animals. Notably, the change in ensemble stability reflected a difference in the number of functional ensembles that could be formed. For animals injected with  $\alpha$ Syn PFFs across both locations, we observed a significant decrease in the number of detected ensembles over time (Supplementary Fig. S5e, S5g). However, the number of functional connections was only different in mice subjected to cortical PFF seeding, suggesting a difference in temporal activity of cortical ensembles not found in striatally seeded animals. (Supplementary Fig. S5f, S5h). Importantly, throughout the 12 week period, there was no change in the number of cells or the number of cells per ensemble

being imaged across mice (Supplementary Fig. S5i–k), suggesting that the effects seen at the population level and the differences observed as a function of seeding location were due to local circuit impairments. This was further substantiated when we compared functional pairwise activity and ensemble stability of monomeric mice to uninjected GCaMP mice (Supplementary Fig. S6a–d.) which revealed similar pairwise and ensemble activity across mice. Overall, the presence of sparse yet strong pairwise synchronicity and optimal ensemble activity is a hallmark of a healthy L2/3<sup>20</sup>, and the phenomenon of systematic deviation in ensemble stability seems to only arise as a function of pathology.

Overall, the change in pairwise hyperactivity measured via two-photon calcium imaging in L2/3 (Fig. 3G, I) tracks the L5 to L2/3 entrainment observed in the electrophysiological measures (Fig. 2E and G) across weeks, eventually leading to seeding location-specific ensemble dysfunction in L2/3. Is there a possible connection between the hyperactivity and dysfunction observed in the superficial L2/3 layer and a pathology-induced effect in the L5 layer? This could be due to their recurrent connectivity. Additionally, the differences seen in ensemble stability may be due to pathway-specific vulnerabilities that are tied to L5 connectivity. Is there a way to identify such an effect in L5?

### Translaminar beta dynamics indicate distinct pathway-specific L5 vulnerability

The disruption of supragranular spatiotemporal neural ensemble dynamics may have significant implications for recurrent translaminar activity. Current understanding across the cortical column posits that the interaction between bottom-up sensory processing and top-down feedback mechanisms generates flexible sensory representations through the coordination of gamma-beta interactions (Fig. 4A), with gamma and beta oscillation predominantly generated by superficial and deep-layer neurons, respectively<sup>55</sup>. This coordination allows for efficient sensory processing under attentional demands<sup>32,55,62,63</sup>. Also, beta oscillations have been implicated in the synchronization of activity across neural ensembles, highlighting their crucial role in cortical processing<sup>30</sup>. Is there a delineating feature of beta that can match the superficial ensemble dynamics differences as a function of seeding locations? We posit that such a marker, possibly driven by L5, alongside ensemble dynamic differences in superficial layers will serve as a powerful handle to differentiate between symptomatically similar synucleinopathies

We analyzed the power distribution of beta and gamma across the cortical depth. Indeed, and consistent with previous studies on motor and working memory in non-human primates<sup>32,55,64,65</sup>, we observed that gamma amplitude was largest in the superficial layers (at depths up to 250  $\mu\text{m}$ , yellow trace), while the beta power occurred in deeper layers (at depths >450  $\mu\text{m}$ , blue trace) (Fig. 4B). Strikingly, in the brains of mice injected with  $\alpha\text{Syn}$  PFF, there was a modest increase in beta power distribution across both seeding locations.

Recent work has also revealed that beta LFP is not a continuous, sustained rhythm, but rather a transient surge in amplitude that occurs throughout individual trials via beta bursts<sup>47,48</sup> (Fig. 4C, D). These bursts have been linked to the initiation and maintenance of movement, as well as the suppression of unwanted movements<sup>29,66</sup>, and disruptions in beta burst activity have been associated with movement disorders such as Parkinson's disease<sup>41,49</sup>. Therefore, we next asked if moment-to-moment features of cortical beta can reflect changing translaminar dynamics in response to  $\alpha\text{Syn}$  pathology initiated from different seeding locations. We found that stochastic beta bursts in the motor cortex emerge as stereotypical waveforms across trials (Fig. 4D, E), typically lasting 40–100 ms, and fluctuating in waveform amplitude (Fig. 4F). These beta bursts span the beta frequency spectrum with a preference towards high beta (Fig. 4E). Here on, we term these stochastic bursts “beta events” ( $\beta$ -event)<sup>48</sup>. To quantify the effect

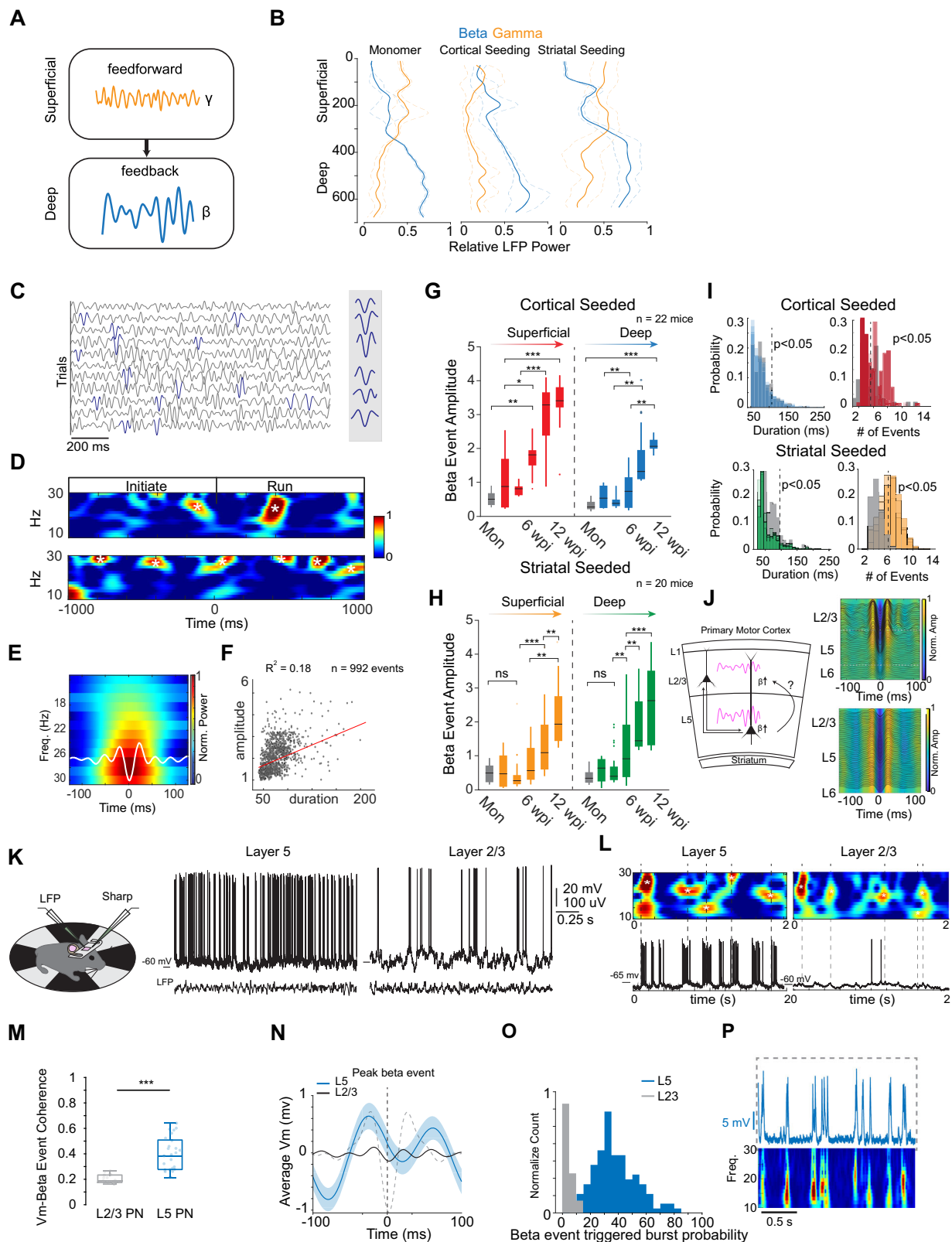
of evolving  $\alpha\text{Syn}$  pathology on  $\beta$ -events, we compared the  $\beta$ -event amplitude in the brains of mice injected with  $\alpha\text{Syn}$  PFFs in the cortex or striatum, as well as monomer controls (Fig. 4G–I) ( $n = 22$  cortically seeded mice in total,  $n = 20$  striatally seeded mice in total; see figure legend for exact mice across weeks). With cortical seeded pathology, as opposed to the broadband beta-ITPC measures across the pathological timeline (Fig. 2G) in which we found an initial increase in amplitude in L5 followed by an L2/3 entrainment, we found a significant increase in  $\beta$ -event amplitude in L2/3 of M1 that preceded a more modest increase in L5 towards the 12 week timepoint (Fig. 4G) (Two-way unbalance ANOVA with Bonferroni comparison test,  $p < 0.001$ ;  $1242 \pm 302$   $\beta$ -events per animal per timepoint);  $n = 22$  cortically seeded mice in total; see figure legend for exact mice across weeks). Under striatal seeding however, the  $\beta$ -event trend tracked the rate previously observed with beta-ITPC in which L5 increased similarly to L2/3 over the 12 week period (Fig. 4H) (Two-way unbalance ANOVA with Bonferroni comparison test,  $p < 0.001$ ;  $1312 \pm 401$   $\beta$ -events per animal per timepoint);  $n = 20$  striatally seeded mice in total; see figure legend for exact mice across weeks). We also noted significant differences in the number of  $\beta$ -events per trial but not the duration of the waveform across weeks and against monomer-injected animals for both seeding locations (Fig. 4I). Importantly, beta events were distributed along the cortical depth for both seeding locations, but with a stronger preference to initiate towards the superficial layers of M1 (Fig. 4J) in cortically seeding animals.

