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Microglial *INPP5D* limits plaque formation and glial reactivity in the PSAPP mouse model of Alzheimer's disease

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Abstract

Introduction: The inositol polyphosphate-5-phosphatase D (*INPP5D*) gene encodes a dual-specificity phosphatase that can dephosphorylate both phospholipids and phosphoproteins. Single

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CONFLICTS OF INTEREST

SAL is a founder of AstronauTx Ltd; all other authors declare no competing interests.

SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

nucleotide polymorphisms in *INPP5D* impact risk for developing late onset sporadic Alzheimer's disease (LOAD).

Methods: To assess the consequences of inducible *Inpp5d* knockdown in microglia of APP^{KM670/671NL}/PSEN1^{exon9} (PSAPP) mice, we injected 3-month-old *Inpp5d*^{fl/fl}/*Cx3cr1*^{CreER/+} and *PSAPP/Inpp5d*^{fl/fl}/*Cx3cr1*^{CreER/+} mice with either tamoxifen (TAM) or corn oil (CO) to induce recombination.

Results: At age 6 months, we found that the percent area of 6E10⁺ deposits and plaque-associated microglia in *Inpp5d* knockdown mice were increased compared to controls. Spatial transcriptomics identified a plaque-specific expression profile that was extensively altered by *Inpp5d* knockdown.

Discussion: These results demonstrate that conditional *Inpp5d* downregulation in the PSAPP mouse increases plaque burden and recruitment of microglia to plaques. Spatial transcriptomics highlighted an extended gene expression signature associated with plaques and identified CST7 (cystatin F) as a novel marker of plaques.

Keywords

Alzheimer's disease; cystatin F; Inpp5d; microglia; oligomer; SHIP1; spatial transcriptomics

Highlights

- *Inpp5d* knockdown increases plaque burden and plaque-associated microglia number.
- Spatial transcriptomics identifies an expanded plaque-specific gene expression profile.
- Plaque-induced gene expression is altered by *Inpp5d* knockdown in microglia.
- Our plaque-associated gene signature overlaps with human Alzheimer's disease gene networks.

1 | BACKGROUND

Late-onset sporadic Alzheimer's disease (LOAD) is the most common form of dementia, characterized by progressive memory decline, leading to loss of cerebrocortical function and eventual death. Genome-wide association studies (GWAS), whole genome sequencing, and gene-expression network analysis have identified hub and driver genes and their networks associated with risk of LOAD.^{1,2} Expression of a large proportion of these genes is specific to, or enriched in, microglia, underscoring a potential role for microglia in AD pathogenesis.³ Multiple approaches have highlighted inositol polyphosphate-5-phosphatase D (*INPP5D*) as an AD risk gene, and its expression in the brain is restricted to microglia.⁴⁻⁶ Current evidence suggests that increased *INPP5D* levels associated with inheritance of the common variant rs35349669 confer increased risk for LOAD.⁷ A second variant in *INPP5D*, rs61068452, was found to be protective against AD, and was significantly associated with lower cerebrospinal fluid (CSF) total tau (t-tau)/amyloid beta (A β)₄₂ ratio.⁸

INPP5D is a dual-specificity phosphatase that can dephosphorylate both phospholipids and phosphoproteins.^{9,10} *INPP5D* encodes the 5' phosphatase SHIP1 (Src homology 2 [SH2] domain-containing inositol polyphosphate 5' phosphatase) protein. SHIP1 negatively regulates targets such as immune cell surface receptors by catalyzing the conversion of phosphatidylinositol (3,4,5)-trisphosphate (PI[3,4,5]P3) to phosphatidylinositol (3,4)-bisphosphate (PI[3,4]P2). This lipid dephosphorylation reaction prevents receptor activation, mitigating signal transduction through the phosphatidylinositol-3-kinase/mammalian target of rapamycin (PI3K/mTOR) pathway.¹⁰ In particular, SHIP1 inhibits phagocytic signal transduction by triggering receptor expressed on myeloid cells 2 (TREM2) and tyrosine kinase binding protein (TYROBP, also known as DAP12).¹¹ *DAP12/TYROBP* was also identified as a key driver gene in AD via an unbiased, multi-scale computational network. Constitutive deletion or pharmacologic inhibition of SHIP1 increases phagocytosis of various substances, including zymosan in SHIP1 knockout (KO) macrophages and A β in SHIP1 inhibitor-treated microglia.^{12,13} Promotion of these functions in microglia has been considered a means for preventing or reducing AD pathology and improving disease outcomes, but loss of the inhibitory signal from SHIP1 could also promote inflammatory processes that are detrimental in AD.¹⁴

These studies strongly implicate a role for *INPP5D* expression or activity of the encoded protein, SHIP1, in AD pathophysiology. Additionally, *INPP5D* expression increases with progression of clinical LOAD and with aging in the 5XFAD mouse model.¹⁵ The specific role(s) that *INPP5D* play(s) in both early or late disease, and the mechanism(s) contributing to these altered functions, remain unknown. We therefore knocked down *Inpp5d* conditionally in microglia to determine early effects as amyloid pathology develops in *APP^{KM670/671NL}/PSEN1^{exon9}* (PSAPP) mice at a time correlating to a preclinical level. This strategy also avoids the developmental effects that can occur in constitutive knockouts.^{16,17} Although the simplest rationale may support hypothesizing that reduction of *INPP5D* would be beneficial in AD, the complex interactions associated with inflammatory signaling and its contribution to disease progression would suggest the alternative; that is, detrimental outcomes are just as likely. Herein, we demonstrate that *Inpp5d* downregulation increases A β fibrillization, plaque burden, and number of microglia associated with A β plaques in the PSAPP amyloid-depositing mouse model of AD.

2 | RESULTS

2.1 | *Inpp5d* knockdown worsened amyloid plaque deposition while increasing recruitment of microglia to the plaque

We used a combination of previously established mouse lines, including the PSAPP mouse, to assess pathological consequences of *Inpp5d* knockdown (Figure 1A, Figure S2A in supporting information). To induce recombination, 3-month-old *Inpp5d^{fl/fl}/Cx3cr1^{CreER/+}* mice, with and without the *PSEN1* and *APP* transgenes, were injected for 5 consecutive days with either tamoxifen (TAM) or corn oil (CO). SHIP1 protein levels were reduced by 75% 1 month post-recombination (Figure S1 in supporting information). At 6 months of age, a reduction in SHIP1 protein levels was maintained in both TAM-treated *PSAPP/Inpp5d^{fl/fl}/Cx3cr1^{CreER+}* (PSAPP-TAM) and wild-type *Inpp5d^{fl/fl}/Cx3cr1^{CreER+}* (WT-TAM) mice

compared to CO-treated mice, *Inpp5d*^{fl/fl}/*Cx3cr1*^{CreER+} (WT-CO), or *PSAPP/Inpp5d*^{fl/fl}/*Cx3cr1*^{CreER+} (PSAPP-CO; Figure 1B,C; Figure S1). Similar to findings in the 5xFAD mouse model,¹⁵ PSAPP-CO mice showed an increase in SHIP1 protein levels.

At 6 months of age, the percent area and number of 6E10 immunoreactive deposits were increased by >66% and 73%, respectively (Figure 1D–F; Figure S2) in the hippocampi of PSAPP-TAM compared to PSAPP-CO littermates. There was no difference in the size distribution of individual amyloid plaques in PSAPP-TAM compared to PSAPP-CO mice (Figure 1G). Staining with the Congo red derivative Methoxy-XO4 (XO4), which binds specifically to β -pleated sheet-containing structures, confirmed the increase in amyloid burden associated with the *Inpp5d*/SHIP1 knockdown (Figure S3A–B in supporting information). Fibrillar deposits were increased despite unchanged levels of A β 40, A β 42, or A β 42/A β 40 ratios as determined by enzyme-linked immunosorbent assays (ELISAs) of Tris-buffered saline (TBS)-, Triton-X-, and formic acid extracts of mouse brain homogenates (Figure S4 in supporting information).

