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# Integrin-deficient T cell leukemia accumulates in the central nervous system

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## eLife Assessment

This study presents an **important** finding that loss or blockade of key integrins unexpectedly enhances central nervous system accumulation of T-cell acute lymphoblastic leukemia cells and may increase their sensitivity to chemotherapy. The evidence is **convincing**, supported by well-designed in vivo models, CRISPR-based perturbations, competitive assays, imaging, and complementary therapeutic experiments. However, the mechanistic basis linking integrin loss, altered spatial distribution, and increased proliferation remains **incompletely** defined, and the translational implications would be strengthened by additional survival studies and validation in more clinically relevant models.

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## Abstract

T-cell acute lymphoblastic leukemia (T-ALL) spreads aggressively to the central nervous system (CNS), particularly the leptomeninges. Children with T-ALL are treated with high-dose, CNS-directed chemotherapy, which can cause lasting neurotoxicity and is not always effective. Little is known about how T-ALL enters and persists within the CNS. However, normal T cell migration into the CNS has been extensively studied. Two integrins—VLA-4 and LFA-1—mediate normal T cell entry to the CNS, and VLA-4 blockade effectively treats multiple sclerosis by excluding T cells from the brain. We hypothesized that these integrins would likewise be required for T-ALL CNS entry. Unexpectedly, not only were VLA-4 and LFA-1 dispensable for T-ALL to reach the CNS, integrin-deficient T-ALL accumulated in the CNS compared to control. Mechanistically, integrin loss accelerated T-ALL proliferation in the CNS, suggesting that integrin-mediated interactions may promote quiescence in this space. Integrin blockade synergized with chemotherapy targeting proliferating cells, raising the possibility that combination therapy might be a powerful strategy.

## Introduction

Acute lymphoblastic leukemia (ALL) is the most common pediatric cancer [1, 2]. In recent years, major advances have improved the five-year survival rate to over 90% [2]. However, ALL's propensity to infiltrate the central nervous system (CNS), particularly the meninges, remains a significant challenge [3, 4]. To eradicate disease, the standard-of-care is to treat all children diagnosed with ALL with high-intensity, CNS-directed chemotherapy [3, 5]. This regimen induces substantial toxicity and can cause irreparable neurological damage. Furthermore, despite aggressive treatment, the CNS is a frequent site of relapse. Our lack of understanding of how ALL enters and persists in the CNS has limited the development of targeted therapies for decades.

The T cell form of ALL, T-ALL, has especially high rates of CNS penetration [6]. Very little is known about how T-ALL infiltrates the CNS, but normal effector T cell trafficking into the CNS has been extensively characterized. T cells cross the blood-endothelial barrier into the CNS through a stepwise process of rolling, sticking, and transmigration [7]. This cascade requires integrins—heterodimeric adhesion molecules consisting of an alpha and beta chain—which allow T cells to attach firmly to the endothelium. The integrin VLA-4 ( $\alpha 4\beta 1$ ) is canonically required for T cells to enter the CNS [8, 9], and in some cases the integrin LFA-1 ( $\alpha L\beta 2$ ) can contribute to this transmigration [7]. In fact, natalizumab, a monoclonal antibody blocking the alpha subunit of VLA-4, is widely used to treat multiple sclerosis by preventing autoreactive T cells from reaching the brain [10]. Additionally, integrins enable B-cell ALL to travel to the meninges via bridging channels between the skull bone marrow and the dura mater [11]. We hypothesized that blocking VLA-4 and LFA-1 would render T-ALL unable to metastasize to the CNS.

## Results

### T-ALL deficient in LFA-1 and VLA-4 preferentially accumulates in the CNS

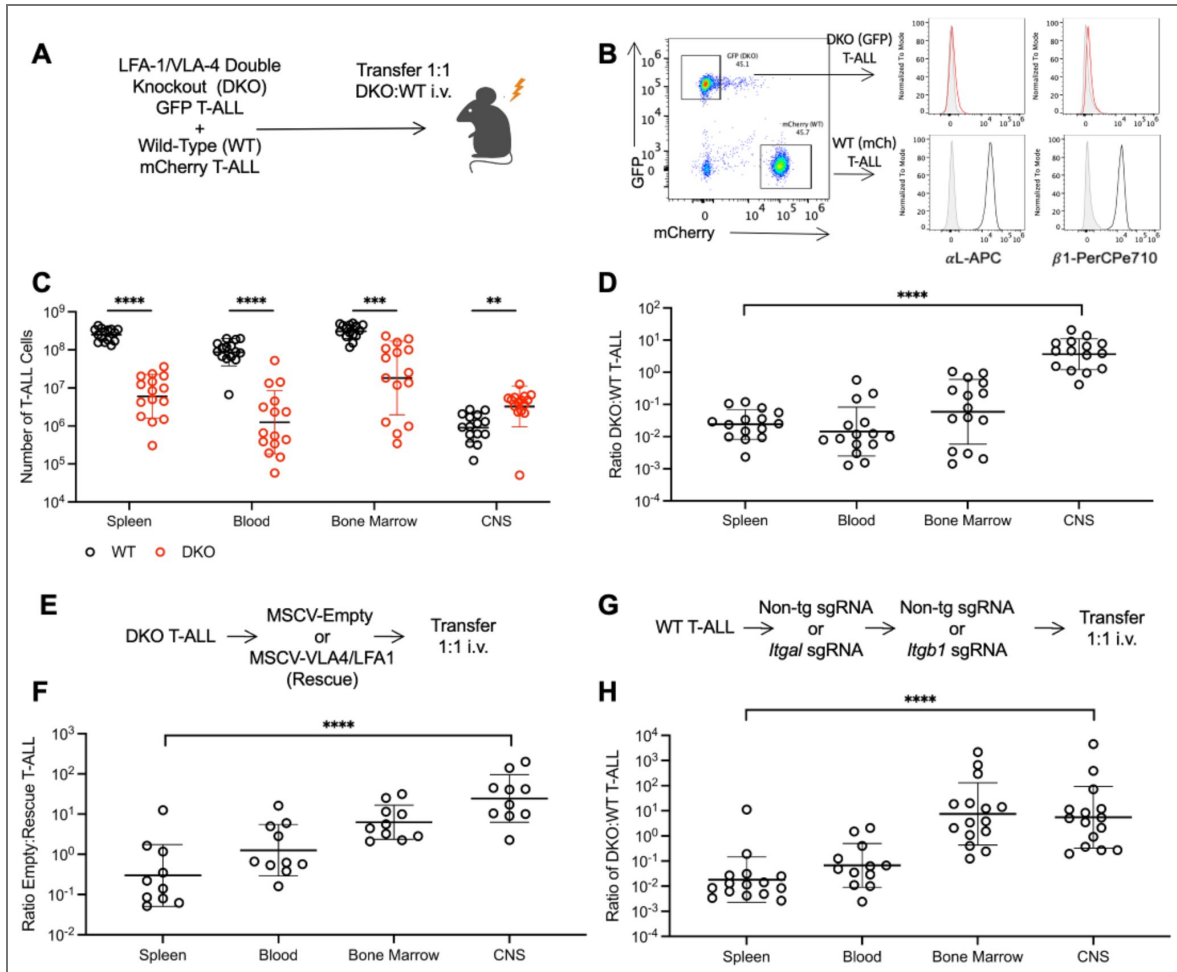
To test whether T-ALL could enter the CNS in the absence of LFA-1 and VLA-4, we developed a T-ALL line by transducing the bone marrow from mice that lacked these integrins (double-knockout, DKO) with a constitutively active NOTCH-1, per an established protocol [12] (S. Fig. 1A [12]). We chose a NOTCH1-driven model because mutations in NOTCH1 and its regulators are found in the majority of human T-ALL cases [13–15]. In the integrin-deficient T-ALL cells, the first and second exons of *Itgal* ( $\alpha L$ , the alpha subunit of LFA-1) were knocked out, and the third exon of *Itgb1* ( $\beta 1$ , the beta subunit of VLA-4) was excised [16, 17]. Of note, B-cell ALL requires  $\alpha 6$  integrin to enter the CNS, which we could not detect on the surface of DKO T-ALL (S. Fig. 1B,C [16]). This is unsurprising as  $\beta 1$  integrin is a canonical partner of  $\alpha 6$ , and we could not detect mRNA or protein for the other canonical binding partner of  $\alpha 6$  integrin,  $\beta 4$  integrin, in these cells (S. Fig. 1B,C [16]; S. Table 1 [16]) [11, 18].

We co-transferred the DKO T-ALL cells at a 1:1 ratio with similarly generated NOTCH1-driven, mCherry<sup>+</sup>, wild-type (WT) T-ALL cells intravenously into sub-lethally irradiated WT adult mice (Fig. 1A,B [16]). At a humane endpoint, we enumerated DKO and WT T-ALL cells in common sites of disease (Fig. 1C,D [16]). As in human patients, CNS T-ALL cells in our model were largely confined to the leptomeninges rather than the brain parenchyma [4, 19]; washing the brain surface yielded 10–20 times more cells than extracting T-ALL cells from the brain parenchyma, and our quantification reflects leptomeningeal disease (S. Fig. 1D [16]). Surprisingly, in the CNS we found nearly 10 times more integrin-deficient T-ALL cells than WT T-ALL cells. This was especially notable considering the DKO T-ALL's relative absence in the spleen, blood and bone marrow, natural reservoirs for leukemia cells (Fig. 1C,D [16]).

