Emerging single cell endothelial heterogeneity supports sprouting tumour angiogenesis and growth

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37 SUMMARY

38 Blood vessels supplying tumors are often dysfunctional and generally heterogeneous. The 39 mechanisms underlying this heterogeneity remain poorly understood. Here, using 40 multicolor lineage tracing, *in vivo* time-lapse imaging and single cell RNA sequencing in 41 a mouse glioma model, we identify tumour-specific blood endothelial cells that originate 42 from cells expressing the receptor for colony stimulating factor 1, Csflr, a cytokine which 43 controls macrophage biology. These *Csf1r* lineage endothelial cells (CLECs) form up to 44 10% of the tumour vasculature and express, besides classical blood endothelial cell 45 markers, a gene signature that is distinct from brain endothelium but shares similarities 46 with lymphatic endothelial cell populations. in silico analysis of pan-cancer single cell 47 RNAseq datasets highlights the presence of a comparable subpopulation in the 48 endothelium of a wide spectrum of human tumours. We show that CLECs actively 49 contribute to sprouting and remodeling of tumour blood vessels and that selective 50 depletion of CLECs reduces vascular branching and tumour growth. Our findings indicate

51 that a non-tumour resident Csf1r-positive population is recruited to tumours, 52 differentiates into blood endothelial cells to contribute to vascularization and, thereby, 53 tumour growth.

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56 INTRODUCTION

57 Endothelial cells form a single cell layer lining the inner walls of blood vessels and play 58 critical roles in organ homeostasis and disease progression. Once formed, following 59 embryonic and early post-natal development, blood vessels retain a high level of 60 adaptability to meet changing metabolic or hemodynamic requirements or enable further 61 tissue growth including in tumours. How local vascular networks respond to these 62 adaptive challenges, rapidly expand, remodel and reestablish homeostasis remains a 63 timely research topic, with many open questions still unresolved. In particular in 64 pathologies, knowledge of the precise nature and origin of new vessel formation, 65 endothelial activation and differentiation steps, and of the mechanisms driving vessel 66 dysmorphia, or preventing effective revascularization in ischemic diabetic complication 67 is critical for new therapeutic approaches. New blood vessel formation in the adult was 68 long believed to arise exclusively by sprouting and proliferation of endothelial cells from 69 local blood vessels without any de novo differentiation from progenitor cells. However, 70 Asahara and coworkers isolated mononuclear cells from human peripheral blood and 71 identified circulating endothelial progenitor cells (EPCs)¹. These cells were shown to 72 derive from the bone marrow and contribute to endothelial cells in blood vessels in 73 hindlimb ischemia and tumour xenograft mouse model ^{2,3}. By fluorescent in situ 74 hybridization with sex chromosome-specific probes in patients with cancer after bone 75 marrow transplantation, Peters and coworkers demonstrated that bone marrow-derived

76 stem cells contribute to endothelial cells in human tumour endothelium⁴. The resulting 77 concept of vasculogenesis in the adult raised the prospect of novel therapeutic approaches for ischemic vascular disease and for targeting tumour angiogenesis ^{5,6}. Interestingly, 78 EPCs also contributed to newly forming lymphatic vessels ^{7,8}. A vast number of 79 publications since have supported or challenged these initial discoveries, and the 80 81 occurrence, origin and significance of EPCs continues to be a matter of debate 9-12. 82 Whereas EPCs and other circulating progenitor cells were long thought to origin from the 83 bone marrow, recent studies challenged this idea and suggested the existence of a 84 different yet unknown vascular niche for endothelial progenitor cells that can contribute 85 to blood vessel formation¹³.

86

87 **RESULTS**

88 A Csf1r cell lineage contributes to blood vascular endothelial cells during glioma 89 growth in mice.

90 In order to study dynamic macrophage recruitment and blood vessel patterning in vivo 91 during glioma growth, we adapted a surgical cranial window model for two-photon 92 microscopy allowing repeated visualization of the same animal over time. We generated spheroids of syngeneic C57BL/6 mouse glioma cells (CT2A or GL261)^{14,15} modified to 93 94 stably express blue fluorescent protein (BFP), and injected them into the mouse cortex, followed by implantation of a glass coverslip ¹⁶. By crossing transgenic Csflr (Colonv 95 stimulating factor 1 receptor)- specific tamoxifen-inducible Cre driver mice ¹⁷ with the 96 97 mTmG Cre reporter mouse line ¹⁸, we induced membrane-targeted GFP expression in 98 Csflr-expressing cells and used this stable lineage trace to follow the cell population over 99 time (Fig. 1a, b). As expected, the vast majority of tumour associated macrophages were 100 labelled by GFP in line with their *Csf1r*-dependent recruitment and expansion¹⁹.