As both microglia and astrocytes contribute to the immune response in AD,^{18,19} we next measured whether *Inpp5d* knockdown altered their numbers. Microglia numbers are increased in *PSAPP* mice with prominent peri-plaque localization.²⁰ In the WT-CO and WT-TAM mice, the number of IBA1⁺ cells or IBA1 immunoreactive areas in the hippocampus was unchanged after *Inpp5d* depletion (Figure 1H–I, Figure S3E). Similarly, these parameters did not change when comparing PSAPP-CO to PSAPP-TAM animals (Figure 1D, H–I). There were more IBA1⁺ cells and IBA1 immunoreactive areas in PSAPP-TAM mice compared to WT-CO mice (Figure 1H–I), whereas the difference between PSAPP-CO and WT-CO was not significant. Most striking was a spatially constrained increase in microglia associated with A β plaques (Figure 1C,J–L). We did not observe a significant increase of glial fibrillary acidic protein (GFAP)⁺ cell numbers or immunoreactive areas in the hippocampus in association with *Inpp5d* knockdown (Figure S3A, C–D).

Given reports of synapse density loss early in AD²¹ and a decrease in synaptogenesis in some AD-associated reactive astrocytes,²² we next investigated changes in synaptic markers in our mice. Western blotting of protein extracts from the hippocampi of female WT-CO, WT-TAM, PSAPP-CO, and PSAPP-TAM mice at age 6 months using anti-PSD95 and anti-synaptophysin antibodies showed no obvious effect of any manipulation or parameter on brain levels of either synaptic marker (Figure S5 in supporting information). However, epitomic characterization²³ of the A β conformers (i.e., A11, OC, NU4) present in the brains of the various groups (Figure S5B) revealed a \approx 50% increase in A11/6E10 ratio (i.e., ratio of prefibrillar oligomer content to total A β content). No such elevation was detected for OC/6E10 or NU4/6E10 (Figure S5).

2.2 | PSAPP mice expressed a unique plaque-induced gene cluster that was exacerbated by *Inpp5d* knockdown

We next performed spatial transcriptomics to analyze brain region–resolved differential gene expression due to the presence of the *APP* and *PSEN1* transgenes and the knockdown of *Inpp5d* (Figure 2, Figures S6–11 in supporting information). We sequenced a total of 12 coronal brain sections from WT-CO, WT-TAM, PSAPP-CO, and PSAPP-TAM

animals (Figure 2A) and reliably identified brain regions based on gene expression profiles (Figure 2B,C). We found that one cluster, Cluster 26, was exclusively expressed in PSAPP mice (Figure 2D). Cluster 26 showed a highly cluster-specific gene expression profile as calculated using a Wilcoxon rank sum test comparing the gene expression in Cluster 26 to all other clusters (Figure 2E) and showed that it co-localized with 6E10-positive plaques of adjacent sections (Figure 2F). Gene ontology (GO) analysis of Cluster 26 genes included “Microglia pathogen phagocytosis pathway” (example genes include: *Clqa/b/c*, *Fcer1g*, *Vamp8*), “Tyrobp causal network” (*Tyrobp*, *Cd37*, *Cd84*, *Itgam*, *Spp1*), “Lysosome” (*Cd63*, *Hexa/b*, *Ly86*), and “Antigen processing and presentation” (*B2m*, *H2-D1/K1*, *Axl*; Figure 2G, Tables S1–S5 in supporting information). Given that A β plaques have been associated with increased inflammation,²⁴ we next asked whether previously reported reactive microglial or astrocyte substates had similar gene expression profiles and were present in the same spatial location as Cluster 26. Indeed, Cluster 26 had a unique gene expression profile, but we found overlapping gene sets compared to previously reported, biologically defined reactive glial substates. This included integration with differentially expressed genes (DEGs) from disease associated microglia, both early (DAM1) and late (DAM2),²⁵ disease associated astrocytes (DAA),²⁶ plaque-induced genes (PIGs),²⁷ and our previously identified interferon-responsive reactive astrocytes (IRRAs;²⁸ Figure 2H). While DAMs and IRRAs co-localized with Cluster 26, DAAs had a broad expression throughout the brain, which was not surprising given the many non-specific homeostatic astrocytic genes present within the module. We also recovered 50 of the 57 previously identified PIGs and found that the PIG module co-localized with plaques and Cluster 26 (Figure 2H).

We next asked whether *Inpp5d* knockdown altered peri-plaque gene expression patterns. We used *muscat*,²⁹ a cluster-resolved, pseudobulk-based DEG analysis approach for multi-sample, multi-condition experiments. We used *edgeR* within *muscat* to calculate genes differentially expressed between PSAPP-CO and PSAPP-TAM brains. We found that the gene expression profile in Cluster 26 was extensively altered by the knockdown of *Inpp5d* (Figure 2I). Of the 231 genes induced in Cluster 26 upon knockdown of *Inpp5d* ($\text{padj} < 0.05$, $\text{l2f} > 1$), 126 were specific to PSAPP-TAM plaques. GO term analysis of these genes included an enrichment for “Defense response to virus” (*Cxcl10*, *Ifit3*, *Oas1a*, *Ccl5*, *Oas1l*), “Type I and II Interferon signaling” (*Stat1*, *Ifit2*, *Irf5/6*, *Isg15*), and “Chemokine signaling pathway” (*Plcb2*, *Rac2*, *Ccl9/12*, *Txnip*, *H2-T23*; Tables S6, S7 in supporting information). While we discovered most DEGs in Cluster 26 due to its plaque association (Figure 2F, Figures S7–S9, Tables S1–S5), *muscat* also uncovered DEGs outside Cluster 26 when comparing WT-TAM to PSAPP-TAM as well as PSAPP-CO to PSAPP-TAM. Given the recent evidence for Cre-mediated activation of microglia,³⁰ we calculated genes that were changed in WT-CO versus WT-TAM mice and found that while tamoxifen injection alone drove a modest numbers of DEGs, they did not fall into categories associated with interferon signaling (Figure S10, Table S2). Last, we used *NICHES*, a recently developed package based on the *phantom5* database,³¹ to interrogate which putative receptor–ligand interactions were enriched within Cluster 26 and therefore plaques. We confirmed that *NICHES* reliably detected receptor–ligand pairs along anatomical regions. In Cluster 26, we observed receptor–ligand pairs primarily involved in phagocytosis and lysosome activity,

including (*ligand – receptor*): *C1qb – Lrp1*, *C3 – Lrp1*, *Timp1 – Cd63*, *Pros1 – Axl*, and *Ly86 – Cd180* (Figure S11, Table S8 in supporting information).

2.3 | Cluster 26 and *Inpp5d* knockdown signatures in PSAPP intersected human AD gene networks

To expand our *NICHES* receptor–ligand interaction spatial transcriptomic maps, we used network analysis on cluster-specific DEG lists. Leveraging high-throughput molecular profiling techniques, we generated a cohort of matched whole-genome sequencing (WGS) and RNA-seq data across four brain regions, from a set of 364 well-characterized AD and control brains in the Mount Sinai Brain Bank cohort.³² Using Bayesian probabilistic causal network (BN) analysis,^{33–35} we built integrative network models to organize genome-wide gene expression features into regulatory networks.³² When projecting the Cluster 26 expression signature and cluster-specific DEG list onto the network neighborhood of *INPP5D* in the parahippocampal gyrus region BN, we found that both were significantly enriched in the subnetwork within a path length of 6 of *INPP5D* (up to 16-fold enrichment, adjusted *P*-value = 3.1E-30; Table S9 in supporting information; Figure 3A). DAM genes like *Gpnmb*, *Trem2*, and *Tyrobp* were among the DEGs present in the network neighborhood of *INPP5D*. Figure 3A displays the expression changes in human AD brain and in *Inpp5d* knockdown mice within the subnetwork of up to four steps from *INPP5D*. In this subnetwork, half of the genes (59/119) were detected in the mouse study, among which 18 were significantly regulated by *Inpp5d* (19.9-fold, adjusted *P*-value = 3.3E-18). We therefore argue that Cluster 26 as well as *Inpp5d* knockdown signatures in PSAPP mice mirrored those observed in human AD gene networks. Indeed, when probing for human AD nodes from the *INPP5D* network, we found that these were regulated by the knockdown of *Inpp5d* and co-localized with the plaque-associated Cluster 26 (Figure 3B, Figure S12 in supporting information).