The results across Fig. 1 thru Fig. 4J lead to a distinct translaminar circuit marker of disease progression across different seeding locations. Specifically, as a function of seeding location (ie. seeding in deep layer cortex but not striatum), we observed both the firing rate and beta-ITPC (spike-field coherence) increase first in L5, subsequently entrain L2/3 (Fig. 2E, G), and this entrainment coincides with L2/3 excitatory ensemble instability (Fig. 3M, O). Importantly, we show that the  $\beta$ -power increase in superficial layers occurs only during pathology while primarily being generated in deeper layers of the cortex in monomer controls (Fig. 4B). Analyzing beta band activity in the form of stochastic moment to moment bursts at the single trial level revealed an increase in  $\beta$ -event amplitude in L2/3 in comparison to L5 under cortical seeding while the opposite was observed under striatal seeding with L5 and L2/3 modestly increasing together (Fig. 4G, H). Combined, these results strongly suggest that  $\beta$ -event generators share a common translaminar locus possibly intrinsic to L5 neurons<sup>48</sup> (Fig. 4J).

To elucidate the mechanism underlying  $\beta$ -event generation, we performed intracellular recordings, albeit separately, across putative L5 neurons and L2/3 neurons (at depths of 450–700  $\mu\text{m}$  and 100–250  $\mu\text{m}$ , respectively) in awake locomoting mice, while simultaneously measuring LFP recordings in superficial layers (at depths of 150–250  $\mu\text{m}$ ) (Fig. 4K). Spike-beta-event coherence measurements revealed a strong link between L5 bursting and extracellularly recorded beta-events, while L2/3 cells revealed a much weaker relationship (Fig. 4L, M) (Layer 5 =  $0.58 \pm 0.08$ ; Layer 2/3 =  $0.19 \pm 0.05$ ;  $p < 0.0001$ ;  $N = 10$  neurons). L5 neurons also displayed a more prominent membrane voltage oscillation compared to L2/3 neurons, indicative of a larger depolarization drive within the peristimulus  $\beta$ -event window (Fig. 4N, Layer 5 membrane potential change =  $0.81 \pm 0.12$  mV; Layer 2/3 membrane potential change =  $0.04 \pm 0.007$  mV;  $p < 0.01$ ;  $n = 6$  neurons). Additionally, L5 neurons were significantly more likely to exhibit intracellular bursts alongside extracellular beta events (Fig. 4O; 2406 beta events across all neurons). Examining the frequency components present in the intracellular recordings further revealed a broad beta-band response during L5 bursting (Fig. 4P). These findings together suggest that extracellular  $\beta$ -events may reflect the translaminar currents responsible for L5 burst mode signaling while conversely, a significantly lower effect is present for L2/3 in the primary motor cortex.

To further substantiate the link between L5 dendritic activity and the generation of translaminar  $\beta$ -events we performed biophysical





modeling using morphologically realistic L5 neurons (Supplementary Fig. S7a, see Methods). With excitation inputs (driven at Poisson rates) across the basal and distal apical dendrites,  $\beta$ -events were indeed generated in the presence of large apical dendritic currents denoted by a somatic burst (Supplementary Fig. S7b). Strikingly,  $\beta$ -event amplitude was observed to be severely reduced under conditions of NMDA and voltage-gated  $\text{Ca}^{2+}$  channel activity block along the L5 apical

dendrite (Supplementary Figs. S7d–f). We further verified this experimentally via application of baclofen, a  $\text{GABA}_B$  agonist, along the calcium rich zone (depth of 150–250  $\mu\text{m}$ ) of L5 neurons (see Methods), which reduced  $\beta$ -event power significantly (Supplementary Fig. S8a, S8b), while application of NMDA agonist (see Methods) along the same region reduced the  $\beta$ -event rate appreciably (Supplementary Fig. S8c, S8d). Beta event dynamics we observed experimentally were largely

**Fig. 4 | Emergence and dysfunction of  $\beta$ -events.** **A** Potential mechanism of narrow-band beta LFP information flow. Beta LFP is modulated by deep layer cortical activity via feedback while gamma dominates in superficial layer via feedforward inputs. **B** Example response of beta/gamma LFP power during voluntary running in monomeric versus  $\alpha$ Syn PFF animals. **C** Filtered beta LFP across velocity triggered trials. Blue traces correspond to putative bursts of beta transients with compiled waveforms of  $\beta$ -events ( $n = 11$  trials). **D** Examples of peristimulus beta band activity across two trials. Asterisks mark detected  $\beta$ -event waveforms. **E** Exemplar  $\beta$ -event spectrogram. **F**  $\beta$ -event amplitude and duration in representative recording of one monomer mouse ( $n = 992$  events). **G, H, Z**-scored  $\beta$ -event amplitude compared to monomeric control animals (indicated by gray box) across depth and pathological timeline for cortical seeded mice (**G**)  $F_{\text{supMon}^{\text{PFF}}}(5, 20) = 8.78, p = 3.15 \times 10^{-7}, F_{\text{deepMon}^{\text{PFF}}}(5, 20) = 13.11, p = 8.76 \times 10^{-11}$  and striatal seeded mice (**H**)  $F_{\text{supMon}^{\text{PFF}}}(5, 18) = 6.78, p = 0.0002, F_{\text{deepMon}^{\text{PFF}}}(5, 18) = 7.04, p = 0.0002$  ( $*p < 0.01, **p < 0.001, ***p < 0.0001$ , two-way unbalanced ANOVA with Bonferroni correction); Cortically seeded:  $1242 \pm 302$   $\beta$ -events per animal per timepoint.  $n = 22$  cortically injected mice in total. Cortical monomer  $n = 4$  mice (2 mice at 2 wpi and 2 mice at 12 wpi), Cortical PFF mice  $n = 18$  mice (3, 4, 4, 3, 4 animals across each 2-week time point, i.e. every 2 weeks starting at week 2 and ending at week 12, respectively). Striatal seeded:  $1312 \pm 401$   $\beta$ -events per animal per timepoint.  $n = 20$  striatally injected mice in total, Striatal monomer  $n = 4$  mice (2 mice at 2 wpi and 2 mice at 12 wpi), Striatal PFF mice  $n = 16$  mice (3, 3, 3, 3, 4 PFF animals across each 2-week time point, i.e. every 2 weeks starting at week 2 and ending at week 12, respectively), wpi weeks post-injection, data shown as median with first