To determine whether DKO T-ALL accumulation in the CNS was solely due to the absence of LFA-1 and VLA-4, we tested whether restoration of VLA-4 and LFA-1 in DKO T-ALL would reverse its accumulation in the CNS. We transduced the DKO T-ALL line with vectors encoding *Itgal* and *Itgb1* or an empty vector to generate “rescue” and “empty” lines (Fig. 1E [16]). We found that restoration of  $\alpha L$  and  $\beta 1$  in the DKO T-ALL (DKO-rescue) reduced the number of cells accumulating in the CNS compared to vector-transduced cells (DKO-empty) (Fig. 1F [16]).

To further test whether DKO T-ALL accumulation in the CNS was solely due to the absence of LFA-1 and VLA-4, we used CRISPR-Cas9 to target the first through third exons of *Itgal* and the first and second exons of *Itgb1* in the WT T-ALL line (Fig. 1G [16], S. Fig. 1E,F [16]). These CRISPR-DKO T-ALL lines recapitulated the original phenotype, with CRISPR-DKO T-ALL preferentially accumulating in the CNS compared to control-targeted cells (CRISPR-WT) (Fig. 1H [16]).

We saw similar results when the DKO and WT T-ALL were transferred singly (S. Fig. 2A, B [16]). Additionally, irradiation of the CNS was not required for DKO T-ALL to accumulate in the CNS (S. Fig. 2C [16]). Taken together, these data suggest that the loss of VLA-4 and LFA-1 results in T-ALL



**Fig. 1. Integrin-deficient T-ALL accumulates in the CNS compared to WT T-ALL.**

(A-D) WT mCherry<sup>+</sup> T-ALL and DKO GFP<sup>+</sup> T-ALL cells were mixed 1:1 and transferred intravenously (i.v.) into sub-lethally irradiated recipients. (A) Experiment design. (B) Left: Representative flow cytometry plot distinguishing WT T-ALL and DKO T-ALL from the CNS. Right: Representative surface stain of  $\beta 1$  and  $\alpha L$  integrins on T-ALL. Black: WT, red: DKO, grey: isotype. Numbers (C) and ratios (D) of DKO (red) and WT (black) T-ALL cells found in the indicated tissues at a humane endpoint. Data compiled from 3 experiments;  $n = 15$ . (E-F) DKO T-ALL cells were transduced with either vectors encoding *Itgal* and *Itgb1* (“rescue”) or empty vector (“empty”) and transferred 1:1 i.v. into sub-lethally irradiated recipients. (E) Experiment design. (F) Ratio of DKO-empty to DKO-rescue T-ALL cells in the indicated tissues at a humane endpoint. Data compiled from two experiments;  $n = 10$ . (G-H) WT T-ALL cells were nucleofected with sgRNAs targeting *Itgal* or non-targeting gRNAs along with recombinant Cas9 protein. The process was repeated for *Itgb1* to generate “CRISPR-DKO” and “CRISPR-WT T-ALL.” The cells were mixed 1:1 and transferred i.v. into sub-lethally irradiated recipients. (G) Experiment design. (H) Ratio of CRISPR-DKO to CRISPR-WT T-ALL cells found in the indicated tissues at a humane endpoint. Data compiled from five experiments;  $n = 15$ .

accumulation within the CNS.

## Integrin loss is unlikely to induce T-ALL accumulation in the CNS by enhancing CNS entry, limiting CNS exit, or promoting immune escape

Having disproven our original, seemingly obvious, hypothesis, we sought to explain how loss of integrins resulted in T-ALL accumulation in the CNS. It was possible that loss of integrins paradoxically improved entry into the CNS, limited exit from the CNS, or promoted T-ALL survival or proliferation within the CNS.

### Entry: Premature release from bone marrow

One explanation for our results was that loss of integrins led to premature release of T-ALL cells from the bone marrow, a key T-ALL niche. Integrins retain normal hematopoietic progenitor cells in the bone marrow; when VLA-4 binding is blocked, these cells exit their niche and move into the blood [20–22]. We hypothesized that this may be occurring in T-ALL, thus allowing DKO to migrate earlier to the CNS and take over the leptomeningeal space. To test this, we conducted a time-course experiment, assessing the ratio of DKO:WT T-ALL in the CNS at different stages of disease.

Contrary to our expectation, we found that WT T-ALL accumulated in the CNS before DKO T-ALL. At around mid-stage disease, DKO T-ALL caught up, and at late-stage disease, DKO T-ALL overtook WT leukemia (Fig. 2A, B [23]).

### Entry: Up-regulation of other integrins

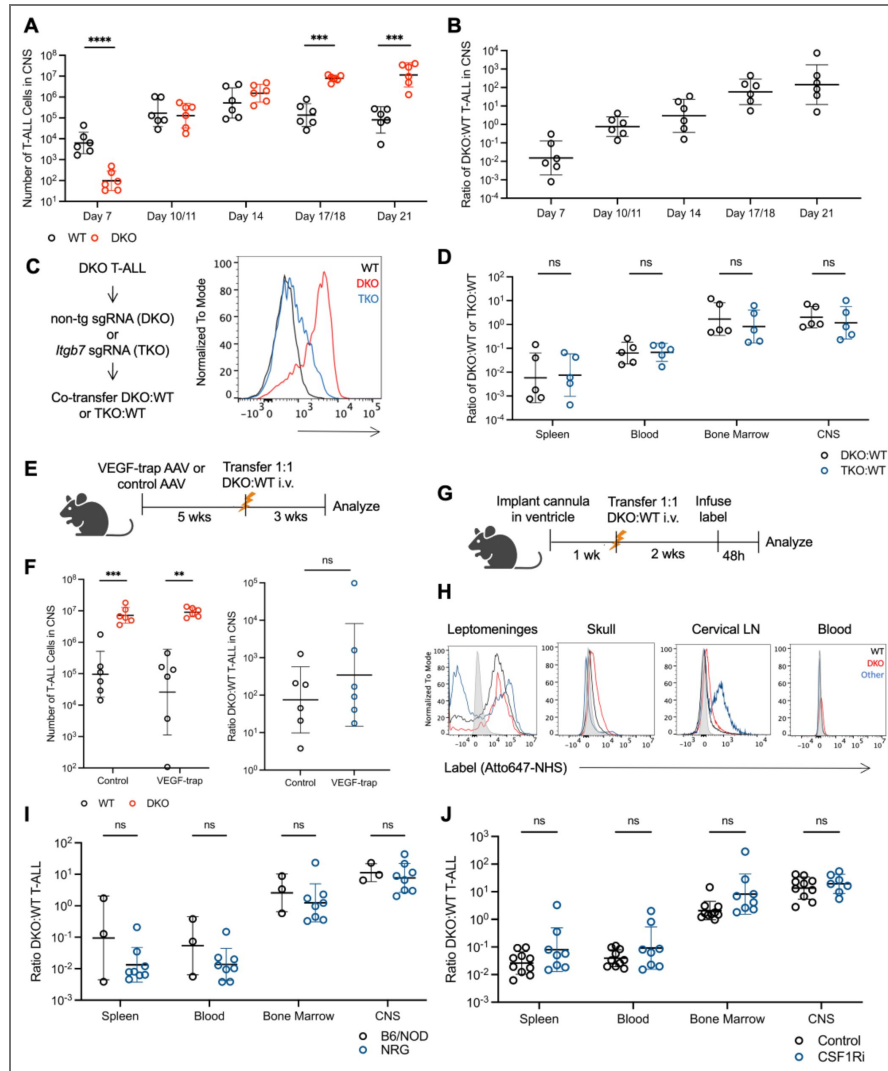
A second attractive explanation for our results was that loss of VLA-4 and LFA-1 resulted in compensatory expression of other integrins that were better matched to adhesion molecules expressed by CNS blood endothelial cells in the context of leukemia. In fact, when we performed bulk RNA sequencing comparing DKO and WT T-ALL isolated from the CNS, we found several differences in integrin expression between DKO and WT T-ALL (S. Fig. 2D,E [24]). Of these, only *Itgb7* was significantly increased in DKO compared to WT T-ALL. We confirmed elevated surface protein expression of  $\beta 7$  integrin (Fig. 2C [25]), which might reflect a combination of increased mRNA and increased availability of its binding partner  $\alpha 4$  integrin upon the loss of  $\beta 1$  integrin. The ligand for  $\alpha 4\beta 7$  integrin, MAdCAM1, has primarily been characterized for its role in T cell migration into the gut [23], but has also been found to be expressed in the choroid plexus during inflammation [24]. Furthermore,  $\alpha 4\beta 7$  integrin has been reported to facilitate T cell entry into the CNS in some settings [25].

To test whether DKO T-ALL was using  $\alpha 4\beta 7$  to cross into the CNS, we triply knocked out  $\alpha L$ ,  $\beta 1$ , and  $\beta 7$  (third exon) in T-ALL cells using CRISPR-Cas9 as described above (Fig. 2C [26]). We found that triple-knockout (TKO) T-ALL was indistinguishable from DKO T-ALL in its accumulation in the CNS (Fig. 2D [27]). These results suggest that integrin  $\alpha 4\beta 7$  is not responsible for the increased accumulation of DKO T-ALL within the CNS.

### Exit

We next hypothesized that DKO T-ALL may accumulate in the CNS due to a defective ability to exit the CNS. The dorsal lymphatics of the meningeal dura mater have been recently characterized as a key route of exit for lymphocytes from the CNS to the draining cervical lymph nodes [26, 27], and integrins in some cases enable T cells to access lymphatics [28, 29].