2.4 | Novel plaque marker *Cst7* was limited to plaque-associated microglia

We next probed for the gene *Cst7* (cystatin F), which is specific to Cluster 26 and which has previously been identified in DAMs and PIGs (Figures 2E,H, 3C–G).^[25,27] In humans, *CST7* is expressed in endothelial cells in addition to microglia and macrophages.³⁶ In our current PSAPP mice cohorts, *Cst7* remained upregulated in both CO- and TAM-treated mice, suggesting that *CST7* might be an excellent marker of plaques. To localize *Cst7* to specific central nervous system cell types, we used RNAScope in situ hybridization (Figure 3E). *Cst7* co-localized with *Aif1*⁺ microglia and with A β plaques in PSAPP, but not in plaque-less WT animals. This co-localization was enabled by the spatial transcriptomic analysis and would not have been detected if we had performed only bulk analysis of all microglia. Further analysis of the spatial transcriptomics data also confirmed the relative specificity of Cluster 26 genes as putative plaque markers, showing that *Cst7* was expressed in 100% of plaque spots and only 8% of non-plaque spots (Figure 3F). *Cst7* therefore outperformed both *Trem2* as well as *Tyrobp* as plaque marker, both of which were highly enriched around plaques but also showed a broad expression in non-plaque tissue. We confirmed the specificity of *Cst7* using immunohistochemistry and show that the *CST7* protein was exclusively present around plaques in PSAPP brains (Figure 3G), leading us to propose that it is a novel marker for A β plaques. Previous studies associated altered

Cst7 expression with microglial dysfunction leading to impairment of A β clearance,³⁷ and upregulation of *Cst7* in A β -containing microglia (but not A β -deficient microglia) from the 5xFAD mouse model of AD.³⁸ These data imply that decrease of SHIP1 function after initiation of AD amyloid pathology leads to an increase in plaque burden and plaque-associated microglial gene expression.

To further assess this in light of our data suggesting an increase in plaque deposition (Figure 1E–F), we performed a phagocytosis assay in vitro to compare microglia from *Inpp5d*-WT to microglia from *Inpp5d*-KO mice. We found that phagocytosis was increased in microglia that lacked *Inpp5d* (Figure 3H), in line with previous reports in the literature.¹² We hypothesize that under chronic stimulation from A β in the mouse model, microglia that lack *Inpp5d* undergo functional impairment in association with A β plaque deposition as was described previously,³⁹ resulting in more plaques.

3 | DISCUSSION

Here we report that conditional *Inpp5d* downregulation in microglia of the adult PSAPP mouse increases plaque burden and recruitment of microglia to plaques. This result was not predicted by the apparent effect of the common *INPP5D* polymorphisms, given that the rs35349669 single nucleotide polymorphism (SNP) confers AD risk,⁷ while rs61068452 is apparently protective against AD.⁸ Notwithstanding, *INPP5D* plays a role in AD pathogenesis, and remains a potential target for microglia-focused AD therapies,¹⁵ but its role in brain and disease progression requires clarification.

There are multiple reasons to have predicted that knockdown of *Inpp5d* might lead to an improved or delayed amyloid phenotype in mice, including perhaps decreased plaque deposition. Although SHIP1 protein has been shown to be elevated in the 5xFAD mouse,¹⁵ protein level has not been assayed in human tissue. Moreover, microglia with decreased *Inpp5d* or in the presence of a SHIP1 inhibitor, demonstrated increased phagocytosis.^{12,13} SHIP1 also blocked TREM2/TYROBP signal transduction, implying that *Inpp5d* knockdown should increase activity of this pathway. Finally, *Inpp5d* was identified as a target of miR155, and constitutive deletion of miR155, which should increase SHIP1, led to an increase in amyloid plaques.^{40,41} However, we observed increased plaque deposition and compaction. The conundrum of predicting the outcome in our study is similar to consideration of TREM2 action for which gain of function is predicted to be beneficial,⁴² yet opposite effects can be seen in a “Jekyll and Hyde” context-dependent scenario.⁴³ Thus, SHIP1 inhibition increased phagocytosis in vitro while *INPP5D* expression has been reported to increase in AD patients as plaque load increases, whereas, in our study, decreased *Inpp5d* levels were associated with an increase in plaques. Further complicating matters, haploinsufficiency and total deficiency of TREM2 can lead to opposite effects,⁴⁴ and differing outcomes have also been reported with microglial depletion.^{45,46} Notably, the timing of gene and/or total microglial manipulation is also critical,⁴⁷ and as we noted, we deliberately used conditional knockdown early in “disease” stage to mimic what might be therapeutically viable.

Detrimental outcomes after loss of *INPP5D* inhibition are not entirely unexpected taken in the context of TREM2/TYROBP signaling. This pathway regulates the switch from a homeostatic to a neurodegenerative molecular signature by inducing miR155 via apolipoprotein E (APOE) signaling,⁴⁸ and TYROBP has also been shown to regulate APOE signaling independent of TREM2.⁴⁹ Support for beneficial outcomes associated with maintenance and/or restoration of homeostatic functions of microglia comes from the numerous studies targeting the members of this pathway, including *Trem2*,^{48,50–52} *Tyrobp*,^{53,54} and miR155.^{41,55} However, as we have previously seen in our studies modulating *Tyrobp* expression, manipulation of molecules involved in immune/microglial function in either direction may result in detrimental or exacerbated pathological outcomes.^{49,53,54,56} A similar interpretation could be developed from the studies of *Trem2* deficiency, as differences were dependent on early versus later disease stages.^{50,51,57,58} It remains to be determined whether the increased plaque burden and alterations in microglia phagocytosis described herein in PSAPP mice after *Inpp5d* knockdown also occur in other mouse models of AD (e.g., 5xFAD), or at different stages during the progression of disease pathology.

Notably, western blots of hippocampal extracts from female mice from each comparison group at age 6 months were probed with anti-PSD95 and anti-synaptophysin antibodies. Despite the increase in amyloid burden, the increase in plaque-associated microglia, and the appearance of DAM genes (including *Clq*), there was no evidence for destruction of hippocampal synapses as determined by the tissue content of these synaptic markers in mice after induction of the *Inpp5d* knockout. However, epitomic characterization of these same extracts revealed a $\approx 50\%$ increase in A11/6E10 ratio (i.e., ratio of prefibrillar oligomer content to total A β content). This observation suggests that *Inpp5d* knockdown may alter the conformational state of A β in such a way as to increase plaque density while apparently reducing synaptic toxicity. Methylene blue⁵⁹ and peroxisome proliferator-activated receptor gamma agonists⁶⁰ have been reported to have similar conformation-altering, toxicity-modifying effects.

All the same factors altered by *Trem2* manipulation are impacted by *Inpp5d* knockdown. Thus, it becomes very difficult to predict what will happen in vivo in the face of incipient plaque deposition, as assayed in this report, and which changes are beneficial or detrimental. Increased plaque compaction may be a protective mechanism,^{59,60} and there are many examples of environmental and/or microglial gene manipulation in mouse models of amyloidosis in which cognition improved in the absence of a decrease, or even in the presence of an increase, in plaques.^{61,62} This may have therapeutic implications, in that inhibition of SHIP1 once disease is initiated might not lead to a clinical benefit. Further, increased association of microglia with plaques reduces neuritic dystrophy.⁶³ The existence of DEGs in regions without plaques after the knockdown of *Inpp5d* indicates that (at least within the time period of this study) these transcriptomic changes do not always result in plaque deposition.

We identified a plaque-specific gene expression profile, Cluster 26, which is extensively altered by the knockdown of *Inpp5d*, moving cells in this region toward a higher inflammatory profile. The latter results and that of our functional phagocytosis assay

showing increased bead phagocytosis by constitutive *Inpp5d* KO primary microglia were consistent with previous results,¹² and with a role for SHIP1 as a negative regulator of immune signaling.