and third quartiles. **I** Histogram distribution of  $\beta$ -event duration and number of events per trial across weeks for cortical (top) and striatal (bottom) seeded mice. Dashed line indicates significance in distribution compared to shuffled surrogates. Distribution of beta power and duration show high variability of  $\beta$ -event behavior. **J** Schematic of proposed emerging changes in narrow-band beta LFP dynamics. Aberrant deep layer beta dynamics may drive superficial beta dynamics via feedback (left). Normalized amplitude of  $\beta$ -event profiles for two example mice of cortically seeded (top) and striatally seeded (bottom) with corresponding depth. **K** Experimental schematic of dual LFP-Intracellular recordings (right). Intracellular recording from putative L5 and L2/3 pyramidal neurons in primary motor cortex (top trace). Wideband filtered LFP (4–40 Hz) (bottom trace). **L** LFP beta spectrogram and spiking response of L5 and L2/3 pyramidal neuron. Detected  $\beta$ -events is noted by white asterisk. **M** Vm and extracellular  $\beta$ -event-triggered coherence for L2/3 and L5 pyramidal neurons. ( $n = 10$  neurons, 7 L5 neurons, 3 L2/3 neurons,  $p = 0.00045, ***p < 0.0005$ , one-sided Kruskal-Wallis, data shown on box plots with 25th, median and 75th percentiles and with whiskers indicating min and max values). **N** Extracellular  $\beta$ -event triggered subthreshold response of L5 and L2/3. Average  $\beta$ -event waveform (dash line), average subthreshold response from L5 pyramidal neurons (blue trace),  $n = 7$  neurons, and subthreshold response to layer 2/3 neurons (black trace,  $n = 3$  neurons). **O**  $\beta$ -event triggered AP bursting probability from L5 and L2/3 pyramidal neurons ( $N = 2406$   $\beta$ -events across 243 trials,  $p = 5.5 \times 10^{-45}$ , Kruskal-Wallis). **P** Intracellular-frequency response during bursting of L5 pyramidal neuron.

absent when we performed the same set of simulations on biophysically modeled L2/3 neurons (Supplementary Fig. S9a–e). Notably, the translaminal profile and beta event amplitude were diminished compared to L5 neurons (Supplementary Fig. S9d, S9e,  $p < 0.005$ , Mann-Whitney U-test). These data suggest that NMDA channel function may be implicated in driving changes of beta event dynamics across a pathological timeline. To assess this, we conducted a phosphoproteomic analysis of the sensorimotor cortex in  $\alpha$ Syn PFF versus monomer injected (monomer control) animals, which revealed differences in levels of phosphoproteins involved in NMDA function and localization (Supplementary Fig. S10). Specifically, we found increased Grin2b phosphorylation in the cortices of mice injected intrastrially with  $\alpha$ Syn PFFs versus monomer, which we further confirmed with immunohistochemical analysis (Supplementary Fig. S11).

### $\beta$ -events power as a function of pathology coincides with aberrant dendritic excitability

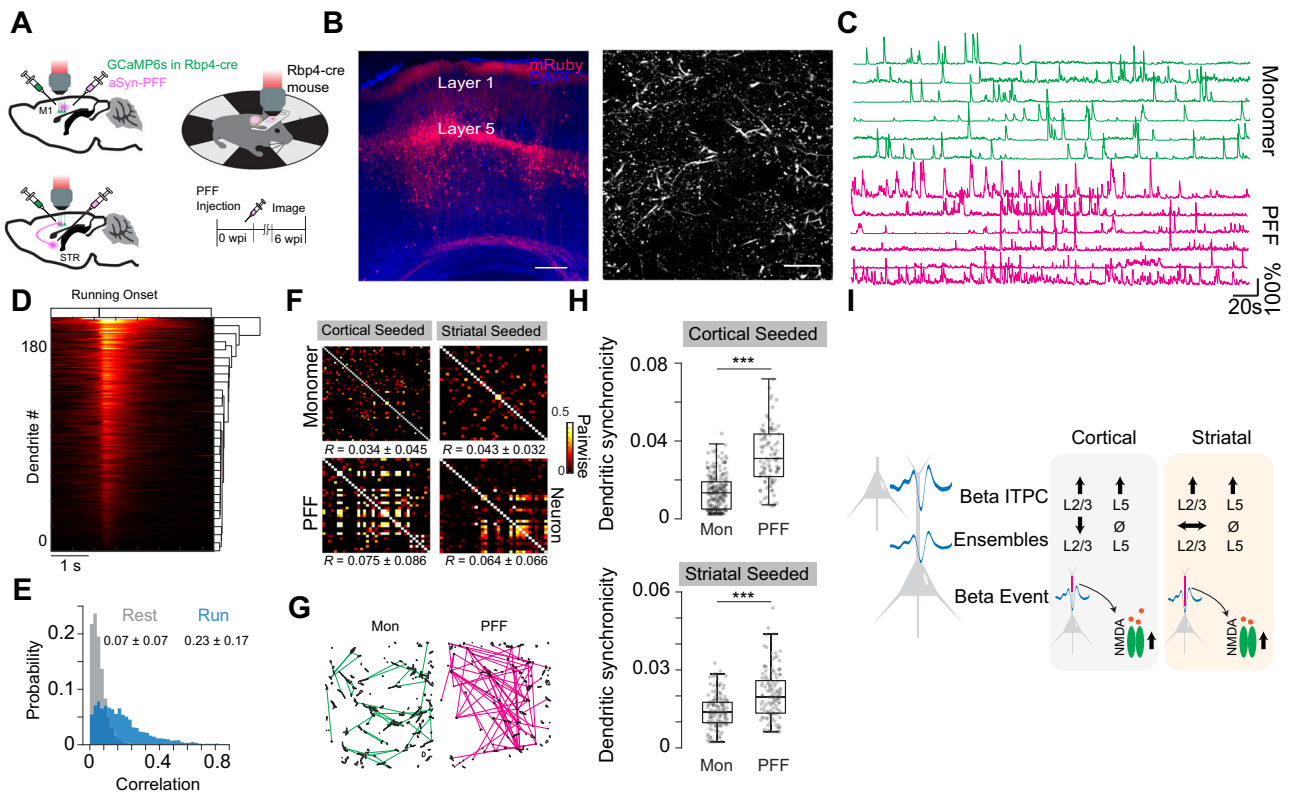
Finally, to corroborate that L5 dendritic activity is indeed aberrant in vivo during high  $\beta$ -event power, we performed  $\alpha$ Syn PFFs seeding across both dorsal striatum and in the primary motor cortex in *Rpb4-cre* mice which expressed cre-flox reporter primarily in L5 pyramidal neurons (Fig. 5A, B). Viral injection of *GCaMP6s-mRuby* allowed for structural reconstruction and functional imaging of L5 tuft dendrites for mice under monomeric and  $\alpha$ Syn PFF pathology under cortical or striatal seeding (Fig. 5C). We imaged L5 tuft dendrites (Fig. 5B, C) (at depths 40  $\mu\text{m}$ –150  $\mu\text{m}$  from the surface) in all 4 cohorts of mice after 6 weeks post-seeding (based on the timeline of L5 dynamics shown in Fig. 2 and Fig. 4) under locomotion (Fig. 5D). L5 tuft dendrites showed clear differences in pairwise population activity across dendrites during rest and locomotory states (rest:  $0.07 \pm 0.07$ ; run:  $0.23 \pm 0.17$ ;  $p < 0.001$ ;  $n = 821$  dendrites; 5 monomer mice) (Fig. 5E). When comparing dendritic populations,  $\alpha$ Syn PFF injected mice exhibited higher pairwise synchronicity than monomer mice (Fig. 5F, G) across the four treatment groups. Notably, distinct and significant changes in L5 dendritic connectivity (Fig. 5H, top, denoting synchronicity) were observed in mice injected with  $\alpha$ Syn PFF in the cortex (monomer:  $0.014 \pm 0.01$ , PFF:  $0.034 \pm 0.02, 3264$  dendritic segments;  $p < 0.001$ ;  $n = 21$  FOVs across 7 mice cortically injected mice) and striatum (monomer:  $0.0137 \pm 0.006$ , PFF:  $0.02 \pm 0.01, 2939$  dendritic segments;  $p < 0.001$ ;  $n = 19$  FOVs across 6 striatally injected mice), with the striatal PFF seeding showing a slightly less but still significance change in

pairwise connectivity compared to cortical PFF seeding ( $p < 0.001$ , one-way ANOVA, 8 mice) (Fig. 5H, bottom). These data support our observation that pathological changes in  $\beta$ -events are closely tied to L5 dendritic aberrancy. Moreover, there is a significant difference in the dendritic activity patterns evoked by  $\alpha$ Syn PFF seeding at different locations, suggesting that L5 pyramidal neurons may play a critical role in differences seen across the translaminal circuit during the evolution of  $\alpha$ Syn pathology.