If DKO T-ALL accumulated in the CNS due to a failure to exit through the dorsal meningeal lymphatics, the ratio of DKO:WT T-ALL cells in the CNS should normalize in the absence of meningeal lymphatics. To ablate the meningeal lymphatics, we took advantage of the fact that, unlike most lymphatic vessels in the body, the dorsal meningeal lymphatics require continuous VEGFR3 signaling to persist [30]. We therefore treated mice with a “VEGF-trap,” an adeno-associated virus encoding the ligand-binding domain of VEGFR3 fused to IgG-Fc, which captures available VEGFC/D and induces dorsal meningeal lymphatic regression. As a control, we treated mice with an adeno-associated virus that encodes the stalk (non-ligand-binding) of VEGFR3 fused



**Fig. 2. Integrin-deficient T-ALL is unlikely to accumulate in the CNS due to enhanced entry, limited exit, or superior immune escape.**

(A, B) Sub-lethally irradiated WT mice were injected i.v. with 1:1 DKO:WT T-ALL and tissues were isolated from 3 mice every 3-4 days. (A) Total number of WT (black) and DKO (red) T-ALL cells in the CNS. (B) Ratio of DKO:WT T-ALL in the CNS. Data compiled from two experiments;  $n = 36$ . (C) Left, CRISPR-TKO T-ALL generation. Right, representative surface staining of integrin  $\beta 7$  on T-ALL cells (blue: TKO, black: WT, red: DKO). (D) Ratio of DKO:WT T-ALL (black) or TKO:WT T-ALL (blue) in indicated tissues at a humane endpoint. Data compiled from two experiments;  $n = 5$ . (E, F) WT mice were infected with AAV encoding a VEGF-trap or control AAV. 5 weeks later, mice were sub-lethally irradiated and injected i.v. with 1:1 DKO:WT T-ALL. At a humane endpoint, tissues were isolated and T-ALL was enumerated. (E) Experiment design. (F) Number of WT (black) and DKO (red) T-ALL or ratio of DKO:WT T-ALL in the CNS of mice treated with either control AAV or AAV-VEGF-trap. Data compiled from 2 experiments;  $n = 6$ . (G, H) WT mice were surgically implanted with an intra-cerebroventricular cannula. 1 week later, mice were sub-lethally irradiated and injected i.v. with 1:1 DKO:WT T-ALL. At mid-stage disease, a reactive label (Atto647-NHS-ester) was infused through the cannula into the left lateral ventricle. 48h later, tissues were isolated and labelled T-ALL cells were enumerated. (G) Experiment design. (H) Representative flow cytometry plots of Atto647 labelling in the indicated tissues. WT T-ALL, black; DKO T-ALL, red; other leukocytes, blue. Representative of 3 experiments;  $n = 6$ . (I) NRG mice or WT controls (B6 or NOD) were sub-lethally irradiated, injected i.v. with 1:1 DKO:WT T-ALL, and analyzed at a humane endpoint. Ratio of DKO:WT T-ALL cells in controls (black;  $n = 3$ ) or NRG mice (blue;  $n = 8$ ). Data compiled from 2 experiments. (J) WT mice were fed either diet supplemented with a CSF1R inhibitor (PLX5622) or control diet for 1 week, sub-lethally irradiated and injected i.v. with 1:1 DKO:WT T-ALL, and maintained on either PLX5622 or control diet until a humane endpoint. Ratio of DKO:WT T-ALL in the indicated tissues of mice fed control diet (black;  $n = 10$ ) or PLX5622 diet (blue;  $n = 8$ ). Data compiled from 2 experiments. Bars represent geometric mean and geometric SD. Statistics calculated on log-transformed data using unpaired t-test with Welch's correction. \* $p \leq 0.05$ , \*\* $p \leq 0.01$ , \*\*\* $p \leq 0.001$ , \*\*\*\* $p \leq 0.0001$

to IgG-Fc [30]. Five weeks later, we induced leukemia in these animals by co-transferring WT and DKO T-ALL (Fig. 2E). We found that the regression of meningeal lymphatics had no effect on the ratio of the DKO to WT T-ALL in the CNS, or on the overall number of cells in the CNS (Fig. 2F). Meningeal lymphatic regression was confirmed by microscopy (S. Fig. 3A). We found similar results when we repeated this experiment using mice that were developmentally lacking in meningeal lymphatics (mice expressing the VEGF-trap under the keratin-14 promoter) [27] (S. Fig. 3B,C). These data suggest that DKO T-ALL does not preferentially accumulate in the CNS due to a requirement for integrins to exit through the meningeal lymphatics.

There may be alternate, uncharacterized pathways of exit from the CNS that could require integrins. To address more generally whether DKO T-ALL cells were defective in CNS egress compared to WT T-ALL cells, we labeled T-ALL cells in the CNS and assessed their location 48 hours later. Mice were surgically implanted with an intraventricular cannula in the lateral ventricle of the CNS prior to leukemia induction. Once mice reached mid-stage disease, we infused a reactive, fluorescent dye into the CNS via the cannula (Fig. 2G). 48 hours later, we found that T-ALL cells in the CNS were efficiently labeled. We found both labeled and non-labeled normal leukocytes in the CNS and the draining cervical lymph nodes, suggesting that normal leukocytes were cycling through the CNS. However, we did not detect labeled T-ALL cells in any other site, including blood, skull bone marrow, or the CNS-draining cervical lymph nodes (Fig. 2H). Overall, we found no evidence that delayed exit causes accumulation of DKO T-ALL within the CNS.

### Survival: Immune escape

As differences in trafficking seemed unlikely to explain the CNS accumulation of the DKO T-ALL, we next hypothesized that differential interactions with the host immune system might be responsible. Integrins play a key role in cementing synapses between leukocytes [31], and cytotoxic T cells can have a powerful anti-tumor effect in ALL [32, 33]. Perhaps the lack of integrins enabled the DKO T-ALL to escape immune surveillance.

If reduced susceptibility to an anti-leukemia immune response explained DKO T-ALL accumulation within the CNS, the ratio of DKO to WT T-ALL in the CNS should normalize in an immune-deficient host. To test this, we co-transferred WT and DKO T-ALL cells into NOD-*Rag1*<sup>-/-</sup>*Il2rg*<sup>-/-</sup> (NRG) mice, which lack B, T, and NK cells, but found no difference in the ratio of the DKO:WT T-ALL in the CNS between the NRG mice and either WT C57BL/6 or NOD controls (Fig. 2I). Interestingly, the B6-derived leukemia was frequently cleared entirely in immune-competent NOD hosts (not shown). T-ALL cells also interact with myeloid cells, although integrin-mediated interactions with myeloid cells promote T-ALL progression in many tissues [34, 35]. We tested a role of myeloid cells in the CNS accumulation of DKO cells by treating mice with the colony-stimulating factor 1 receptor (CSF1R) inhibitor PLX5622, which has been widely used to deplete microglia [36] as well as meningeal macrophages [37, 38], prior to induction of leukemia. We again found no difference in the ratio of DKO:WT T-ALL in the CNS between mice treated with the CSF1R inhibitor and controls (Fig. 2J). Consistent with previous findings that myeloid cells support T-ALL persistence in the CNS, we found that treatment with the CSF1R inhibitor resulted in an overall reduction in the number of T-ALL cells (S. Fig. 3D-F) [39]. More generally, we did not observe a difference in the rate of apoptosis between DKO and WT T-ALL in the CNS, as assessed by flow cytometry of cells stained for cleaved caspase 3 (S. Fig. 3G,H). Taken together, these data suggest that DKO T-ALL is not accumulating in the CNS because of its avoidance of an anti-leukemia immune response.

## The loss of integrins in T-ALL confers a proliferation advantage in the CNS

### DKO T-ALL proliferates more rapidly than control T-ALL in the CNS

Within the dura mater, we observed that the DKO and WT T-ALL localized differently—the DKO cells tended to spread through the tissue while the WT cells tended to cluster near blood vessels (Fig. 3A). This is consistent with observations of increased effector T cell motility within the spinal cord meninges upon intrathecal injection of integrin-blocking antibodies [40].