Increased microglial phagocytosis could be predictive of a beneficial reduction in plaque burden, which is the opposite of the pathological outcomes we found in vivo. Notably, previous studies have shown that microglial function may be adversely affected by A β , particularly as a result of chronic exposure to pro-inflammatory cytokines, resulting in increased A β accumulation and potentially underpinning the contradiction between our in vitro and in vivo results.^{39,40} Indeed, our *Inpp5d* knockdown exacerbated the plaque-induced Cluster 26 profile, which overlaps with previously reported reactive glial substates, including DEGs from both early and late DAM profiles,²⁵ and our previously identified IRRAs.²⁸ Further studies will be required to elucidate the mechanisms underlying *Inpp5d*-mediated phagocytosis and plaque deposition changes in the context of AD.

There are several important caveats to consider in the interpretation of our results. SHIP1 is under investigation as a potential therapeutic target for treatment of AD, with exploration of both inhibitors and agonists. Recent work showed that a SHIP1 agonist can enhance phagocytic degradation of synaptosomes and dead neurons by microglia.⁶⁴ The focus of our investigation described herein was on the inhibition of SHIP1, with the expectation that this would boost microglial functions (e.g., phagocytosis of amyloid) with the goal of reducing plaque burden: but in fact, plaque burden increased. These seemingly conflicting data emphasize the complexity in identifying an effective SHIP1-targeted therapeutic strategy that is underpinned by our results in vivo. Although our findings have multiple therapeutic implications, a caveat is our inability to associate this altered pathology with any detectable changes in brain function, which may be a result of our focus on the earliest stages of pathology and/or a dissociation between plaque load and function, which has been observed in other settings.^{21,60,62} Assessment of the impact of intervening with SHIP1 modulation at a range of disease stages will be critical in determining whether boosting or inhibiting SHIP1 activity is most likely to lead to a meaningful clinical benefit in patients at high risk for, or already suffering from, the clinical dementia of AD. Translating results from mouse models to human disease has been notoriously difficult, especially for AD, which has seen multiple failures of potential drugs and therapies once at the clinical trial stage. Further validation, such as with inducible CRISPR interference against *INPP5D* in induced pluripotent stem cell-derived microglia, could be a promising approach.⁶⁵

Nonetheless, we show here that manipulating microglial SHIP1 has a profound impact on the amyloidosis and neuroinflammation of AD. This, combined with the GWAS-associated SNPs in *INPP5D*, highlights microglial function and/or dysfunction as a central contributor to early AD pathology. Ultimately, further research into the disease-stage-specific functional consequences of manipulating SHIP1 activity is required to determine whether SHIP1 is a viable therapeutic target to prevent or treat the clinical dementia associated with AD. Importantly, our spatial transcriptomics analysis highlights an extended DEG signature associated with plaques and identifies *Cst7* as a potentially highly specific marker of plaques in the AD brain. We also took advantage of cross-species gene regulatory network analysis to highlight crossover of the *INPP5D*/SHIP1 pathway in human AD, as some aspects

of plaque pathology can be species-specific. The network analysis and extensive spatial transcriptomics data should prove helpful for other groups seeking to localize key genes of interest in the PSAPP mouse brain for their own research.

4 | METHODS

4.1 | Animals

All animal procedures were conducted in accordance with the National Institutes of Health Guidelines for Animal Research and were approved by the Institutional Animal Care and Use Committee (IACUC) at the Icahn School of Medicine at Mount Sinai. Mice were housed on a 12 hour light/dark cycle and had access to food and water ad libitum. *APP^{KM670/671NL} x PSEN1^{exon9}* (PSAPP), B6.129S6-*Inpp5d^{tm1Wgk/J}* (*Inpp5d^{fl/fl}*; stock no.: 028255), B6.129P2(Cg)-*Cx3cr1^{tm2.1(cre/ERT2)Litt/WganJ}* (*Cx3cr1^{CreER/CreER}*; stock no.: 021160), and B6.C-Tg(CMV-cre)1Cgn/J (CMV-Cre; stock no.: 006054) were purchased from The Jackson Laboratory. The inducible *Cx3cr1^{CreER/CreER}* mice were crossed with mice carrying *loxP*-flanked *Inpp5d* to obtain *Inpp5d^{fl/fl}/Cx3cr1^{CreER/+}* mice,^{66,67} in which TAM-induced Cre expression results in chronic recombination in microglia, but not in transient myeloid cells. These *Inpp5d^{fl/fl}/Cx3cr1^{CreER/+}* mice were crossed with *PSAPP* mice to obtain *PSAPP/Inpp5d^{fl/fl}/Cx3cr1^{CreER/+}* and their WT, *Inpp5d^{fl/fl}/Cx3cr1^{CreER/+}* littermates.⁵³ Constitutive *Inpp5d* KO mice were generated by crossing *Inpp5d^{fl/fl}* animals to a germline Cre strain, CMV-Cre, then backcrossed to remove the CMV-Cre to obtain only *Inpp5d^{KO/WT}* mice. All mice were on the C57BL/6J background and experimental groups consisted of both sexes, unless otherwise noted. Spatial transcriptomic analysis was completed on brains from females.

4.2 | Tamoxifen preparation and treatment

Tamoxifen was dissolved in CO at a concentration of 20 mg/ml by shaking overnight at 37°C. For induction of Cre recombinase, 3-month-old male and female mice received daily intraperitoneal (i.p.) injections of either TAM at 75 mg/kg body weight or CO for 5 days.

4.3 | Tissue processing

Mice were anesthetized at 6 months of age with pentobarbital (30 mg/kg, i.p.) and transcardially perfused with 20 ml of ice-cold 0.1 M phosphate-buffered saline (PBS), pH 7.4. One hemisphere was immersion-fixed for 48 hours in 4% paraformaldehyde for immunohistochemistry analysis. The remaining hemisphere was dissected for isolation of hippocampus and cortex and snap-frozen for biochemical analysis.

4.4 | Immunohistochemistry

Fixed hemibrains were sectioned sagittally using a vibratome (Leica) at 30 μ m. Free-floating brain sections stained with 6E10 anti-A β antibody were pre-treated with 70% formic acid for 15 minutes. Sections were blocked for 1 hour in PBS containing 0.2% (v/v) Triton-X and 5% normal goat (or donkey) serum (NGS), then incubated overnight at 4°C in primary antibody in PBS-0.2% Triton-X with 1% NGS. The tissues were subsequently washed in PBS-0.2% Triton-X, then incubated in PBS with 1% NGS and Alexafluor-conjugated secondary antibodies for 2 hours. Sections were then counterstained

as necessary and mounted onto slides. For methoxy-XO4 staining, mounted sections were dried at room temperature, then incubated sequentially (5 minutes per incubation) in PBS, 40% ethanol (EtOH), 1 μ M methoxy-XO4 (Cayman Chemical) in 40% EtOH, and then in 90% EtOH before coverslipping. To perform 3,3'-diaminobenzidine (DAB) staining for CST7, sections were washed briefly in PBS, then underwent antigen retrieval using 1x Reveal Decloaker (Biocare Medical) at 80°C for 15 minutes before being allowed to cool for 5 minutes. Sections were washed again in PBS before quenching with 3% H₂O₂, followed by permeabilization with PBS-0.2% Triton-X. Sections were blocked with 5% NGS in PBS-0.2% Triton-X for 1 hour, then incubated overnight at 4°C in primary antibody in PBS-0.2% Triton-X with 1% NGS. The tissues were subsequently washed in PBS-0.2% Triton-X, then incubated in PBS with 1% NGS and biotinylated secondary antibody. Sections were washed in PBS, then the signal was amplified using the ABC system (VECTASTAIN Elite ABC kit, Vector Laboratories), as per the manufacturer's instructions. The signal was then developed with a DAB Reagent (KPL), as per the manufacturer's instructions. Sections were then mounted, dried, and dehydrated before coverslipping and imaging. Adjacent sections were used to perform 6E10 immunostaining with DAB detection, with the sections quenched as before, then treated with 70% formic acid for 15 minutes. All steps were then performed as listed previously, starting from the permeabilization step. The tissues were stained with the following primary antibodies: rabbit anti-IBA1 (1:500; Cat. # 019-19741, Wako), mouse 6E10 anti-A β (1:1000; Cat. #9320-500, Covance), chicken anti-GFAP (1:1000; NBP1-05198), and mouse anti-Cst7 (1:200; antibody F010; a gift from Dr. Adriano Aguzzi). Secondary antibodies included: goat anti-mouse immunoglobulin (IgG) AlexaFluor 594 or 350 (1:1000; Invitrogen), goat anti-rabbit IgG AlexaFluor 488 (1:1000; Invitrogen), goat anti-chicken 594 (1:1000; Invitrogen), and biotinylated goat anti-mouse (1:1000, Vector Laboratories). Images were acquired on a Biotek Cytation 5 Multimode Reader (Agilent) with the same gains and exposures across all animals for each stain. Automated quantification of areas and cell counts was performed using the Biotek Gen5 software (Agilent) by creating an object mask with thresholds set for intensity and object size. For total microglia counts, adjustments were performed for plaque-associated microglia based on 4',6-diamidino-2-phenylindole (DAPI) number. Confocal images were acquired using a Confocal Zeiss LSM 510. Images were converted to Z Projects in Fiji (National Institutes of Health) based on sum intensity. Plaque-associated IBA1⁺ microglia were manually counted from Z Projects based on contact with a 6E10⁺ immunoreactive deposit.