Collectively, our results suggest that burst-mode activity in L5 neurons that is mediated by NMDA and  $\text{Ca}^{2+}$  signaling is a crucial element in the creation of  $\beta$ -events. Additionally, it appears that the abnormal phosphorylation of NMDA-related proteins could be a significant factor in the increased amplification of  $\beta$ -event power that is indicative of dendritic abnormalities in the local cortical circuits of the sensorimotor cortex in mice with  $\alpha$ Syn pathology. The differences in  $\beta$ -event amplitude between seeding locations further suggest that L5 dendritic channels are a vulnerable target, which not only helps generate a pathway-specific translaminal signature of disease progression (Fig. 5i), but could also serve as therapeutic targets for treating synaptopathies.

## Discussion

In this study, we introduced  $\alpha$ Syn PFFs into the dorsal striatum and deep-layer motor cortex of mice, resulting in the formation of inclusions similar to those found in the brains of individuals with PDD and DLB<sup>11,12</sup>. Over a period of 3 months (Fig. 1), we then examined the motor cortical activity patterns in awake locomoting mice and elucidated how circuit patterns reveal distinct and differentiable biomarkers. We discovered that there is a competition between cortical L2/3 and L5 excitatory activity in terms of phase entrainment to beta frequencies (Fig. 2), but not gamma or theta, despite the similarities in spread of Lewy pathology induced by PFF injection across the two sites. Specifically, with cortical seeding, L5 activity initially dominates, but L2/3 subsequently outpaces L5 beta-phase-entrainment. With striatal seeding, we found that L2/3 entrainment by L5 in the cortex is weaker in comparison. We also observed that cortical seeding destabilizes superficial L2/3 ensembles (Fig. 3), which is not the case with striatal seeding. Importantly, this destabilization is reflected by changes in beta-band dynamics across superficial layers, and analyzing these dynamics in a moment-to-moment fashion reveals a seeding location-dependent increase in stochastic  $\beta$ -event dynamics (Fig. 4).



**Fig. 5 | Layer-5 dendritic dysfunction under  $\alpha$ Syn pathology is seeding-location dependent.** **A** Experimental schematic and injection strategy for dendritic imaging during  $\alpha$ Syn pathology. **B** Representative images of L5 dendrites in M1 motor cortex immunohistochemically (left) and in vivo calcium imaging (right) (scale bars = 100  $\mu$ m and 20  $\mu$ m, respectively). **C** Fluorescence traces of dendritic tuft activity for single field of view of monomer (green) and cortically seeded mice (magenta). **D** Average dendritic population response during motor initiation from example mouse ( $n = 182$  dendrites, 2 FOV from cortical monomer mouse). **E** Pairwise population response during rest and running states ( $n = 801$  dendrites, 7 FOV, 4 mice total, 2 mice for each monomer group). **F** Example pairwise plot for each pathological condition

( $n = 281$  dendritic segments, 4 mice). **G** Sørensen-Dice dendritic synchronicity map, observed for mice injected with monomeric  $\alpha$ Syn (Monomer) or  $\alpha$ Syn PFFs (note increased connectivity in the case of cortically seeded PFF animals). **H** Dendritic population synchronicity observed for mice in each treatment group during locomotion. ( $n = 13$  animals, Cortical monomer  $n = 3$  mice, Cortical PFF  $n = 4$  mice, Striatal monomer  $n = 2$  mice, Striatal PFF  $n = 4$  mice,  $*p < 0.01$ ,  $**p < 0.001$ ,  $***p < 0.0001$ , One-way ANOVA, data shown on box plots with 25th, median and 75th percentiles and with whiskers indicating min and max values). **I** Pathological outcomes of cortical and striatal seeding lead to differing cellular and subcellular responses.

Importantly,  $\beta$ -event dynamics was discovered to be tightly locked to apical L5 dendritic  $\text{Ca}^{2+}$  currents and aberrancies were reflected at this level of dendritic signaling (Fig. 5). Markedly, these distinct changes in cortical circuit function occurred before behavioral dysfunctions observed at later timepoints (Supplementary Fig. S2). The fact that beta dynamics are primarily driven by the feedback-layers in deep cortex<sup>55</sup>, a feature we clearly observed (Fig. 4B), and that top-down feedback to L5 neurons targets apical dendritic activity in the cortex<sup>25,67</sup>, suggests a mechanism in which top-down control of motor output might be a potent target of synuclein dysfunction during the prodromal phase.

$\alpha$ Syn pathology progressively spreads between anatomically connected regions and this spread may result in different stressors among distinct neuronal networks over time<sup>12,17,43</sup>. Here, we show that both striatal and cortical seeding of  $\alpha$ Syn PFFs leads to widespread inclusion formation across the cortex (Fig. 1A–F), although with distinct circuit hyperactivity patterns possibly governed by anatomical connectivity. Surprisingly, the nature of circuit dysfunction is at odds with previous reports. For example, acute application of  $\alpha$ Syn PFFs in vitro was shown to result in impaired synaptogenesis and reduced activity<sup>4,51</sup>, while in vivo injection of  $\alpha$ Syn PFFs in the striatum led to (i) cortical hyperactivity attributed to a net change in excitation-inhibition balance exacerbated by gabaergic neuron loss, but with minimal Lewy neurite formation in the cortex after 9 months<sup>17,68</sup>, or (ii) a decrease in synapse and spine density as well as spontaneous

excitatory post-synaptic current frequency in L5 of the somatosensory cortex<sup>51</sup>. Our results (Fig. 1G, H), show that the motor cortex is indeed able to exhibit marked Lewy pathology and that intricate recurrent circuits in vivo might actually amplify circuit dysfunction (Fig. 2. and Fig. 3), before full-scale neurodegeneration.

With high-density electrophysiological recordings (Fig. 2A, B) we observed distinct differences in spike-field entrainment, specifically in the beta-frequency (Fig. 2C–G), as a result of cortical versus striatal  $\alpha$ Syn-PFF seeding across over a 12 week period. Notably, a L5-dependent entrainment of spike-field coherence in L2/3 was observed in response to both cortical and striatal  $\alpha$ Syn-PFF seeding (Fig. 2E, G), but with distinct differences in rates of increase. This result has important implications within the context of recurrent cortical and cortical-basal ganglia circuits and strongly underscores cortical L5 neurons as a vulnerable node between differing pathological seeding strategies<sup>9,13,28</sup>. Studies in non-human primate models have proposed that beta-frequency dynamics are driven by deep-layer neurons reflecting top-down feedback, with superficial layers driving gamma-band activity reflecting feedforward signaling<sup>31,55</sup>—a result we recapitulated in this study (Fig. 4A, B). Moreover, we found that beta-frequency dynamics were switched (with greater  $\beta$ -event localization in superficial layers) and heavily modulated under both striatal and cortical  $\alpha$ Syn-PFF seeding, yet with distinct differences. What are the circuit metrics that enable such differences?

Cortical microcircuits are hierarchically organized through feedforward, feedback, and lateral connections<sup>13</sup>. In the primary motor cortex, the L2/3 → L5A/B connection is the strongest, accounting for one-third of the total synaptic current in the circuit, followed by the reciprocal L5A → L2/3 connection as the second largest<sup>25</sup>. Therefore, strong recurrent inputs between superficial and deep layers can likely amplify any modulation in excitability due to pathophysiology<sup>26</sup>. This is highly likely given that we observed, for both cortical and striatal seeding, an initial increase in L5 beta ITPC followed by entrainment in L2/3 (Fig. 2E, G).

The differences in spike-field entrainment between mice injected intracortically versus intrastrially could reflect the greater distance over which the propagation of pathology must occur to affect L5 when PFFs are administered in the striatum, or the likelihood that seed injection at the two sites influences non-identical circuit pathways across the L5 → L2/3 axis given where they are seeded (Fig. 1G–I). For example, Layer 5A (intratelencephalic neurons projecting to the striatum) and Layer 5B neurons (pyramidal tract neurons) will be impacted in different ways across the two seeding locations, thereby impacting L2/3 differently. Notably, under cortical seeding, both Layer 5A and 5B could have a direct insult, while under striatal seeding layer 5B could be delayed in comparison to L5A due to multiple synaptic delays.