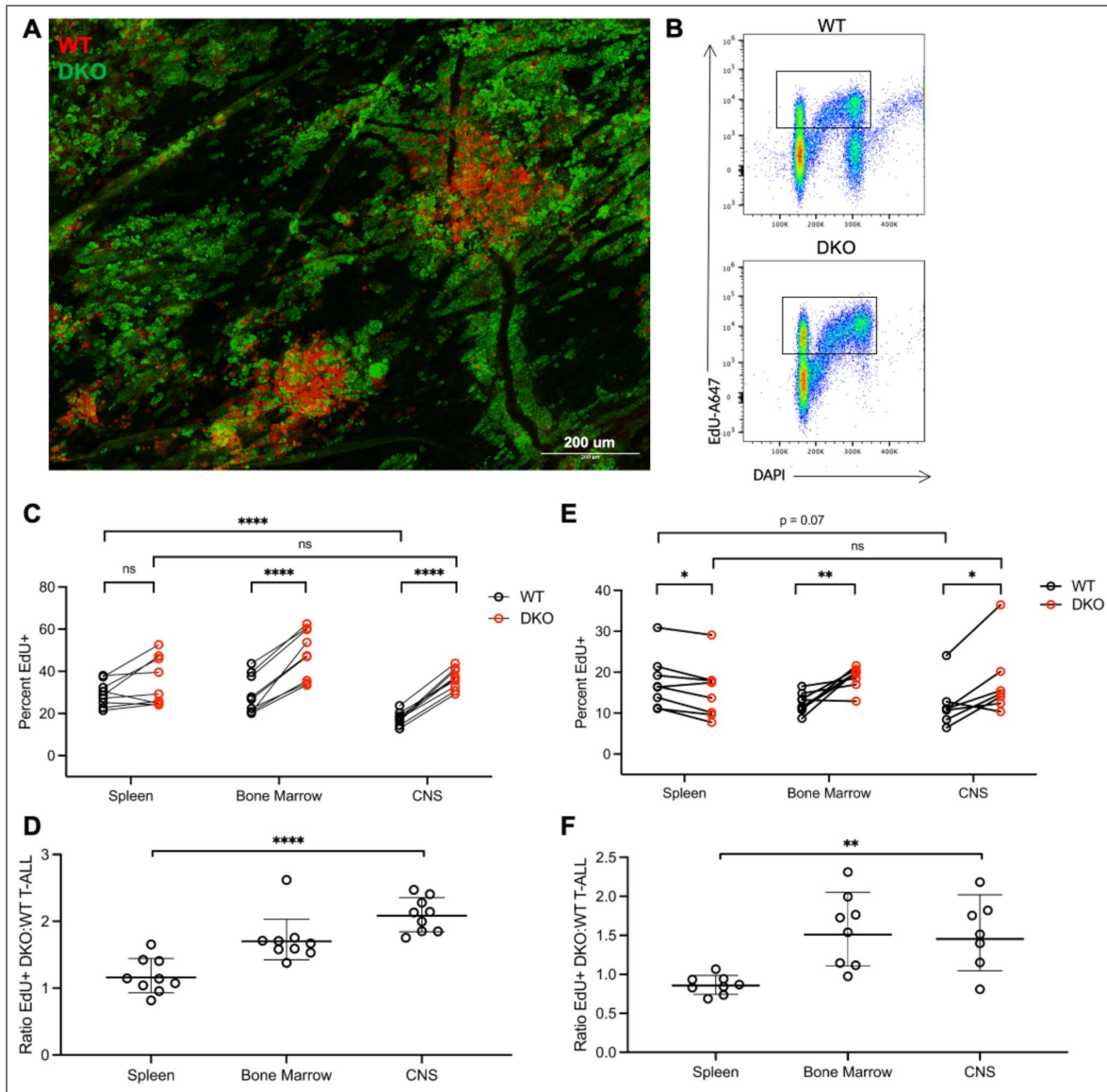
We hypothesized that a difference in distribution might result in differential access to nutrients or differential signals from the stromal niche and in turn affect proliferation. Indeed, interactions with meningeal stroma have been shown to limit ALL proliferation in culture and xenotransplants [41]. We therefore performed an *in vivo* pulse-labelling experiment with 5-ethynyl-2'-deoxyuridine (EdU) and assessed the incorporation of EdU and the abundance of DNA (cell cycle stage) by flow cytometry (Fig. 3B). We observed proliferating DKO and WT T-ALL cells in the CNS, bone marrow, and spleen, and found that DKO T-ALL was more proliferative than WT in both the bone marrow and CNS (Fig. 3C). Furthermore, the greatest difference between DKO and WT T-ALL proliferation was present in the CNS (Fig. 3D). When we repeated this experiment with the CRISPR-DKO and WT T-ALL pair, we again observed increased proliferation of DKO T-ALL in the CNS (Fig. 3E,F). Differences between the DKO and WT T-ALL in other tissues were less consistent between the two pairs of lines. Interestingly, when comparing the proliferation rate of either DKO or WT T-ALL across tissues, there was little difference in the DKO T-ALL proliferation rate between spleen, bone marrow, and CNS; rather, the proliferation of the WT T-ALL was suppressed in the CNS when compared to spleen (Fig. 3C,E).

We hypothesized that the unique stroma of the CNS might be responsible for suppressing the proliferation of WT T-ALL. Fibroblasts are abundant in the meninges and can powerfully modulate immune responses [42–47]. We expanded dural fibroblasts from mice with T-ALL and assessed their transcriptome by RNA-sequencing [43]. We used the NicheNet program to match ligands expressed by the fibroblasts with altered signaling in T-ALL indicated by differentially expressed genes in control versus DKO T-ALL isolated from the CNS [48, 49]. NicheNet highlighted TGF-beta as most likely to be responsible for differences between control and DKO T-ALL, followed by oncostatin M (S. Fig. 4A, B). Future work will test these candidates.

## Integrin blockade synergizes with chemotherapy targeting proliferating cells to limit CNS T-ALL

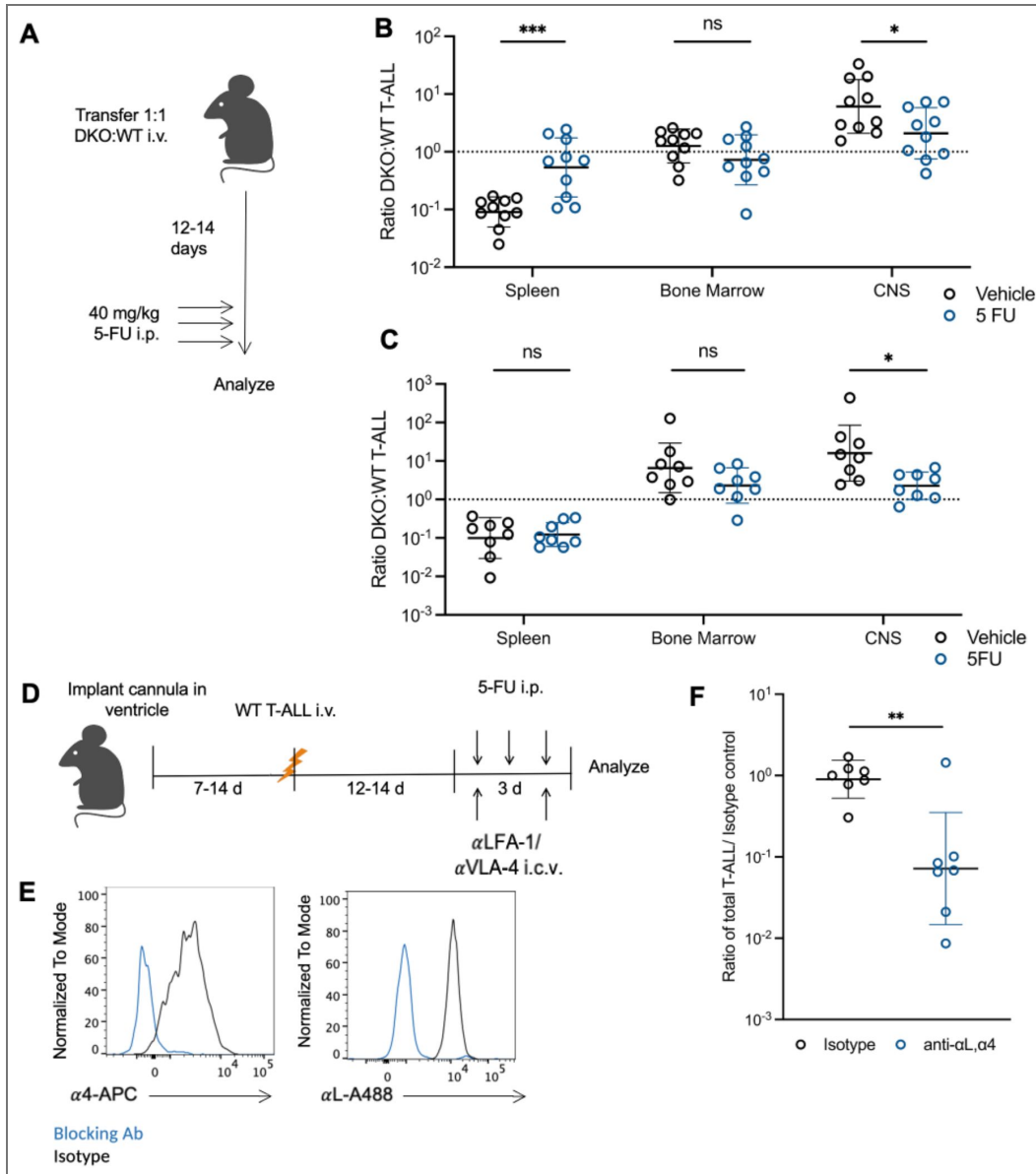
One of the mainstays of therapy for ALL is to target rapidly-proliferating cells with nucleotide-synthesis inhibitors [3]. We hypothesized that if increased proliferation of DKO compared to WT T-ALL cells in the CNS was in fact responsible for the increased accumulation of DKO compared to WT T-ALL in the CNS, the ratio of DKO:WT T-ALL should normalize if proliferation was limited. To test this, we treated mice at mid-stage disease with the thymidylate synthase inhibitor 5-fluorouracil (5FU), which reaches the cerebrospinal fluid, for three days [50–53] (Fig. 4A). We found that DKO T-ALL was depleted more than WT T-ALL in the CNS, where the ratio of DKO:WT T-ALL became nearly 1:1 in just three days (Fig. 4B). When we repeated this experiment with the CRISPR-DKO and WT T-ALL pair, we again observed a reduction in the ratio of DKO:WT T-ALL in the CNS with a short treatment (Fig. 4C). The efficacy of 5FU treatment was confirmed by enumerating total T-ALL in key tissues compared to vehicle-treated mice (S. Fig. 4C,D).