4.5 | Western blot

Protein was extracted from cortical tissue by homogenization in RIPA buffer (Thermo Fisher Scientific) containing phosphatase/protease inhibitor cocktail (Thermo Fisher Scientific) and centrifuged at 15,000 $\times g$ for 20 minutes at 4°C before collecting the supernatant. A 50 μ g aliquot of total protein was separated by electrophoresis in precast 4% to 12% Bis-Tris Gels (Bio-Rad) and transferred to nitrocellulose membranes. Membranes were blocked in 5% milk before probing with the following antibodies: rabbit anti-SHIP1 (1:1000, Cell Signaling, D1163), rabbit anti- β -actin (1:2500, Sigma, A-2066), mouse anti-PSD95 (clone 6G6-1C9, 1:2500, EMD Millipore, MAB1596), and rabbit anti-synaptophysin (1:1000, Abcam, ab16659). The secondary antibody used was peroxidase-labelled anti-rabbit IgG

(1:5000, Vector Laboratories). SuperSignal West Femto ECL (Pierce, #34096) was used to reveal the immunoreactive proteins, and images were acquired using a Fujifilm ImageReader LAS-4000. Luminescent immunoreactive protein bands were quantified using Fiji software (ImageJ).

4.6 | Spatial transcriptomics

Spatial transcriptomics analysis was performed using the 10xGenomics' Visium platform following the manufacturer's instructions, as we have reported.²⁸ In short, fresh frozen brain sections from *PSAPP/Inpp5d^{fl/fl}/Cx3cr1^{CreER+}* and *Inpp5d^{fl/fl}/Cx3cr1^{CreER+}* mice either treated with CO or TAM were cryosectioned coronally to 10 μm and mounted on Visium capture areas ($n = 3$ for each condition, 12 sections total). After methanol fixation at -20°C for 30 minutes, sections were stained with hematoxylin and eosin, imaged on a Keyence BZ-X710 using a 20x objective, and tiled using BZ-X710 software. Sections were then enzymatically permeabilized for 12 minutes, and spatial barcodes and unique molecular identifiers were added to the captured polyA mRNA. The 12 Illumina libraries were generated using Dual Index primers and sequenced on a NovaSeq6000 using a S4 200 cycle flow cell v.1.5. The sequencing output was aligned to the mouse reference genome mm10 using SpaceRanger.

The following libraries were used to analyze and visualize the data in R (4.1.0): *Seurat* (4.0.6), *SingleCellExperiment* (1.16.0), *BayesSpace* (1.2.1),⁶⁸ *EnhancedVolcano* (1.10.0), *ComplexHeatmap* (2.10.), *viridis* (0.6.2), *muscat* (1.5.2),²⁹ *ggplot2* (3.3.5), *dplyr* (1.0.7), *purrr* (0.3.4), *scatter* (1.22.0), *UpSetR* (1.4.0), *ggraph* (2.0.5), *igraph* (1.2.11), *gtools* (3.9.2), *cowplot* (1.1.1), *ggpubr* (0.4.0), *NICHES* (0.0.2),³¹ *CIDER* (0.99.0). *SCTransform* was applied to each sample separately before merging. For all downstream clustering analyses, variable features from all 12 samples were used. Thirty principal components and a resolution of 0.8 were used to cluster the spots across all samples. *Seurat's FindAllMarker* function was used to calculate enriched genes in each cluster and *muscat* was used to calculate cluster-resolved differentially expressed genes across conditions using *edgeR* on the sum of counts per cluster. *BayesSpace*⁶⁸ was used to identify sub-spot level gene expression. *NICHES* was used to calculate ligand-receptor pair enrichment using predefined clusters and the *fantom5* database.³¹ *Seurat's AddModuleScore* was used to visualize groups of genes in Visium brain sections.

4.7 | Network analysis

To verify if the present mouse model gene signature mirrored that in the human AD gene network, we used the BN we previously constructed from the parahippocampal gyrus region of the Mount Sinai Brain Bank AD cohort.^{32,69} The cluster signatures and cluster-specific DEGs from the spatial transcriptomics analysis were intersected with the network neighborhood within a path length of up to six steps from *INPP5D* in the human AD BN. *P* value significance of the intersections at different path lengths was computed by hypergeometric test in R (v4.0.5). A subnetwork up to four steps from *INPP5D* was visualized by *Cytoscape* (v3.9.0).

4.8 | RNA in situ hybridization

RNA in situ hybridization (ISH) was performed using the RNAscope Multiplex Fluorescent kit (ACD) as before with slight modification.⁴⁹ The same hemibrains collected for spatial transcriptomics were sectioned at 10 μm on a cryostat and mounted on glass slides. Slides were fixed in 4% paraformaldehyde for 15 minutes at 4°C. After fixation, sections were dehydrated and stained with methoxy-XO4 by immersion in 50% EtOH for 2 minutes, 50% 5 μM methoxy-XO4 (Cayman Chemical) in 50% EtOH for 2 minutes, then 50% EtOH for 1 minute, before proceeding through 70% EtOH and two rounds of 100% EtOH for 5 minutes each. The mounted sections were quenched with H_2O_2 for 10 minutes then protease-permeabilized for 20 minutes at room temperature using reagents provided in the kit (Protease IV). Brain sections were then incubated for 2 hours at 40°C with probes followed by signal amplification steps. When the ISH assay was complete, slides were incubated with fluorophore-conjugated secondary antibody for 30 minutes, then counterstained with DAPI. The following probes were used: *Cst7*-C2 (Cat. #498711-C2) and *Aif1*-C4 (Cat. #319141-C4).

4.9 | A β assays

Tissue was processed via serial detergent fractionation with ultracentrifugation to produce TBS-soluble, Triton-X-soluble, and formic-acid-soluble fractions as described previously.⁵⁴ In brief, tissue was homogenized in ice-cold TBS containing protease/phosphatase inhibitor cocktail, then ultracentrifuged at $100,000 \times g$ for 1 hour at 4°C. The supernatant containing the TBS-soluble fraction was collected, and the pellet was subsequently homogenized in Triton-X solution (TBS with 1% [v/v] Triton-X-100) containing protease/phosphatase inhibitor cocktail and ultracentrifuged as before. The supernatant containing the Triton-X-soluble fraction was collected, and the pellet was homogenized in 70% formic acid before the ultracentrifugation was repeated. The resulting formic acid fraction was neutralized with 1 M Tris buffer. To quantify A β levels, human/rat A β 1-40 and 1-42 ELISA kits (Wako, Cat. #294-64701 and #290-62601) were used according to the manufacturer's instructions. Absolute concentrations of A β were normalized to initial tissue weight.