In line with changes at the circuit level, cell-type specific inhibition can play a major role in shaping local pathological changes as a function of aSyn uptake. Recent studies have described the role of changes in excitation-inhibition balance in L2/3 as a critical foundation for optimal sensory coding<sup>61,69</sup>, including the role of somatostatin interneurons (SOM+) in controlling L2/3 beta synchrony<sup>70,71</sup>. Given that these adapting inhibitory interneurons receive a strong excitatory drive from both deep-layer pyramidal neurons and laterally from within L2/3<sup>69</sup>, αSyn PFF induced changes in L5 excitatory activity observed in this study could further amplify and alter superficial beta synchrony both via anterograde signaling (L5 → SOM+ and L5 → L2/3) and retrograde propagation (via PFF uptake by L2/3 axons)<sup>23,69</sup>. Future experiments should focus on disentangling such connections in the context of αSyn pathology.

By monitoring the average pairwise level (correlations) measured via calcium dynamics and (tabulated as the mean correlation between any two cells) in L2/3 of mice injected with αSyn PFFs at both seeding locations, we observed an apparent increase in network synchronicity (Fig. 3A–I) mirroring the rate of change in firing rates over the 12-week period observed with high-density electrophysiology probes (Fig. 2E and G). Yet, where these seeding strategies diverged was at the ensemble level (Fig. 3J, K), where local persistent multi-neuronal activity patterns were measured across time<sup>19</sup>. Cortical but not striatal seeding of αSyn PFFs revealed a systematic decrease in ensemble stability (Fig. 3L–O), highlighting the gradual inability for ensembles to reliably activate within their respective groups during sensorimotor behavior, which in this study was reaching a set speed of locomotion. Importantly, this decrease was not a simple consequence of single-neuron activity level changes or functional connectivity, since neither unit spike rate, beta spike-field coherence, nor pairwise correlation was a good predictor of ensemble stability. Instead, our results likely reflect a disorganization of interconnected neuronal dynamics relegated to cortical-originating pathology—synonymous with dementia with Lewy body staging. This disorganization may arise from a local loss of inhibition<sup>1,17,72</sup> or impaired inhibitory drive as previously mentioned, which can dynamically alter spatio-temporal activation patterns of multiple neurons in L2/3, thus influencing L5 dynamics<sup>23,25,55</sup>. Our results support a model in which the pervasiveness of pathology is not an indicator of how disease propagation sculpts multi-neuronal activity patterns but rather that the variability of αSyn pathology can impact specific vulnerable

recurrent synaptic pathways across the L2/3–L5 axis in a connectivity-dependent manner<sup>4,17,43</sup>, giving rise to distinct spatio-temporal activity patterns that amplify circuit dysfunction (Fig. 5I). The fact that the maximal change in spike-field coherence occurred at beta frequencies (Fig. 2C and G) suggests that top-down control of motor cortical dynamics is impacted in a pathway-specific manner possibly via amplification through the L5 → L2/3 recurrent circuit<sup>13,25,27</sup>, and could be an early-stage marker of disease progression across vulnerable cortical pathways<sup>9,12,73</sup>. This work also highlights the potential interplay of how cortical dysfunction can act as stressors to downstream target regions involved in motor function including the generation of aberrant beta oscillations in human patients.

Mechanistically, such beta dysfunction manifests in the form of transaminar stochastic β-events<sup>48</sup> (Fig. 4A–F) with amplitudes that grow with time over the prodromal phase (Fig. 4G–J). We show that these β-events are strongly modulated by L5 pyramidal neurons, specifically thick-tufted tract neurons and not L2/3 pyramidal neurons (Fig. 4K–P). This is not surprising, given that L5B thick-tufted pyramidal neurons receive strong contextual feedback across their distal dendrites from higher-order areas<sup>67,74</sup>, are strongly modulated by L5 SOM+ Martinotti cells<sup>69</sup>, and receive inputs from L5A neurons (which project to the striatum)<sup>25,26</sup>, L2/3 pyramidal neurons, and the thalamus<sup>24</sup>. The combination of these inputs can drive complex bursts in L5B neurons through glutamatergic and voltage-gated calcium channels<sup>74,75</sup>. The presence of NMDA-driven nonlinearities in the distal tuft of L5 dendrites plays a crucial role in dendritic electrogenesis and influences the regulation of calcium-induced bursting<sup>76</sup>, a potential vulnerability. Indeed, we observed increased dendritic coactivity across L5 neurons<sup>67,74,75</sup> in PFF-injected mice (Fig. 5A–G). Importantly, NMDA currents last about 50 ms–80 ms which is approximately one beta period<sup>76</sup>. Using a compartmentalized model (Supplementary Fig. S7, 9), pharmacological controls (Supplementary Fig. S8), and phosphoproteome analysis (Supplementary Figs. S10, 11), we conclude that β-events are strongly tied to dendritic spiking and increased phosphorylation of NMDA-receptors, and is present within 3 months in both striatally and cortically seeded mice. This is in large agreement with previous studies that have reported an early impact of αSyn on glutamatergic neurotransmission<sup>77</sup> and calcium-influx in neurons<sup>4</sup>. Specifically, aberrant αSyn impacted the subunit composition and function of both NMDARs and AMPARs<sup>77–79</sup>. However, the implications of such alterations were not explored within cortical microcircuit activity. Our results unambiguously establish that glutamatergic receptor function is significantly altered in the brains of mice injected with αSyn PFFs in both seeding locations and triggers aberrant L5 dendritic population activity in a location-dependent manner. Impaired somatic output resulting from this dendritic vulnerability can also alter L2/3 ensemble activity through recurrent connections. Importantly, these subcellular changes can be traced through β-events and can lead to demarcated behavioral changes. Thus, these findings support the idea that NMDA receptor hyperfunction and resultant aberrant L5 dendritic calcium spikes may be an early consequence in the propagation of αSyn pathology and provides a synaptic signature of such dysfunction in the form of β-events, much before full-scale neurodegeneration.

## Summary

In summary, our results highlight a mechanism by which αSyn pathology originating from either the dorsal striatum or primary motor cortex, characteristic of pathology propagation in PDD and DLB, can initiate distinct transaminar cortical circuit dysfunction. Critically, our results show that L5 pyramidal neurons are vulnerable targets, and play a key role in generating pathologically aberrant β-events, a potential marker of top-down input-dependent dysfunction,

and drives superficial cortical ensemble stability. Our findings suggest that this cellular vulnerability is uniquely positioned to act as a stressor leading to recurrently amplified local cortical activity, and sets a path towards future investigations centered on how temporal and spatial activity in the cortex can combinatorically inform on pathway-dependent Lewy pathology propagation. Our work also highlights the need to further explore the role of NMDA receptor and calcium channel dysfunction with regard to early-stage  $\alpha$ Syn-induced changes within cortical microcircuits and investigate potential therapeutic strategies targeting these channels to mitigate the downstream effects of  $\alpha$ Syn pathology on behavior.

## Methods

All procedures were in accordance with protocols set forth by the NIH, Purdue Institutional Animal Care and Use Committee (IACUC), and the Purdue Laboratory Animal Program (LAP). Mice were housed in a 12 h:12 h light:dark cycle (temperature, 22–23 °C; humidity, 40%). For all recordings, we used mice with a C57BL/6J (The Jackson Laboratory, #000664) background kept on a 12 h light/dark cycle in conventional housing with unrestricted access to food and water. Male and female mice were used in approximately equal numbers. Surgical procedures were carried out aseptically under 1–2% isoflurane anesthesia. Carprofen and dexamethasone was administered subcutaneously. Custom titanium headplate was attached to the skull using dental cement (Parkell Metabond). Mice were allowed to recover in their home cages for 3 days. Mice underwent habituation of head fixation and running on a 6-inch rotating wheel for at least 1 week before experiments were performed.