Given these findings, we hypothesized that in wild-type T-ALL disease, the efficacy of chemotherapy in the CNS could be improved by simultaneously blocking integrins; this strategy might encourage T-ALL cells to proliferate and thereby render them sensitive to the drugs. We used an intraventricular cannula to administer integrin-blocking antibodies to the CNS of mice with established WT T-ALL at the same time that they received systemic chemotherapy (5FU) (Fig. 4D). We confirmed the efficacy of blocking antibodies by flow cytometry (Fig. 4E). We found that the combination of intracranial blocking antibodies and systemic chemotherapy was far more effective at reducing total CNS T-ALL than chemotherapy alone (Fig. 4F). Mice that received intracranial antibodies without chemotherapy had no significant difference in CNS T-ALL accumulation from those that received isotype (S. Fig. 4E); however, administration of antibodies without reducing the T-ALL burden with 5FU resulted in a high rate of mortality (3/8 vs 1/16).



**Fig. 3. Integrin-deficient T-ALL proliferates more than WT T-ALL in the CNS.**

(A) Representative (of  $n = 12$ ) immunofluorescence image of whole-mounted dural meninges of mice injected i.v. with 1:1 DKO:WT T-ALL. Mice were analyzed at mid-stage disease. Red, WT T-ALL; green, DKO T-ALL; scale bar, 200 $\mu$ m. (B) Representative flow cytometry gating to identify proliferating cells isolated from tissues of mice that received EdU 4 hours before analysis (in two intraperitoneal doses, 100  $\mu$ g at -4h and 100  $\mu$ g at -2h). (C, D) Sub-lethally irradiated WT mice received 1:1 DKO:WT T-ALL cells i.v., and were analyzed at mid-stage disease. (C) The percent EdU+ of WT (black) or of DKO (red) T-ALL cells and (D) the ratio of % EdU+ DKO: % EdU+ WT T-ALL cells in the indicated tissues. Data compiled from 3 experiments;  $n = 9$ . (E, F) As in (C, D) but for the CRISPR-WT and CRISPR-DKO pair of T-ALL lines. Data compiled from 4 experiments;  $n = 8$ . For (C, E), statistics comparing WT and DKO within a tissue were calculated on logit-transformed data using paired t-test. Statistics comparing WT or DKO from CNS to spleen were calculated on logit-transformed data using unpaired t-test with Welch's correction. For (D, F), statistics calculated using unpaired t-test with Welch's correction. Bars represent geometric mean and geometric SD. \* $p \leq 0.05$ , \*\* $p \leq 0.01$ , \*\*\* $p \leq 0.001$ , \*\*\*\* $p \leq 0.0001$



**Fig. 4. Integrin-deficient T-ALL is more susceptible to chemotherapy targeting dividing cells and integrin blockade synergizes with chemotherapy to limit CNS T-ALL**

(A-C) Sub-lethally irradiated WT mice received 1:1 DKO:WT T-ALL i.v., were treated daily i.p. with 40mg/kg 5-fluorouracil (5-FU) for 3 days at mid-stage disease (starting 12-14 days after T-ALL transfer), and were analyzed 24h after the last dose of 5-FU. (A) Experiment design. (B) The ratio of DKO:WT T-ALL cells in vehicle-treated (black;  $n = 10$ ) or 5FU-treated (blue;  $n = 10$ ) mice. Data compiled from 2 experiments. (C) As in (B) but for the CRISPR-WT and CRISPR-DKO pair of T-ALL lines. Vehicle-treated (black;  $n = 8$ ), 5FU-treated (blue;  $n = 8$ ). Data compiled from 2 experiments. (D-F) Mice were surgically implanted with a cannula in the left lateral ventricle. At least 7 days later, they were sub-lethally irradiated and injected i.v. with WT T-ALL. At mid-stage disease (12-14 days after T-ALL transfer), mice were injected i.p. with 40mg/kg 5-FU daily for three days. On days 1 and 3 of 5-FU treatment, mice received 5 microliter intraventricular infusions of isotype (47 microgram) or anti-LFA-1 and anti-VLA-4 (26 micrograms each) blocking antibodies. 24h after the last dose, tissues were isolated and T-ALL cells were enumerated. (D) Experiment design. (E) Surface expression of  $\alpha 4$  and  $\alpha L$  integrins on WT T-ALL from the CNS after intracerebroventricular injection of LFA-1- and VLA-4- blocking antibodies (blue; representative of  $n = 7$ ) or isotype (black; representative of  $n = 7$ ). Representative of 4 experiments. (F) Total number of WT T-ALL in the CNS normalized to the average isotype control per experiment. Data compiled from 4 experiments;  $n = 7$  pairs. Bars represent geometric mean and geometric SD. Statistics calculated on log-transformed data using unpaired t-test with Welch's correction. \* $p \leq 0.05$ , \*\* $p \leq 0.01$ , \*\*\* $p \leq 0.001$ , \*\*\*\* $p \leq 0.0001$

## Discussion

Much to our surprise, the loss of LFA-1 and VLA-4 in T-ALL did not block leukemia access to the CNS but instead promoted T-ALL accumulation in the CNS. It now seems improbable that T-ALL enters the CNS by crossing an intact endothelial barrier. Perhaps T-ALL either damages the endothelium to enter the CNS or migrates into the CNS via adjacent tissues, such as the bridging channels linking the skull bone marrow to the dura mater, although B-ALL transit through the bridging channels has been shown to require integrins that T-ALL does not express in our model [11]. An important future question is how T-ALL accesses the CNS: what route does it take into the meninges, and what are the mechanical requirements for this migration? Mechanistically, integrin expression suppresses proliferation of T-ALL within the CNS, likely by mediating interactions with the meningeal stroma. A second key question is how integrins limit T-ALL proliferation in the CNS. The CNS is a canonically immune-suppressive environment, and the nature of CNS immune responses is an area of active research [42, 47, 54, 55]. We hypothesize that interactions between the meningeal stroma and normal T cells may limit over-exuberant responses and incidentally restrain T-ALL division.

Finally, these findings have important implications for the design of therapeutic strategies for ALL. We have found strong synergy between integrin blockade and anti-proliferative chemotherapy to treat CNS T-ALL. An analysis of 261 blood and bone marrow biopsies from pediatric and young adult T-ALL patients revealed consistently high integrin expression [13] (S. Fig. 5D [↗](#)). High integrin expression by B-ALL cells independently predicts poor prognosis [35, 56–59], and VLA-4 expression has been found to mediate chemoresistance of ALL in the bone marrow [58, 60–64]. Within the CNS, proteomic and transcriptomic analyses of ALL indicate altered expression of adhesion proteins [65–67]. Furthermore, upon xenotransplantation of Jurkat T-ALL cells into mice, combined treatment with the chemotherapeutic drug cytarabine and Tris[2-(dimethylamino)ethyl]amine, which disrupts adhesion of Jurkat cells to meningeal cells in culture, reduced the number of Jurkat cells in the meninges compared to cytarabine alone [41]. Our findings highlight the need to understand what the fate of leukemia cells is when they are untethered from their niche. This might vary drastically from tissue to tissue, and understanding these cell dynamics will inform the optimal combination of treatments. Thus, for example, integrin loss alone was toxic in our model, but integrin blockade was highly effective in combination with 5FU. Ultimately, targeting integrins in T-ALL may be a powerful strategy to reduce the need for prolonged neurotoxic therapies.

## Materials and methods

### Animals

C57BL/6 (CD45.2), B6.SJL-PtprcaPepcb/BoyJ (CD45.1), Mx1-cre [68], *Itgb1*<sup>fl/fl</sup> [17], *Itgat*<sup>-/-</sup> [16] and NRG (NOD background, *Rag1*<sup>-/-</sup>*Il2rg*<sup>-/-</sup> [69]) mice were obtained from Jackson Laboratory. K14-VEGFR3-Ig mice were kindly provided by Kari Alitalo [70]. Mice were compared to littermate controls, unless otherwise specified. For CD45.2 or CD45.1/2 hosts, female adult mice (6–12 weeks) were used. For all other hosts, male and female adult mice (6–12 weeks) were used depending on availability, as sex did not seem to affect the results. Mice were housed in specific pathogen-free conditions at New York University School of Medicine animal facilities. All cages were on a 12-h light/dark cycle (light 06:00–18:00) in a temperature-controlled and humidity-controlled room. Room temperature was maintained at 72 ± 2 °F (22.2 ± 1.1 °C), and room humidity was maintained at 30–70%. All animal experiments were performed in accordance with protocols approved by the New York University Grossman School of Medicine Institutional Animal Care and Use Committee.