For analysis of native oligomeric A β protein structure, 2 μl protein samples from the TBS-soluble fraction were spotted onto activated PVDF membrane (0.22 μm ; Millipore). Membranes were incubated with either rabbit pAb A11 (anti-prefibrillar oligomers; 0.5 $\mu\text{g}/\text{ml}$; gift from Charles Glabe), rabbit pAb OC (anti-fibrillar oligomers and fibrils; 0.25 $\mu\text{g}/\text{ml}$; gift from Charles Glabe), or mouse mAb NU-4 (anti-oligomers; 1 $\mu\text{g}/\text{ml}$; gift from William Klein, Northwestern University). Generation, purification, and characterization of A11, OC, and NU-4 have been described previously.^{70,71} Membranes were incubated with appropriate anti-mouse or -rabbit horseradish peroxidase-conjugated secondary antibody (1:20,000; Vector Laboratories) and developed as described in the western blot section. Normalization to total APP/A β signal was achieved by detection of human APP transgene metabolites with the anti-A β antibody 6E10 (1:1000, Covance).

4.10 | Purification of primary mouse microglia and phagocytosis assay

Heterozygous *Inpp5d*^{KO/WT} mice were crossed to obtain *Inpp5d*^{KO/KO} (KO) and *Inpp5d*^{WT/WT} mice for isolation of primary microglia. Primary microglia were isolated from

cerebral cortices and hippocampi from postnatal day P0-P3 mice as described previously⁷² with slight modification. In brief, tissue was homogenized in ice-cold Hibernate A (BrainBits), then centrifuged at $300 \times g$ for 5 minutes at 4°C. The pellet was resuspended in high-glucose Dulbecco's Modified Eagle Medium (DMEM) supplemented with 10% heat-inactive fetal bovine serum (FBS; Gibco), 2 mM glutamine, penicillin/streptomycin (100 U/ml and 0.1 mg/ml, respectively) and seeded in poly-L-lysine precoated flasks. Cells were maintained at 37°C and 5% CO₂, during which media was changed twice a week. Microglia were collected after approximately 8 to 10 days by agitating the flasks on a orbital shaker at $\sim 1 \times g$ (180 rpm) for 30 minutes to detach microglial cells from the astrocytic monolayer.

Green fluorescent latex beads (Sigma #L1030) were used to assess phagocytosis in the primary microglia. Beads were pre-opsionized in heat inactivated FBS (Gibco) for 1 hour at 37°C before use.⁷² Cells were submitted to serum deprivation for 1 hour prior to addition of the beads, and then fluorescent latex beads were added to the media for an additional 1 or 4 hours. The final concentrations for beads and FBS in serum-free DMEM were 0.01% (v/v) and 0.05% (v/v), respectively. Cultures were then washed three times with PBS and fixed in 4% paraformaldehyde before cells were stained with anti-IBA1 and DAPI. Images were obtained using a Biotek Cytation 5 Multimode Reader (Agilent) to generate (at minimum) a 3×3 tiled montage of the well. Automated quantification of primary microglia number and dual labeling was performed using the Biotek Gen5 software (Agilent) by creating an object mask with thresholds set for intensity and object size for IBA1⁺ cells with a secondary mask to count the subpopulation of dual-labeled cells for their content of green fluorescent beads. Quantification was performed from three to four wells per group; data points represent experimental mean normalized to *Inpp5d*-WT with $N = 3$ independent experiments.

4.11 | Human brain information and experiments

For human AD gene network analysis, we used the multi-Omics data we previously generated from a set of 364 well-characterized AD and control brains spanning the full spectrum of LOAD-related cognitive and neuropathological disease severities represented in the Mount Sinai Brain Bank AD cohort. Details about the cohort population information, generation of -Omics data (including WGS and RNA sequencing), and data preprocessing are described in Wang et al.⁶⁹

4.12 | Statistical analyses

All data, excluding spatial transcriptomics results, were analyzed with GraphPad Prism 9 and are presented as means \pm standard error of the mean. Sample sizes and statistical tests are indicated in figure legends. Analyses used include one-way analyses of variance, and Student *t* test.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Data availability

All sequencing data generated in association with this study are available on our searchable website portal https://liddelowlab.shinyapps.io/Hasel_Castranio_AD_Spatial/. Raw data are available in the Gene Expression Omnibus (GSE No. 203424).

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RESEARCH IN CONTEXT

5. **Systematic review:** The authors reviewed previous publications using traditional sources, including PubMed and preprint server databases. The mechanism of *INPP5D*/SHIP1 has been more frequently studied in the periphery. There are relatively few peer-reviewed publications regarding *INPP5D*/SHIP1 biology in the brain and/or within the context of Alzheimer's disease (AD). We have cited relevant publications.
6. **Interpretation:** Our results show that conditional *Inpp5d* downregulation increases plaque deposition, microglial recruitment to plaques, and plaque-induced gene expression during early stages in disease development in the *APP^{KM670/671NL}/PSEN1^{exon9}* (PSAPP) mouse model of AD.
7. **Future directions:** Our observation that specific knockdown of *Inpp5d* in the microglia of the PSAPP mouse model of cerebral amyloid beta amyloidosis causes complex changes in the neuropathological features associated with human AD should provide context for further investigation of its biology. These insights should help clarify role(s) for *INPP5D* in the brain and in AD that have diagnostic and/or therapeutic implications.

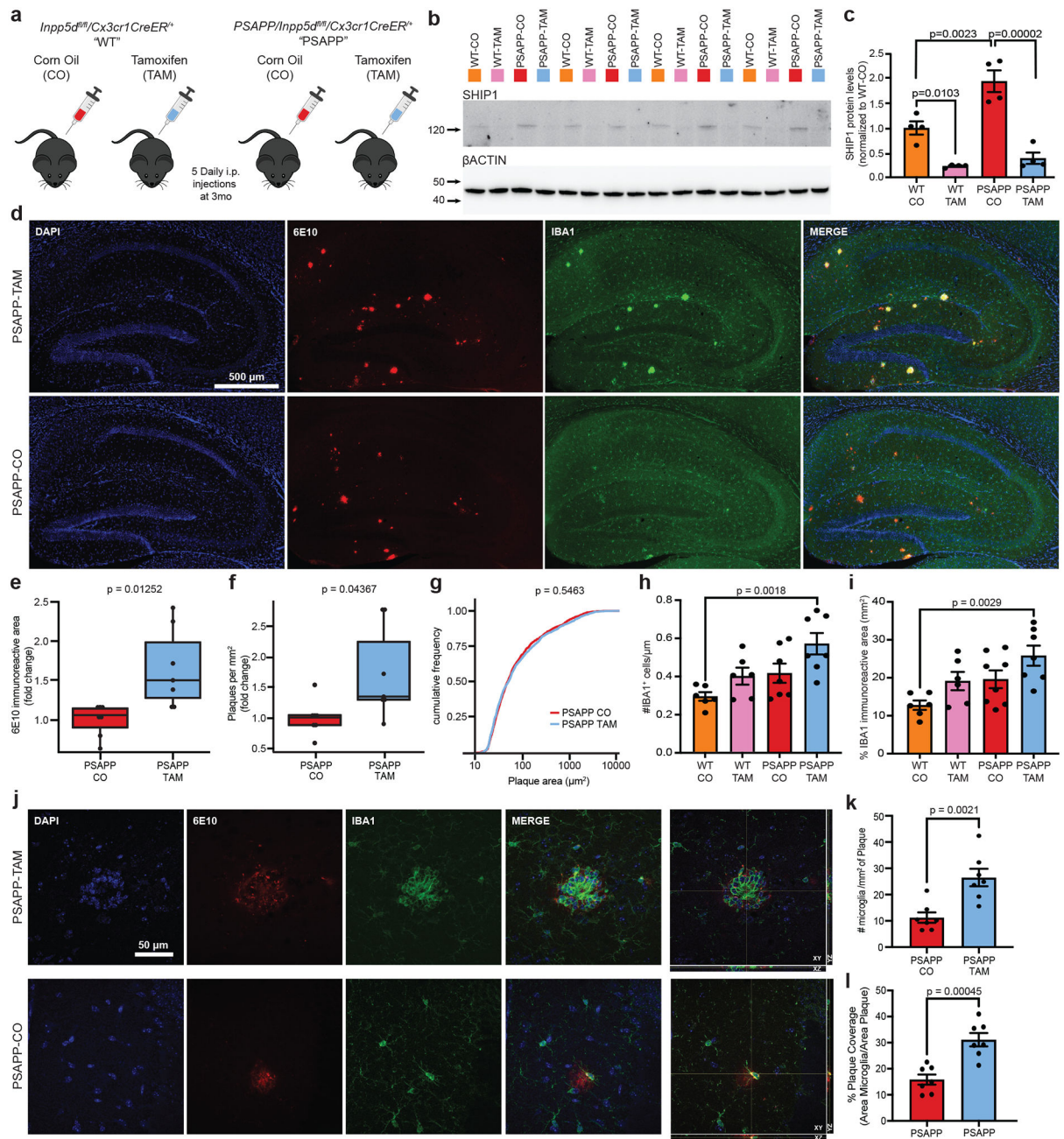


FIGURE 1. *Inpp5d*/SHIP1 knockdown increased amyloid pathology and plaque-associated microglia number.