## Production of preformed fibrils

Mouse recombinant  $\alpha$ Syn was purified as described previously by Zhang et al. and Volpicelli et al.<sup>50,80</sup>. The expression of untagged protein variants was induced using isopropyl  $\beta$ -D-1-thiogalactopyranoside (IPTG) at 37 °C using *E. coli* BL21-DE3 cells transformed with pT7-7- $\alpha$ Syn plasmids.  $\alpha$ Syn in the bacterial lysate was concentrated using an ammonium sulfate salting-out method and enriched via a boiling step. The boiled supernatant containing  $\alpha$ Syn was passed through a HiLoad 16/600 Superdex 200 size exclusion column (Cytiva) and a HiPrepQ HP 16/10 (Cytiva) or DEAE (GE Healthcare Bio-Sciences) anion exchange column sequentially. Fractions enriched with  $\alpha$ Syn were pooled, and endotoxin was removed using the Pierce high-capacity endotoxin removal resin (Thermo Fisher Scientific) so that residual endotoxin levels were below 0.5 endotoxin units (EU) per mL<sup>81</sup>. The protein was dialyzed against phosphate-buffered saline (PBS; pH 7.4) and stored at –80 °C for 6–12 months.

$\alpha$ Syn PFFs was generated by filtering 500  $\mu$ L of a 5 mg/mL monomeric  $\alpha$ Syn solution through a 0.22  $\mu$ m filter and incubating the filtered protein solution in 1.5 mL sterile microcentrifuge tubes at 37 °C with agitation at 1,000 rpm for 7 days. After fibrillization,  $\alpha$ Syn fibrils were concentrated by centrifuging the fibril suspension at 13,000  $\times$  g for 10 min and removing the supernatant containing the unfibrillized, monomeric protein. Pelleted fibrils were resuspended in 250  $\mu$ L of Dulbecco's phosphate-buffered saline (DPBS; Cytiva). An aliquot of the fibril suspension (5  $\mu$ L) was incubated with 8 M guanidine hydrochloride for 1 h at 22 °C to dissociate fibrils into monomers. The  $\alpha$ Syn concentration was determined via absorbance measurements at 280 nm using a nanodrop instrument, with an extinction coefficient of 7450 M<sup>-1</sup> cm<sup>-1</sup>.  $\alpha$ Syn fibrils were stored at –80 °C as 25  $\mu$ L of aliquots at a concentration of 5 mg/mL. Prior to injection,  $\alpha$ Syn fibrils were sonicated in ethanol-sterilized sonicating tubes (Active Motif) using a cup horn sonicator (Qsonica q700) at 30% power (~100 W/s), with a cycle of 3 s on and 2 s off, for a total 'on' time of 15 min, while keeping the bath temperature constant between 5 and 15 °C.

## Injections

Mice of both sexes (3–6 months) were injected with PFFs and subsequently analyzed over a period of 12 weeks. Animals were anesthetized using isoflurane with the animal properly secured on the stereotaxic frame. A suspension of sonicated  $\alpha$ Syn PFFs (1.5  $\mu$ L, 5 mg/mL) or a solution of control. Monomeric protein and PFFs was injected in primary motor cortex  $\pm$  1.8 mm lateral and 0.3 mm anterior to bregma, 0.7 mm from dura and striatum  $\pm$  2 mm lateral and 1 mm posterior to bregma, 2.5 mm from dura at a constant flow rate of 0.1  $\mu$ L/min using a 10  $\mu$ L syringe (Hamilton) fitted with a 33-gauge needle with a 45 angled tip. Equal volume was administered between monomeric and PFF injected mice. Total volume was 1 and 1.5  $\mu$ L for cortical and striatal injections, respectively. The needle was left in place for 5 min post-infusion to avoid any backflow. Injected animals were administered 5 mg/kg carprofen in normal saline immediately post-surgery and 2 days following the surgery.

## Probe recordings

A small craniotomy (<1 mm) was made over the primary motor cortex (0.4 mm AP, 1.5 mm ML). A 64-channel electrode (Masmanidis Lab, UCLA) was inserted perpendicular to the surface of the pia using a micromanipulator (Sensapex 4-uMP, Finland). The microelectrode was positioned 1 mm into the cortex and was allowed to settle for 20 min before recordings began. Signals were digitized at a bandwidth of 0.1–10 kHz and sampled at 20 kHz (Intan RHD, USA).

## Intracellular recordings in vivo

Pipettes for intracellular recording (100–150 M $\Omega$  were pulled from borosilicate glass, 1 mm OD, outer diameter, 0.5 mm ID; Sutter Instruments, CA, USA), using a P-1000 micropipette puller (Sutter Instruments, CA, USA) and filled with 1 or 3 M potassium acetate. Current clamp signals from sharp pipettes were filtered by a 4-pole Bessel filter (4 kHz) and acquired at 20 kHz using a Digidata 1440. Tip potential offsets were zeros upon brain entry and prior to cell entry. The electrode was slowly advanced using a 3-axis manipulator (Luigs-Neumann, Germany) until an increase in electrode impedance was observed. Break-in was achieved through typical "buzzing" procedure<sup>82</sup>.

## Two-photon imaging

For two-photon imaging, a 3 mm cranial window was slowly drilled over the motor cortex (0.4 mm AP, 1.5 mm ML). To avoid excessive tissue heating, the skull was moistened with 0.9% sodium chloride. The dura was left intact for L2/3 imaging. Animals were imaged using two-photon microscopes (Sutter MOM, Sutter Instruments, CA, USA), and Ti:Sapphire laser (Spectra X3, Spectra-Physics, CA, USA) at 920 nm through a 16 $\times$  water immersion objective (Nikon, NA: 0.8, WD:3 mm). Resonance galvanometers were controlled through MATLAB ScanImage (Vidrio tech, MBF, USA). Each imaging session lasted 50–70 min with an acquisition rate of 30.048 Hz under resonance scanning, 512  $\times$  512 pixels, and typically with a field of view of 400  $\times$  400  $\mu$ m. For L2/3 imaging, animals expressing GCaMP6s under the Thy1-promoter was used. For dendritic imaging of L5 pyramidal neurons, 400 nl of flex-gcamp6s-mRuby (Addgene, Massachusetts, USA) was injected in the primary motor cortex (0.1 mm AP, 1.5 mm ML).

## Pharmacology

An injection micropipette was filled with 100  $\mu$ M baclofen or 500  $\mu$ M APV in standard extracellular saline. The pipette was inserted into M1 near the silicon probe and placed at a depth of ~200  $\mu$ m beneath the pial surface. The drug was delivered by gentle pressure injection. Injections (100 nl each) were made every 20 min throughout the recording session. Sessions with no injections or saline injections followed the same time course as during baclofen and APV injections.

### Signal processing

For LFP analysis, data was down sampled to 1024 Hz and bandpass filtered using a Butterworth filter from 1 to 200 Hz. Theta (4–12 Hz), Beta (12–30 Hz), and Gamma (30–90 Hz) spectrograms were extracted using a linear regression filter<sup>83</sup>. Spearman correlation was calculated across all channels, LFP above 90% similarity was considered for the analysis<sup>84</sup>. A parent LFP signal was derived by taking the median of the correlated LFPs to calculate spike-field coherence (see below). From here, locomotion-triggered value was obtained using a sliding window 2 s before and after locomotion onset.

### Beta event detection

B-events are detected based on their LFP spectral characteristics. Event candidates are determined if their power is 3 standard deviations above mean beta power, event profile is >20 ms, and <250 ms. Parameters were chosen based on the robustness of detection across pathological weeks. This routine is applied for each electrode across depth per trial.

### Spike sorting

After the recording session, we subjected the electrical signals to processing, including the sorting of action potentials from individual neurons. This sorting was accomplished using Kilosort (<https://github.com/MouseLand/Kilosort>)<sup>85</sup>, followed by a meticulous manual review in Phy2 (<https://github.com/cortex-lab/phy>). Neurons were evaluated based on waveform characteristics, stability of firing rates observed throughout the recording session, and auto-correlograms. Spiking analysis and spike-field coherence was conducted on neurons exhibiting sustained activity throughout the entire recording session. Average neural activity for each animal per pathological timepoint were smoothed by calculating the mean spiking activity across multiple trials per recording and convolving by a Gaussian filter with a full-width at half-maximum (FWHM) width of 10 ms. The depth at which each spike occurred was determined based on the electrode that recorded the maximum waveform amplitude.