## Bone marrow transduction and transplantation for generation of T-ALL

To generate the original GFP+ DKO and mCherry+ WT T-ALL lines, hematopoietic cells were isolated from bone marrow by flushing femurs and tibias. Stem and progenitor cells were enriched by magnetic selection of cells expressing c-Kit (STEMCELL EasySep Mouse Biotin Positive Selection Kit II). Cells were cultured in the presence of 50 ng/mL Flt3 ligand, 50 ng/mL stem cell factor (SCF), 10 ng/mL interleukin-3 (IL-3), and 10 ng/mL interleukin-6 (IL-6) (PeproTech) in OptiMEM + GlutaMAX (Invitrogen) supplemented with 10% fetal bovine serum (FBS)(Cytiva), 100 IU/mL penicillin and 100 ug/mL streptomycin (Gibco or Corning), and 55  $\mu$ M  $\beta$ -mercaptoethanol (Sigma). 24 and 48 hr after enrichment, c-Kit+ cells were infected with retrovirus (MSCV Notch1 $\Delta$ E-IRES-GFP or Notch1 $\Delta$ E-IRES-mCherry), generated by transfection of Platinum E (Plat-E) retroviral packaging cells (Cell BioLabs Inc.), and concentrated by filtration according to the manufacturer's instructions (Amicon Ultra-15; 100,000 NMWL). Transduction efficiency was determined by reporter fluorescence at 96 hr. Irradiated C57BL/6 mice (two rounds of 500 rads) received  $35 \times 10^3$  Notch1- $\Delta$ E-IRES-GFP/mCherry+, lineage- (lineage markers: CD4, CD8, B220, Gr1, Ter119, Cd11b) cells, together with  $500 \times 10^3$  unfractionated bone marrow cells for hemogenic support, intravenously by retro-orbital injection. Mice were bled regularly to monitor leukemia progression. The DKO donor hematopoietic cells were *Itgal*<sup>-/-</sup> *Itgb1*<sup>fl/fl</sup> *Mx1-Cre*. To delete *Itgb1*, when GFP+ blasts reached >10% of peripheral blood, mice received three intraperitoneal injections of poly(I:C) (10 ug/g, GE Healthcare) in PBS, administered every other day. Notch1 $\Delta$ E-IRES-GFP *Itgb1*<sup>fl/fl</sup> *Itgal*<sup>-/-</sup> *Mx1-cre* cells (DKO T-ALL) or Notch1 $\Delta$ E-IRES-mCherry cells (WT T-ALL) were isolated from bone marrow at a humane endpoint.

For secondary model induction T-ALL cells were maintained *in vitro* (see below). For induction, recipients were sub-lethally irradiated (550 cGy) and retro-orbitally injected with  $1-2 \times 10^6$  T-ALL cells. Radiation was performed with either Cs or X-ray irradiator (X-rad 320/350).

### T-ALL cell culture

Primary mouse T-ALL cells isolated from bone marrow were maintained on OP9 stromal cells [71], kindly provided by Iannis Aifantis, in OptiMEM + GlutaMAX (Invitrogen) supplemented with 10% fetal bovine serum (FBS), 5 ng/ml IL-7, 100 IU/mL penicillin and 100 ug/mL streptomycin (Gibco or Corning), and 55  $\mu$ M  $\beta$ -mercaptoethanol. Cells were passaged every 3–4 days onto a fresh feeder layer. OP9 cells were cultured in 75cm<sup>2</sup> flasks in MEM $\alpha$  + GlutaMAX supplemented with 20% FBS and 100 IU/mL penicillin and 100 ug/mL streptomycin (Gibco or Corning), and passaged every 2-3 days. For co-culture, OP9s were pre-treated with 7.5 ng/mL mitomycin C (Sigma) for 3-4 hours to prevent feeder cell division. T-ALL cells for leukemia induction were isolated from co-culture by density gradient (Ficoll-paque Premium, Cytiva).

### Retroviral transduction of T-ALL cells

The coding regions of *Itgb1* and *Itgal* (Dharmacon) were cloned into the pMXs-IRES-mCherry retroviral vector, and Plat-E packaging cells were used for the generation of retrovirus. DKO T-ALL cells, seeded on mitomycin C-treated OP9s, were transduced with a retrovirus encoding either *Itgb1* or *Itgal* or empty vector. 48-72 hours after transduction, cells were sorted on a BD FACSAria II for positive surface expression of  $\alpha$ L or  $\beta$ 1. Cells were allowed to recover on MMC-treated OP9s for 48 hours and another round of transduction was performed using the plasmid carrying the other gene. After 48-72 hours, T-ALL cells double-positive for surface expression of  $\alpha$ L and  $\beta$ 1 were sorted on the BD FACSAria II.

### CRISPR deletion of genes in T-ALL cells

gRNAs targeting *Itgal*, *Itgb1* and *Itgb7* were ordered from IDT (sequences below) for use with the IDT Alt-R CRISPR-Cas9 system. ALT-R CRISPR-Cas9 Negative Control crRNA#1 was ordered as a non-targeting control (IDT). On day of nucleofection, gRNAs were resuspended in sterile tris-EDTA pH 8.0 to a concentration of 100uM. 1.2 uL of each gRNA for either *Itgal* or *Itgb1* or *Itgb7* were mixed

with 1.7 uL of Alt-R S.p. HiFi Cas9 Nuclease V3, and 0.9 uL of PBS (total volume of 5 uL), and incubated for 20 minutes at room temperature, per manufacturer's instructions.  $1-2 \times 10^6$  T-ALL cells were resuspended in 100uL P3 nucleofector solution (82 uL nucleofection solution + 18 uL supplement; Lonza P3 Primary Cell 4D-Nucleofector™ X Kit), combined with gRNA-Cas9 complex, and nucleofected using program "activated T cells" using the Lonza Amaxa 4D Nucleofector. Immediately after, 500 uL warm media was added to cells in cuvette and incubated for 10 min at 37C before cells were returned to culture. After 48-72 hours, T-ALL cells were sorted based on surface expression of  $\alpha$ L or  $\beta$ 1 on BD FACSAria II. T-ALL cells were then subjected to a second round to delete both *Itgal* and *Itgb1* and in some cases a third round to delete *Itgb7*.

*Itgb1:*

Mm.Cas9.ITBG1.1.AB: 5' ACA UUC UCC GCA AGA UUG UUU UAG AGC UAG AAA UAG CAA GUU  
AAA AUA AGG CUA GUC CGU UAU CAA CUU GAA AAA GUG GCA CCG AGU  
CGG UGC 3'

Mm.Cas9.ITBG1.1.AC: 5' GUC ACC AAU CGC AGC AAG UUU UAG AGC UAG AAA UAG CAA GUU  
AAA AUA AGG CUA GUC CGU UAU CAA CUU GAA AAA GUG GCA CCG AGU  
CGG UGC 3'

*Itgal:*

Mm.Cas9.ITGAL.1.AC: 5' GUA CUU GGG AAU GAC GCG UUU UAG AGC UAG AAA UAG CAA GUU  
AAA AUA AGG CUA GUC CGU UAU CAA CUU GAA AAA GUG GCA CCG AGU  
CGG UGC 3'

Mm.Cas9.ITGAL.1.AE: 5' GCC AAG AAG CUC UGC GUG UUU UAG AGC UAG AAA UAG CAA GUU  
AAA AUA AGG CUA GUC CGU UAU CAA CUU GAA AAA GUG GCA CCG AGU  
CGG UGC 3'

*Itgb7:*

Mm.Cas9.ITGB7.1.AA: 5' CAG CAC CGU UUU GUC CAC GAG UUU UAG AGC UAG AAA UAG CAA  
GUU AAA AUA AGG CUA GUC CGU UAU CAA CUU GAA AAA GUG GCA CCG  
AGU CGG UGC 3'

Mm.Cas9.ITGB7.1.AC: 5' CGU GGA CCU AUA UUA CCU UAG UUU UAG AGC UAG AAA UAG CAA  
GUU AAA AUA AGG CUA GUC CGU UAU CAA CUU GAA AAA GUG GCA CCG  
AGU CGG UGC 3'

## Tissue isolation and processing for flow cytometry

All mice were euthanized and perfused with 5-10mL PBS. Unless otherwise specified, after isolation, all tissues and cell suspensions were stored in PBS/2% FBS on ice.

*Blood:* collected via cardiac puncture and stored in blood collection tube with K<sub>2</sub>EDTA (BD Microtainer).

*Bone marrow:* right tibia and femur were flushed with PBS/2% FBS.

*Skull bone marrow:* isolated from calvarium by mechanical disruption with glass mortar and pestle. *Spleen and lymph nodes:* mechanical disruption and filtration through a 100um cell strainer with the flat end of a 1mL syringe plunger.

*Central nervous system (CNS):* brain and inside of the dura attached to skull were washed with PBS/2% FBS via p1000 pipette and cells were collected. The wash and whole brain were placed in a 50mL conical tube and pulse-vortexed on low speed for 15 seconds. Brain was collected and

mechanically disrupted and filtered through a 100µm cell strainer with the flat end of a 1mL syringe plunger. Lymphocytes were purified by centrifugation using a 33% isotonic density gradient (Percoll; GE Healthcare). Red blood cells were depleted using tris-ammonium chloride solution. For later experiments (Fig. 1K,L, 2D-4), we determined that brain parenchyma processing was dispensable: >95% of T-ALL cells were collected with just wash of inside of dura and vortex of whole brain, and ratio was unaffected. All mice in each experiment were processed using the same protocol.

*Cell counting*, single-cell suspensions were counted using a Multisizer 3 (Beckman Coulter) set to detect nuclei between 3.5-10µm. Total cell counts from bone marrow were obtained by assuming that cells isolated from tibia and femur of one leg represent 10% of the total bone marrow per mouse (unless otherwise noted). Total cell counts from blood were obtained by assuming that the total amount of blood per mouse is 2mL.

## Immunostaining and flow cytometry

*Immunostaining*: Unless otherwise specified, surface stains were performed on live cells with fluorophore-conjugated antibodies at 1:200 dilution in PBS supplemented with 2% FBS, 2% Normal Mouse Serum (NMS), and 2% Normal Rat Serum (NRS), for 20 minutes on ice and protected from light.