A, *Inpp5d^{fl/fl}/Cx3cr1CreER⁺* mice were crossed with the amyloid-depositing *APP^{KM670/671NL}/PSEN1^{exon9}* mouse model of Alzheimer's disease (AD) to obtain either heterozygous PSAPP mice, that is, *PSAPP/Inpp5d^{fl/fl}/Cx3cr1CreER⁺* (PSAPP) or wild-type littermates *Inpp5d^{fl/fl}/Cx3cr1CreER⁺* (WT). At age 3 months, male and female mice of both genotypes received daily injections of tamoxifen (TAM) or corn oil (CO) for 5 days before aging to the experimental timepoint of ages 6 to 6.5 months. B, Western blot of hippocampal tissue at 6 to 6.5 months of age using anti-SHIP1 (D1163) antibody normalized to β -actin. Quantification in (C). D, Representative images from 6-month PSAPP-TAM (top)

and PSAPP-CO (bottom) mice stained with 6E10 (amyloid, red) or anti-IBA1 (myeloid cells, green). Scale bar = 500 μm . E, Relative fold change of 6E10-immunoreactive area in the hippocampi of male and female PSAPP-CO and PSAPP-TAM mice ($n = 7$ mice per group). F, Relative fold change of plaque numbers per mm^2 in the hippocampi of male and female PSAPP-CO and PSAPP-TAM mice ($n = 7$ mice per group). For both (F) and (G), fold changes were calculated separately for each sex. G, Cumulative frequency of plaque areas of male and female PSAPP-CO and PSAPP-TAM mice (plaques from 8 mice per group). H, Quantification of number of IBA1⁺ microglia and (I) area of IBA1⁺ area in hippocampus from WT-CO, WT-TAM, PSAPP-CO and PSAPP-TAM ($n = 6$ to 7 mice per group). J, Confocal Z-stack and orthogonal views of plaques from PSAPP-TAM and PSAPP-CO mice. Quantification of plaque-associated microglia shown in (K). Scale bar = 50 μm . L, Quantification of plaque coverage by microglia (% area of IBA1 staining overlapping area of 6E10⁺ plaques) for PSAPP-CO and PSAPP-TAM ($n = 7$ mice per group). Bar graphs represent the mean and error bars represent standard error of the mean. *P*-values were calculated using unpaired t test (B, E, F, K, L), one-way analysis of variance with Tukey's multiple comparison (C H, I), or by a Kolmogorov–Smirnov test (G)

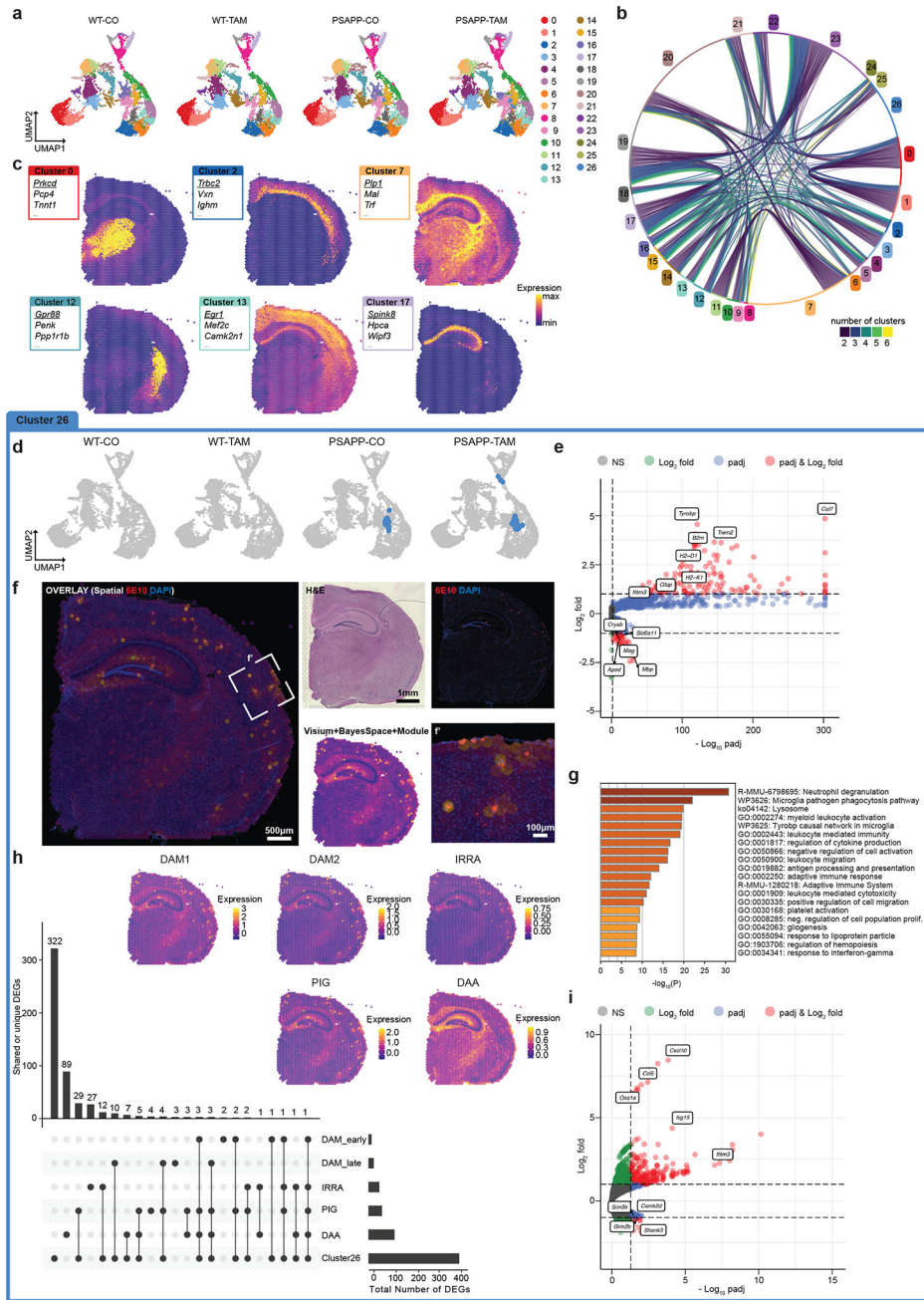


FIGURE 2. Spatial transcriptomics identified plaque-specific gene expression profiles exacerbated in *Inpp5d*/SHIP1 knockdown animals.