### Spike-field coherence

Spike-field coherence of beta LFP was analyzed using a combination of custom coding routines and Chronux<sup>86</sup>, an open-source MATLAB package for analyzing spike-field coherence. Peristimulus beta LFP and single units was extracted using the same locomotion onset window described above. A three-dimensional array was generated for each trial containing the frequency response, time, and coherent (phase-vector) value. Phase inter-trial synchronization was derived using inter-trial phase clustering defined as

$$ITPC = |n^{-1} \sum_{r=1}^n e^{ik}| \quad (1)$$

where  $n$  is the number of trials,  $k$  is the phase angle of the LFP in which a given spike occurs.

### Image analysis

Imaging datasets were imported as Tiff stacks into ImageJ (v1.5) for batch preprocessing. High frequency denoising was done using a median filter and Kalman stack filter. Motion correction and fluorescence extraction of neurons were done using NormCorr (<https://github.com/flatironinstitute/NoRMCorre>) and CalmAn, (<https://github.com/flatironinstitute/CalmAn-MATLAB>), respectively<sup>87,88</sup>. Features extracted were refined through a convolved neural network (CNN) classifier and manual curation. Lastly, calcium spike rasters were generated by thresholding the first derivative of  $\Delta F/F$  at 3 standard deviations above the baseline fluorescence activity. This spike matrix was used for subsequent population analysis described below.

### Pairwise calcium analysis

A  $N \times T$  binary spike matrix was constructed, where  $N$  denotes the number of neurons and  $T$  denotes the total number of frames for each calcium video. Each row ( $N$ ) represents spiking events of a neuron. Coactive events were quantified by binning fluorescence traces to binary spiking events. Sørensen-Dice correlation (SDC) was used to quantify synchronous firing events of individual neurons<sup>89,90</sup>. Unique neuronal pairs identified through the binary matrix ( $N \times T$ ) are linearized and filtered through a function:

$$SDC = \frac{1}{n_{max}} \sum [S_i > 0.25] \in [P] = \begin{cases} 1 & \text{if } P \text{ is true,} \\ 0 & \text{if } P \text{ is false} \end{cases} \quad (2)$$

This metric is bound between 0 and 1 (where a value of 1 denotes that neuron pairs are identical). Thus, high cell-cell similarities represented high coactivation probabilities at any given frame between cells during periods of active locomotion. To calculate average pairwise synchronicity across pathological weeks, a bootstrapping procedure was used wherein each individual neuron's spike activity was circularly shifted by random amounts of time. This was done 1000 times across all neurons and the significant pairwise connection was calculated based on a 99% cutoff to the shuffled distribution.

### Ensemble detection and analysis

To detect cortical ensembles, we again carried out a bootstrapping procedure—1000 times across all neurons to determine periods of highly coactive activity patterns, which was reflected by the 99% percentile of fraction of total neurons activate in a given frame. Once defined, we computed the cosine similarity across the population using binned timed vectors to quantify periods of reactivation within similar neurons. The time component of the binary vector was binned every 150 ms (5 acquisition frames) and event spikes were summed to produce a time vector  $t_i = (N_i, T_i)$ , where  $N_i$  is the summed event spikes of time bin  $T_i$ . This vector series represents temporal changes of grouped neurons. To correlate changes of network states, we constructed a similarity index using a cosine-similarity function, defined as the similarity index between vector pairs  $t_i$  and  $t_j$  as<sup>20</sup>:

$$\text{Similarity Index}(SI) = \frac{t_i \cdot t_j}{\|t_i \times t_j\|} \quad (3)$$

To extract the most informative cortical ensembles across pathological times points, principle components were extracted from cosine similarity maps using singular value decomposition (SVD)<sup>19,20</sup>. SVD allows for the identification of components that contribute the most to a given network state. The SVD of a matrix can be represented as the multiplication of 3 decomposed vectors  $U, \Sigma, V^T$ . Since the multiplication of  $UV^T$  yields an identity matrix, the SVD of our vectorized matrix  $M$  was defined as  $M = VV^T\Sigma$ , where  $\Sigma$  contains eigenvalues which represent modes of matrix  $M$ . Typically >90% of the overall network activity can explained from 4–8% of the total dimension of the data whose reactivation time became candidates for ensembles.

To refine neurons involved in the ensembles, a between neuron “similarity index” as described above was calculated. The proportion of neurons within an ensemble reactivation was defined as the ensemble stability.

### Neuron simulations of Layer 5 dynamics and beta events

Morphologically detailed biophysical L5 pyramidal neuron model was implemented in the NEURON simulation environment with python (3.7) interface<sup>91</sup>. Excitatory synapses and inhibitory synapses (1200 excitatory and 300 inhibitory synapses) were randomly distributed on the basal and apical dendrites, with an AMPAR to NMDAR ratio of 1:2. The synaptic conductance was simulated with a double-exponential kinetic with following rise and decay time constant:  $\tau_{1,AMPA} = 0.1$  ms,  $\tau_{2,AMPA} = 2$  ms,

$\tau_{1,NMDA} = 2$  ms,  $\tau_{2,NMDA} = 75$  ms,  $\tau_{1,GABA} = 1$  ms,  $\tau_{2,GABA} = 5$  ms. The fully open synaptic conductance is:  $G_{AMPA} = 0.2$  nS,  $G_{NMDA} = 0.4$  nS,  $G_{GABA} = 0.8$  nS. The reversal potential for excitatory and inhibitory synapses is 0 mV and  $-75$  mV respectively. For NMDAR, a sigmoidal voltage dependent term was implemented<sup>92</sup>.

$$g_{AMPA/GABA}(t) = G_{AMPA/GABA} (\exp(-(t - t_0)/\tau_1) - \exp(-(t - t_0)/\tau_2)) \quad (4)$$

$$g_{NMDA}(t, V) = \frac{G_{NMDA}}{1 + 1/3.75 \exp(-0.062V)} (\exp(-(t - t_0)/\tau_1) - \exp(-(t - t_0)/\tau_2)) \quad (5)$$

Each synapse receives a presynaptic Poisson spike train with a lognormal distribution of average firing rate ( $\mu = 0.8-1$ ,  $\sigma = 1$ ). Single trials of 20 s simulation was used for statistical analysis of beta-event amplitude. For apical NMDA-R block, the NMDA-R conductance was reduced by 50% compared to the control condition to maintain spike generation. We applied the same condition when investigating the effect of voltage gated calcium channels in beta event generation. To observe the effect of decrease apical inhibition to beta event amplitude, the inhibitory input rate to apical dendrites was decreased 5-fold. The local field potential was computed from the transmembrane current of each compartment along the neuron using LFPy<sup>93</sup>. The simulated LFP was subsequently filtered at 12–30 Hz and  $\beta$ -events were detected using the same procedure described above.

### Brain tissue preparation for phosphoproteomic analysis

Brain tissue samples ( $\sim 15$ – $20$  mg) isolated from  $\alpha$ Syn PFF- or monomer-injected mice were resuspended in 300  $\mu$ L of 25 mM ammonium bicarbonate (ABC) supplemented with protease and phosphatase inhibitors and homogenized in a Precellys Evolution tissue homogenizer using soft-tissue homogenizer CK14 tubes (Bertin Technologies SAS, France). The total protein concentration was then measured using the bicinchoninic acid (BCA) assay (Pierce Chemical Co., USA). The Bligh and Dyer (B & D) extraction method was then performed using lysate volumes equivalent to 25  $\mu$ g of total protein. Four volumes of cold ( $-20$  °C) methanol were then added to the water-methanol layer, and samples were centrifuged at 17,200  $\times g$  for 10 min. The supernatant was removed, and pelleted proteins were dried in a vacuum centrifuge. The protein samples were resuspended in 10  $\mu$ L 8 M urea solution supplemented with 10 mM DTT and incubated for 1 h at 37 °C. Next, 10  $\mu$ L of alkylation reagent mixture (97.5% acetonitrile (ACN), 0.5% triethyl phosphine, 2% iodoethanol) was added, and samples were incubated for 1 h at 37 °C. After alkylation, samples were dried in a vacuum centrifuge and subsequently resuspended in 80  $\mu$ L of 0.025  $\mu$ g/ $\mu$ L trypsin (Pierce Chemical Co., USA). The digestion was carried out using a bar cycler (50 °C, 60 cycles; 50 s at 20 kPSI and 10 s at 1 ATM). Peptides were then desalted with a C18 silica MicroSpin column (The Nest Group Inc, USA). Phosphopeptide enrichment was then performed with PolyMac spin tips (Tymora Analytical, USA), following the manufacturer's recommendations for phosphoproteomic analysis.