*Antibodies purchased from Biolegend*: anti-B220 (clone RA3-6B2), anti-CD4 (clone GK1.1), anti-CD8 (clone 53-6.7), anti-CD11a (clone M17/4), anti-CD11b (clone M1/70), anti-CD25 (clone PC61), anti-CD31 (clone MEC13.3), anti-CD44 (clone IM7), anti-CD45 (clone 30-F11), anti-CD45.1 (clone A20), anti-CD45.2 (clone 104), anti-CD49d (clone R1-2), anti-CD104 (clone 346-11A), anti-β7 (clone FIB504), anti-CX3CR1 (SA011F11), anti-GR1 (clone RB6-8C5), anti-Ter119 (clone TER-119), Streptavidin, mouse IgG2a isotype control (clone MOPC-173) and rat IgG2a, k isotype control (clone RTK2758).

*Antibodies purchased from eBioscience*: anti-CD4 (clone GK1.5), anti-CD29 (clone eBioHMb1-1), anti-Lyve1 (clone ALY7), and Armenian Hamster IgG isotype control (clone eBio299Arm).

*Apoptosis stains*: anti-Active Caspase-3 (clone C92-605) was purchased from BD Biosciences and cells were fixed and permeabilized according to kit instructions (BD Biosciences Cytotfix/Cytoperm Cat: BDB554714). Cells were stained at a concentration of 1:100 for 20 minutes at 4°C.

*Viability stains*: “LIVE/DEAD™ Fixable Far Red Dead Cell Stain Kit, for 633 or 635 nm excitation” was purchased and used according to kit instructions (ThermoFisher).

*EdU labelling*: “Click-iT™ Plus EdU Alexa Fluor™ 488 Flow Cytometry Assay Kit” and “Click-iT™ Plus EdU Alexa Fluor™ 647 Flow Cytometry Assay Kit” were purchased and cells were stained according to kit instructions (ThermoFisher). DNA quantification was performed by staining with DAPI (Sigma) at 2µg/mL with 5µg/ml RNase A (Thermo Scientific) for 15 minutes at RT.

*Flow cytometry*: data was acquired using a BD LSRII or Beckman Coulter Cytoflex and analyzed using FlowJo v10.

## Mouse treatments

*For partial irradiation/shielding of CNS*: 1/16” sheet lead (McMaster-Carr) was formed into a half cylinder (6” long, 2” radius) with two semi-circular cut outs in the middle of the long sides (approx. 1-1.5” deep). Four of these half-cylinders were nested to achieve a total thickness of ¼” inch lead and secured with tape. Mice were anesthetized with ketamine/xylazine (100mg/kg /10 mg/kg) and placed on a rigid, irradiator-safe surface with their limbs taped out perpendicular to their body. The lead shielding was placed over the mice, ensuring that their taped limbs were exposed by the semi-circular cut outs, and that all other parts of their body, including the tail, were underneath the lead. Mice were irradiated with 550 cGy using an X-ray irradiator with directional collimator and SnAlCu filter (X-rad320, Precision X-Ray).

*For ablation of meningeal lymphatics*, 5–8-week-old female C57BL/6 mice were injected intraperitoneally with  $1 \times 10^{12}$  PFU of either AAV-VEGFR3(1-4)-Ig (active) or AAV-VEGFR3(4-7)-Ig (control) (provided by Kari Alitalo).

*For intraventricular infusion*, mice were surgically implanted with an intraventricular cannula (P1 Technologies, C315GS-4/SP GUIDE 26GA 38834 4MM Pedestal, 2MM Projection) in the left lateral ventricle (-0.5mm R/C, -1.0mm M/L, -1.8mm D/V relative to bregma). For infusions, >5uL was drawn up into a 33G air-tight Hamilton Syringe (Hamilton, 1705 series) and syringe was mounted in a Remote Infuse/Withdraw Pump 11 Elite Nanomite Programmable Syringe Pump (Harvard Apparatus) mounted on a stereotaxic instrument (Model 940, Kopf). Mice were anesthetized with isoflurane (1.5%-2.5%) or ketamine/xylazine (100mg/kg / 10mg/kg) and mounted via bite bar/nose cone. Syringe needle was advanced into cannula 8.5mm from opening (approx. 0.3-0.5mm into lateral ventricle) and 5uL infusion was conducted at 250nL/min with a 10min rest before needle was withdrawn. *For dye infusion*, NHS-ester-Atto647 (Sigma) was dissolved in DMSO at 10mg/mL (stored at -20C), then further diluted 1:5 in PBS to 2mg/mL for infusions.

*For myeloid cell depletion*, CSF1R inhibitor PLX5622 was formulated at 1200ppm into rodent diet AIN-76A by Research Diets, Inc. (New Brunswick, NJ). Animals received either the PLX5622 formulated diet or a normal chow diet (AIN-76A) a week prior to leukemia induction and for the duration of the experiment (total four weeks). Both PLX5622 and control diet were irradiated and contained red dye.

*For EdU labelling experiments*, mice were administered 100ug EdU (ThermoFisher) in 100uL PBS intraperitoneally at 4h and 2h before euthanasia for a total dose of 200ug EdU.

*For 5-fluorouracil experiments*, 5-fluorouracil (5FU) (Sigma) was dissolved in DMSO to a concentration of 50mg/mL and stored at -20C. For injections, 5FU was diluted 1:10 in PBS to 5mg/mL and mice were injected intraperitoneally at a dose of 40mg/kg. Vehicle-treated mice were injected with 150uL of 10% DMSO/PBS.

*For blocking antibody experiments*, intracerebral ventricular infusions were performed as described above. Anti-CD49d (PS/2; Bio-Xcell) and anti-CD11a (M17/4; Bio-Xcell) were combined at equal quantities for a total of 26ug of each in 5uL. Isotype control mice received 5uL (47ug) of Rat IgG2a, anti-trinitrophenol (clone 2A3; Bio-Xcell). Infusions were performed three days and one day before tissue isolation and analysis.

## Immunofluorescence

*Meninges whole mounts*, adapted from [72]. The top of the skull was removed with surgical scissors. While still attached to the skull, whole-mount meninges were fixed in 2% PFA overnight at 4C. The dura/arachnoid was then peeled from the skullcap. Meninges were incubated in blocking solution [PBS containing 2% normal serum (goat and/or donkey), 3% bovine serum albumin (BSA) (Fisher), anti-CD16/32 (clone 93, eBioscience, 1:1,000), 0.1% Triton X-100, and 0.05% Tween 20] supplemented with avidin block (Vector Laboratories) for one hour at RT on a shaker; washed (with PBS 3 times for 5 min each at RT); followed by another one-hour block at RT with blocking buffer supplemented with biotin block (Vector Laboratories), and washed again. Primary antibodies were applied at 1:200 dilution in staining buffer [PBS containing 3% BSA and 0.5% Triton X-100] and meninges were incubated overnight at 4C. Meninges were washed then incubated in appropriate dilutions of secondary antibodies in staining buffer for 2 hours at RT or overnight at 4C. Meninges were washed, then wet mounted on a Superfrost Plus glass slide, and allowed to dry for 10-15 minutes. Fluoromount G and a coverslip were added and slides were allowed to fully dry at RT (16h) before edges were sealed with clear nail polish.

*Primary antibodies*: anti-Lyve-1 biotinylated (clone ALY7; eBioscience), anti-CD31-PE (clone MEC13.3; Biolegend), anti-GFP (ab13970; Abcam), anti-RFP (600-401-379, Rockland).

*Secondary antibodies*, Streptavidin-BV421 1:500 (Biolegend), Alexa Fluor 488 AffiniPure Goat Anti-Chicken IgY (IgG) (H+L) 1:1000 (Jackson ImmunoResearch), Alexa Fluor 647 AffiniPure Donkey Anti-Rabbit IgG (H+L) 1:1000 (Jackson ImmunoResearch), Cy3 Affinipure Donkey anti-Rabbit IgG (H+L) 1:1000 (Jackson ImmunoResearch).

Images were acquired using confocal microscope (LSM710 (Carl Zeiss) or Stellaris 8 (Leica)) with a 10x dry or 20x dry objective using ZEN (Zeiss) or LasX (Leica) software. Images were processed and analyzed using ImageJ (version 1.53; National Institutes of Health) and FIJI (version 2.14.0,

Schindelin et al, 2012).

## RNA extraction and sequencing

*For T-ALL cell sequencing:* DKO, WT, CRISPR-DKO and CRISPR-WT (both CD45.2) cells were isolated from the CNS and spleen of 1:1 DKO:WT co-transfer mice (congenically-marked, CD45.1/2) at mid-stage disease, stained for the presence of the integrins and CD45.1 and CD45.2, and sorted using a BD FACS Aria.

*For fibroblast sequencing:* Dural meninges were isolated from naïve wild-type mice as described in [26] and placed immediately in microcentrifuge tubes of digestion buffer containing collagenase D (1mg/ml, MilliporeSigma), DNase I (50ug/mL, Roche), and Dispase (5 U/mL, Roche) in serum-free, high-glucose DMEM (Invitrogen) (modified from [43]) and stored on ice. Tissues in microtubes were placed on rotator for 2h at 37C. Suspension was agitated by pipetting until fully dissociated and filtered through 40um cell strainer. Single cell suspension was pelleted at 400g for 4 minutes and seeded on 6-well tissue culture treated plates in meningeal fibroblast media (high-glucose DMEM supplemented with 15% FBS (Cytiva) and 100 IU/mL penicillin and 100 ug/mL streptomycin (Gibco or Corning)). 16h after seeding, media and non-attached cell fraction was aspirated, leaving fibroblasts behind, and media was replaced and changed every 2-3 days. Meningeal fibroblasts were cultured for 10-14 days before being lifted from the plate with Trypsin/EDTA and snap-frozen in Trizol for RNA extraction.