A, Uniform manifold approximation and projection (UMAP) dimensional reduction of wild-type (WT)-corn oil (CO) (8037 spots, 3 sections), WT-tamoxifen (TAM; 8882 spots, 3 sections), *PSAPP/Inpp5d^{fl/fl}/Cx3xrf^{CreER+}* (PSAPP)-CO (7748 spots, 3 sections), and PSAPP-TAM (9366 spots, 3 sections). Each dot represents a Visium transcriptomics spot. The 27 color-coded clusters correspond to brain areas/regions (see below). B, Chord diagram of genes enriched > 1 log₂ fold (12f) in each cluster and adjusted *P*-value < 0.05. Each string represents a gene shared between two and six other clusters. C, Example clusters highlighted in a Visium brain section. Boxes show top three enriched genes per

cluster. Underlined gene is highlighted in the SpatialFeaturePlot. D, UMAP with Cluster 26 highlighted showing that it is only present in PSAPP animals. E, Volcano plot of genes enriched in Cluster 26 as calculated with *Seurat*'s FindAllMarkers function. Red genes are $l2f > 1$ and $padj < 0.05$, blue $l2f < 1$ and $padj < 0.05$, green genes are $l2f > 1$ and $padj > 0.05$, gray genes are $l2f < 1$ and $padj > 0.05$. F, Module score of Cluster 26 and 6E10 (amyloid) staining of the adjacent brain section shows colocalization. f' shows higher magnification of box in (F). G, Gene Ontology term analysis of genes enriched $l2f > 1$ in Cluster 26 using metaspice.⁷³ H, UpSet plot highlighting the overlapping and unique genes between previously defined reactive microglia and astrocyte subtypes: Disease-associated microglia, DAM1 (early) and DAM2 (late),²⁵ interferon-responsive reactive astrocytes (IRRA),²⁸ plaque-induced genes (PIGs),²⁷ as well as disease associated astrocytes (DAA).²⁶ Insets are module scores of the genes in each reactive subtype highlighted on a PSAPP-TAM spatial transcriptomics brain section. For IRRA, DAA and Cluster 26, genes were filtered for $padj < 0.05$ and $l2f > 0.5$. For DAM1, DAM2, and PIGs, genes were selected based on gene sets provided in the original publication. I, Genes differentially expressed in Cluster 26 between APP-CO and APP-TAM as calculated with *edgeR* on sum of counts using *muscat* (red genes are $l2f > 1$ and $padj < 0.05$, blue $l2f < 1$ and $padj < 0.05$, green genes are $l2f > 1$ and $padj > 0.05$, gray genes are $l2f < 1$ and $padj > 0.05$).

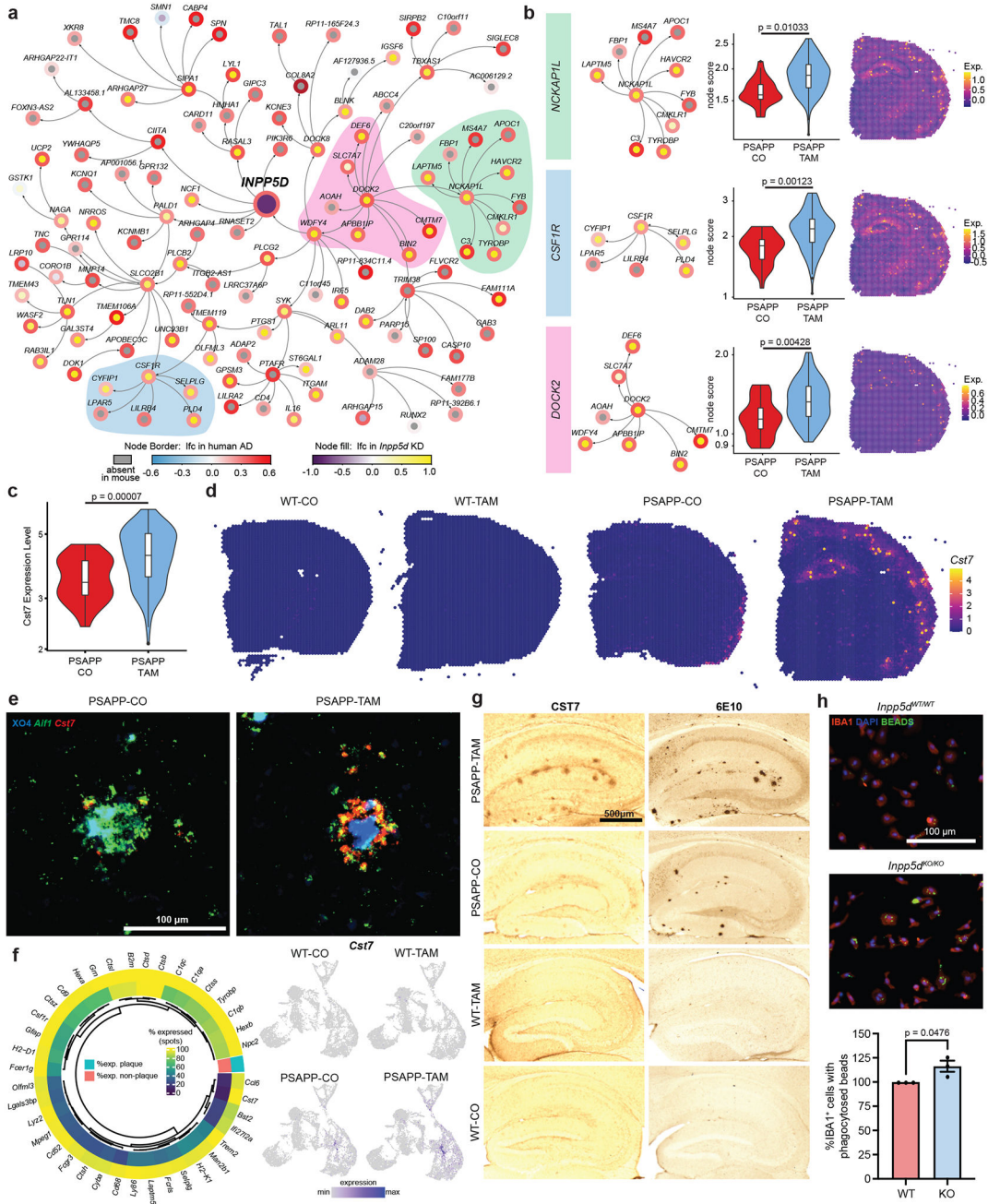


FIGURE 3. Top Cluster 26 differentially expressed gene *Cst7* was a plaque-specific marker and was exacerbated by *Inpp5d* knockdown.

A, Network showing the gene expression change in human Alzheimer’s disease (AD) and *Inpp5d* knockdown mice within the neighborhood of four steps of *INPP5D*. Node border paint denotes the log₂ fold-change (lfc) in human AD brains compared to control brains. Node fill paint denotes the log₂ fold-change caused by *Inpp5d* in mice, with gray color denoting absence in mouse data. B, Highlight of specific nodes from network in (A). Genes associated with each node were enriched in cluster 26 from *PSAPP/Inpp5d*^{fl/fl}/*Cx3xr1*^{CreER+} (PSAPP)-tamoxifen (TAM) compared to PSAPP-corn oil (CO) mice (violin plots, middle, *P*-values from unpaired *t* test) and can be spatially resolved by probing for this module in

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spatial transcriptomic brain sections (right). C, Violin plots of *Cst7* in Cluster 26 across PSAPP-CO and PSAPP-TAM shows increased expression in PSAPP-TAM plaque spots as calculated with *edgeR* on sum of counts using *muscat*. D, Cluster 26 marker gene *Cst7* highlighted in wild-type (WT)-CO, WT-TAM, PSAPP-CO, and PSAPP-TAM brain sections. E, RNAScope of *Cst7* (red) and *Aif1* (green) with Methoxy-XO4 (blue) staining in PSAPP-CO and PSAPP-TAM mouse cortex. Scale bar = 100 μm . F, Left: Percentage expression of the top Cluster 26 enriched genes in Cluster 26 (outer ring) versus the average percentage expression in all other clusters (inner ring). Right: FeaturePlots showing expression of *Cst7* in all spots across WT-CO, WT-TAM, PSAPP-CO, and PSAPP-TAM brain sections. G, CST7 antibody staining detected with DAB and 6E10 in adjacent section from WT-CO, WT-TAM, PSAPP-CO, and PSAPP-TAM hippocampus. Scale bar = 500 μm . H, Representative images of latex bead phagocytosis assay on *Inpp5d*-WT mouse primary microglia and *Inpp5d*-knockout (KO) primary microglia with quantification of percentage IBA1⁺ cells with phagocytosed fluorescent bead (right). Scale bar = 100 μm . Quantification was performed from three to four wells per group; with $N = 3$ independent experiments. Data points represent mean normalized to *Inpp5d*-WT per experiment. P -values calculated with unpaired t test.