### Mass spectrometry analysis

Peptides were separated using the UltiMate 3000 RSLC nano HPLC system (Thermo Fisher Scientific) connected to an Orbitrap Fusion Lumos mass spectrometer (Thermo Fisher Scientific). Peptide separation was carried out in a PepMap trap column (50 mm  $\times$  300 mm ID, with a 5  $\mu$ m particle size and 100 Å pore size), followed by an Acclaim PepMap C18 silica column (500 mm  $\times$  75  $\mu$ m ID, with a 2  $\mu$ m particle size and 100 Å pore size). Samples were eluted using a gradient between two buffers, buffer A (0.1% formic acid in water) and buffer B

(0.1% formic acid in 80% ACN). Samples were injected into the trap column for 5 min, using a flow rate of 5  $\mu$ L/min. Then, the flow rate was set to 150 nL/min, and samples were separated in the analytical column using a 160 min gradient. Briefly, mobile phase B was set to 2% and increased to 27% for 110 min, after which the gradient was increased to 40% B within 15 min. The percentage of mobile phase B was then increased to 100% in 10 min, and held constant for an additional 10 min, after which the mobile phase B concentration was reset to 2%.

MS analysis was performed with the orbitrap detector, with a MS1 resolution of 60,000 and MS2 resolution of 15,000. Quadrupole isolation was set to "True". The scan range was set to 375–1600 m/z. The Radio Frequency (RF) lenses were set to 30%, Automatic Gain Control (AGC) target was set to "Standard", with a maximum injection time "Auto", and 1 microscan. A dynamic exclusion duration of the 60 s was used, with the exclusion of isotopes. Data was collected using Data-Dependent Acquisition (DDA) mode with cycle time of 3 s, between scans. The precursor ion (MS1) intensity threshold was set to  $5.0 \times 10^3$  and peptides were fragmented using high energy collisional dissociation (HCD) with 30% collision energy. MS/MS data were also collected using "Orbitrap" as a detector, with AGC target "Standard", maximum injection time "Auto" and 1 microscan and isolation window of 1.2 m/z. The procedures listed above is consistent across previously published works<sup>94–97</sup>.

### Bioinformatics analysis

Raw LC-MS/MS data were searched against the UniProt<sup>98</sup> *Mus musculus* processed using the Maxquant platform<sup>99</sup> (Ver 2.0.3.1; [www.maxquant.org](http://www.maxquant.org)). Searches were performed using trypsin as an enzyme allowing up to 2 missed cleavages. Variable modifications were set for "methionine oxidation" and for phosphoproteomics "STY phosphorylation" and "carbamidomethyl" as a fixed modification. Other search parameters and data filtering were similar as described in ref. 100. A downstream analysis of Maxquant search results was performed using Perseus computational platform<sup>101</sup>. Raw data files were filtered based on contaminants, and intensity values were log 2-transformed. Protein/peptide entries present in >70% of the samples were selected for downstream analysis. Data imputation was performed for the replacement of missing values and data quality validation. A two-sample *t*-test was performed on the samples with permutation-based false discovery rate (FDR) correction. Heatmaps were generated in MATLAB, and sample clustering based on principal component analysis was performed utilizing MetaboAnalyst 5.0 (<http://www.metaboanalyst.ca/>).

### Behavior assays

To evaluate the effects of transmissible  $\alpha$ Syn-synuclein pathology on motor, cognitive learning and memory, and emotional behavior, mice were tested on a comprehensive behavior for 2 weeks<sup>2</sup>. The order of tests was randomized, and all tests were conducted by an experimenter blinded to treatment group. All tests were conducted between 08:00 and 4:00 in the lights-on cycle. Mice were habituated to the testing room 1 h before tests, and the apparatus were cleaned with 70% ethanol in between animals to minimize odor cues.

### Open field test

To assess general activity, locomotion, and anxiety, mice were tested in the open field. Mice were individually placed in the center and allowed to freely explore for 15 min while the trial was videotaped. Subsequent video scoring was completed by an observer blind to treatment groups using Bonsai and custom coded MATLAB scripts to analyze exploration index and rearing time. Exploration index was calculated as the time each mouse spends in a defined area within the arena (a value of  $-1$  denotes very high occupation in a single region while 1 denotes brief occupation in a single region).

### Wire hang test

The wire hang test of motor function was conducted to observe mouse grip strength. The mice were placed onto a 24-gauge steel wire 60 cm above a Plexiglas case. The wire was lightly shaken to cause the animals to grip the wires and then released. The latency of mice to fall off the steel wire was measured, and average values were computed from two trials (10 min apart).

**Statistics.** For comparisons across two or more unpaired groups, the single-sided, Kruskal–Wallis ANOVA test was used. Multiple (3 or more) groups were compared using a Two-way unbalanced ANOVA. We conducted nested statistics of neurons->mouse->treatment group to report statistical values. In cases where this yielded significance ( $p < 0.01$ ), post-hoc Bonferroni test was reported for comparisons between pairs of groups. In the case of paired groups, a Wilcoxon–Mann–Whitney U-test was used.

**Immunohistochemistry (IHC).** Animals were euthanized by an overdose with isoflurane and perfused transcardially with cold PBS. Brains were removed quickly after decapitation and post-fixed overnight in the same fixative, and then cryoprotected in water containing 30% (w/v) sucrose at 4°C. Coronal sections (40 µm thick) were prepared from the brains of perfused mice using a frozen sliding microtome (Thermo Scientific Microm HM430). Sections were stored at -20°C in cryoprotectant (30% w/v sucrose and 30% v/v ethylene glycol in 0.1 M phosphate buffer) prior to IHC. Sections were blocked with a solution of 10% (v/v) normal donkey serum in PBS supplemented with Triton-X100 (0.3% v/v) (PBST) for 90 min and then incubated with primary antibody solution prepared in PBST supplemented with 1% (v/v) normal donkey serum overnight at 4°C. After washing in PBS, sections were incubated with secondary antibodies conjugated with Alexa fluorophores (Jackson ImmunoResearch Laboratories, 1:500 dilution) for 90 min at 22°C and then washed 3 x 10 min in PBS. The tissues were mounted on slides, allowed to dry overnight, and sealed with a coverslip using DPX mounting media (cat. no. 13512, Electron Microscopy Sciences). Primary antibodies used in this study were: anti-HuC/HuD (cat. no. A21271, Invitrogen, 1:50), anti-NeuN (cat. no. CTX00837, GeneTex, 1:1000) anti-pSer129-aSyn (EP1536Y) (cat. no. ab51253, Abcam, 1:500), anti-p-Ser129-aSyn (81a) (cat. no. MABN826, Millipore Sigma, 1:500), anti-GRIN2B (cat. no. 54739, Signalway Antibody, 1:100), and anti-p-Ser1303-GRIN2B (cat. no. 11821, Signalway Antibody, 1:100). Quantitation of pathology from IHC sections was done using fluorescence analysis algorithms of annotated brain regions.

### Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

### Data availability

Calcium imaging and electrophysiological processed datasets generated in this study have been deposited in the Zenodo database under the accession code: 10.5281/zenodo.14029419. The mass spectrometry proteomics data pertaining to this study have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier [PXD058091](https://doi.org/10.26434/chemrxiv-2024-pxd05). We have also uploaded phosphopeptide spectra to MS Viewer. These can be accessed using the key mbua9pixto. Data generated in this study are provided in Source Data file. Source data are provided with this paper.

### Code availability

Code and example MATLAB scripts corresponding to figures, analysis of cortical ensembles, and detection of beta events are available at: <https://github.com/HammadFKhan/SiteSpecificLewyPathologyCircuits>.

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## Author contributions

H.F.K. designed, performed, and analyzed experiments. S.D. designed/optimized the synuclein seeding strategy, designed and performed histological and proteomic analysis. A.N.S. assisted with proteomic analysis and injections. S.X. and S.Y. assisted with biophysical modeling and microscope build-out. X.C. assisted with experiments early in the study. U.K.A. performed detailed proteomic

analysis. T.L.K.-U., J.-C.R., and K.J. advised on experiments. J.-C.R. and K.J. conceived the study. K.J. designed experiments and oversaw the project. H.F.K. and K.J. wrote the paper with contributions and inputs from all authors.

### Competing interests

The authors declare no competing interests.

### Additional information

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