Total RNA was extracted using RNeasy Plus Micro Kit (Qiagen) and purified with RNase-Free DNase (Qiagen). Total RNA was quantified on an Agilent BioAnalyzer via the Pico kit (Agilent). SMART-Seq HT Kit (Takara) was used to construct cDNA and libraries. cDNA was constructed using 1ng input of RNA and amplified with 12 cycles. cDNA product was purified with DNA AMPure beads (Beckman-Coulter) and quantified via Agilent Quant-IT. 300pg of each cDNA sample was used to construct libraries via Nextera XT DNA library preparation kit. 12 PCR cycles were used to amplify the libraries and to add the barcodes. Libraries were then cleaned, quantified, and pooled for sequencing on an S1 (100 cycles) flow cell on Illumina NovaSeq sequencer.

Sequencing reads were mapped to the reference genome (mm10) using the STAR aligner (v2.5.0c) [73]. Alignments were guided by a Gene Transfer Format (GTF) file. The mean read insert sizes and their standard deviations were calculated using Picard tools (v.1.126) (<http://broadinstitute.github.io/picard>). The read count tables were generated using HTSeq (v0.6.0) [74], normalized based on their library size factors using DESeq2 [75], and differential expression analysis was performed. The Read Per Million (RPM) normalized BigWig files were generated using BEDTools (v2.17.0) [76] and bedGraphToBigWig tool (v4). All the downstream statistical analyses and generating plots were performed in R environment (v3.1.1) (<https://www.r-project.org/>).

Nichenet analysis was performed according to [48, 49] and using the code graciously provided on [https://github.com/saeyslab/nichenetr/blob/master/vignettes/ligand\\_activity\\_geneset.md](https://github.com/saeyslab/nichenetr/blob/master/vignettes/ligand_activity_geneset.md). The list of differentially expressed genes was identified by FDR <0.1 from the DESeq2 analysis performed on the WT and DKO T-ALL isolated from the CNS (described above). The mouse reference libraries were used for analysis.

Packages used to generate plots were TidyVerse [77], ggplot, and circlize [78].

## Statistical analysis

Statistical analysis (excluding RNA-seq experiments) was conducted using the PRISM program (GraphPad). Experiments comparing the total number of WT to the total number of DKO T-ALL cells, and experiments comparing the ratio of DKO:WT between groups, were analyzed using a two-tailed, unpaired t-test with Welch's correction. Experiments comparing the survival or proliferation of WT to DKO T-ALL cells were analyzed using a two-tailed, paired t-test. For data displayed on a log-scale,  $\log_{10}$ -transformed data was used to calculate significance. For data presented as percentages, logit-transformed data was used to calculate significance. Significance was defined as \*p < 0.05, \*\*p < 0.01, \*\*\*p < 0.001, \*\*\*\*p < 0.0001.

## Data availability

RNA sequencing expression data included in this study have been deposited in the NCBI Gene Expression Omnibus (GEO), accession number GSE318579. Additional data not in the article itself are available upon reasonable request from the corresponding author.

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### Author contributions

Conceptualization: SYL, CC, SRS. Methodology: MO, KAM, AYC, JKS, RCF, MC. Investigation: SYL, CC, HC, BSS, JBV. Formal analysis: SYL, CC. Writing, original draft: SYL, SRS. Writing, review & editing: SYL, CC, HC, BSS, SRS. Supervision: SRS.

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## Additional files

[Supplementary material](#) 

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## Peer reviews

### Reviewer #1 (Public review):

The manuscript by Lux et al. addresses how T-cell acute lymphoblastic leukemia (T-ALL) cells migrate into the central nervous system (leptomeninges), specifically through VLA-4 and LFA-1 integrins. VLA-4 and LFA-1 are important regulators of normal T-cell migration into the CNS, so the authors tested whether they also mediate T-ALL infiltration. They generated an intracellular NOTCH1 T-ALL mouse model and then used CRISPR/Cas9 gene targeting to delete VLA-4 and LFA-1. They show that integrin-deficient T-ALL cells accumulate in the CNS compared to control T-ALL cells. The authors performed a time course experiment and found that although WT T-ALL cells accumulated in the CNS before DKO T-ALL cells, over time, DKO T-ALL cells outgrew the WT T-ALL cells. Subsequently, they performed bulk RNA-sequencing and revealed that Integrin beta 7 (Itgb7) was upregulated in the DKO T-ALL cells. To test whether Itgb7 was compensating for the loss of VLA-4 and LFA-1, the authors generated a triple KO (TKO). The TKO T-ALL cells migrated to the CNS; however, CNS accumulation between the TKO and the DKO was not significantly different. To evaluate if there is reduced

exit of T-ALL DKO cells from the meninges, they inhibited T-ALL exit via the dorsal meningeal lymphatics by generating an AAV VEGF-trap encoding the binding domain of VEGFR3, and then co-injected WT: DKO cells weeks later. There was no effect on the WT:DKO T-ALL ratio or on the overall number of T-ALL cells in the CNS with meningeal lymphatics regression, suggesting that the DKO does not preferentially accumulate in the CNS, or that delayed exit results in DKO T-ALL accumulation in the CNS.

Additionally, the authors tested whether DKO affected immune surveillance by injecting DKO:WT T-ALL cells into NRG mice. DKO T-ALL cells localized in the dura mater and were spread throughout the tissue, whereas WT T-ALL cells clustered near blood vessels. These observations lead the authors to hypothesize that differential access to nutrients or other signals may influence leukemic cell proliferation. However, EdU labeling revealed no differences, leading the authors to hypothesize that the unique stromal cell layer in the meninges supports the DKO proliferative advantage. Finally, the authors tested whether integrin blockade and chemotherapy might chemosensitize T-ALL cells in the CNS. After a single treatment with 5FU, DKO cells were depleted faster than the WT cells; however, a single treatment with integrin blockade was toxic. After combining 5FU with the integrin antibodies, the authors showed that T-ALL cells in the CNS were significantly more depleted than in treatment with either single therapy.

These data highlight how challenging it is to identify regulators of T-ALL migration and adherence. This study highlights the importance of these experiments and the clinical need to identify the molecules that influence leukemic infiltration into the CNS.

Overall, this study was well performed with appropriate statistical power to implicate integrins in T-ALL CNS infiltration and proliferation.

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## Reviewer #2 (Public review):

### Summary:

In this study, the authors set out to understand how T cell leukemia cells enter and persist in the CNS, with a particular focus on the role of adhesion molecules known to regulate normal immune cell trafficking. Contrary to expectations, they find that loss of two key adhesion molecules does not impair CNS entry but instead leads to increased accumulation of leukemia cells, which is associated with enhanced cell proliferation in this environment. These findings challenge prevailing assumptions about how leukemia cells interact with tissue niches and suggest a potential therapeutic strategy combining adhesion blockade with chemotherapy.

### Strengths:

The study addresses an important and longstanding question in leukemia biology using well-designed *in vivo* models and multiple complementary approaches. The key observation is robust and consistently supported across genetic models and experimental systems. The authors systematically test alternative explanations, including altered entry, exit, and immune evasion, which strengthens the interpretation that proliferation differences underlie the phenotype. The work has potential translational relevance, particularly in highlighting a possible strategy to enhance the efficacy of anti-proliferative therapies in the CNS.

### Weaknesses:

While the central phenotype is clear, the mechanistic basis remains incompletely defined. Addressing the following points would strengthen the manuscript.

### Major critiques:

(1) The central claim that integrin loss enhances CNS accumulation via increased proliferation is not mechanistically resolved; current data are correlative (EdU incorporation, distribution patterns) and do not establish that integrin-mediated signaling directly restrains cell cycle progression in the CNS niche. The authors should perform functional perturbation of candidate pathways identified (e.g., TGF- $\beta$ ) using pharmacologic inhibitors or genetic approaches (dominant-negative receptor or CRISPR knockdown) in vivo or in ex vivo CNS-derived T-ALL co-culture systems to test whether blocking this pathway rescues the WT proliferation phenotype; if not feasible, the mechanistic claims should be toned down and clearly presented as hypotheses.

(2) The relationship between altered spatial distribution and proliferation is suggestive but not directly demonstrated. The imaging data indicate differences in localization, but these observations are not quantitatively linked to cell cycle status. The authors could strengthen this point by incorporating spatially resolved proliferation analyses, such as combining EdU labeling with imaging or quantifying proximity to stromal or vascular niches, or alternatively by providing additional quantitative analysis of the existing imaging data.

(3) The conclusion that CNS accumulation is not due to altered trafficking (entry/exit) is suggestive but not definitive, as early seeding dynamics are not directly assessed. Authors should perform short-term homing or early time-point competitive trafficking assays (e.g., CNS quantification at 6-48h post-transfer) to rigorously exclude differences in entry kinetics; if such experiments are not feasible, this limitation should be explicitly acknowledged in the discussion.

(4) The therapeutic claim that integrin blockade synergizes with chemotherapy is promising but underdeveloped, as it lacks survival outcomes and a broader translational context. The authors should include survival analyses and, if possible, test combination treatment in a more clinically relevant setting (e.g., delayed intervention or alternative standard-of-care agents), or otherwise temper translational conclusions and discuss risks such as inducing proliferation in the absence of chemotherapy.

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