101 Unexpectedly, however, through the combination of longitudinal live two-photon 102 imaging and lineage tracing, we discovered GFP-positive cells contributing to the tumour 103 vasculature during glioma growth (Fig. 1c). These Csflr lineage endothelial cells 104 (CLECs) frequently emerged at tip cell positions of the growing vasculature, heading 105 blood vessel sprouts in the mouse glioma (Fig. 1d, Supplementary Movie1). Additionally, 106 CLECs were found bridging adjacent capillaries during anastomosis (Fig. 1e). 107 Fluorescent dextran injection confirmed that CLECs contributed to - and were 108 incorporated in the endothelial lining of - functionally perfused blood vessels including 109 lumenized sprouts headed by endothelial tip cells (Fig. 1f). Counterstaining on fixed 110 samples identified that all cells forming blood vessels including the lineage-traced CLECs 111 expressed typical endothelial markers (CDH5, CD31) and lacked macrophage and 112 myeloid cell markers (F4/80, CD45) (Fig. 1g, h, Extended Data Fig. 1). BFP was not 113 expressed in CLECs, demonstrating that they were not derived from glioma cells (Fig. 114 1g).

115 To gain more insight into the nature and significance of CLECs, we isolated and 116 quantified the cell population in the mouse glioma model. By 4 weeks of tumour 117 development (close to the specified humane endpoint of 5 weeks) up to 10% of the tumour 118 vasculature was comprised of cells carrying the *Csfr1* lineage trace. Similar frequencies 119 were found in tumours derived from either of the mouse glioma cell lines, CT2A (Fig. 2a, 120 b), or GL261 (Fig. 2c). Polyclonal antibody staining indicated that Csflr may also be expressed in some brain capillary endothelial cells ^{20,21}. Recent single cell sequencing of 121 122 adult mouse brain endothelial cells however did not confirm this notion²², suggesting that 123 CLECs are unlikely to originate from brain endothelium. Indeed, tamoxifen mediated 124 activation of our transgenic *Csfr1* lineage trace did not label brain endothelial cells in 125 mice lacking tumours, sham operated as well as in the healthy brain parenchyma

126 surrounding implanted tumours (Fig. 2d, e, f). Csflr is important for the embryonic 127 development of microglia and its expression is also detected in adult microglia by single 128 RNA sequencing. Surprisingly however, Csf1r-Mer-iCre-Mer mediated cell 129 recombination of the Cre-Reporter expression was also not observed in microglia of the 130 healthy adult mouse brain. Irrespective of this apparent discrepancy between reported 131 microglial Csflr expression and the lack of Csflr- Mer-iCre-Mer mediated recombination 132 in microglia, these data also exclude microglia as a potential source of CLECs. 133 Furthermore, we observed CLECs also in tumours originating from B16F1 melanoma 134 cells implanted under the skin; thus, this Csfr1-lineage endothelial cell contribution not 135 only occurs in brain tumours with microglia in their environment, but also in other tumour 136 types and locations (Fig. 2g, h). Notably, we also found Csflr-lineage lymphatic 137 endothelial cells in B16F1 skin melanomas indicating that CLECs can contribute to both 138 blood and lymphatic vessels (Extended Data Fig. 2).

139 CLECs do not originate from hematopoietic niche in bone-marrow or spleen.

140 Given the abundant labelling of macrophages and the important role of Csflr in the 141 recruitment of bone-marrow derived monocytes and differentiated macrophages, we 142 asked whether CLECs are also derived from the bone-marrow. To test this possibility, we 143 generated bone marrow chimeras. Flow cytometric analysis confirmed 97.3 \pm 1.6 % 144 reconstitution of wild-type recipient bone marrow by transplanted Csf1r-Mer2.Cre^{mTmG} 145 bone marrow after 8 weeks post transplantation (Extended Data Fig. 3). Analysis of glioma blood vessels in these bone marrow chimeras (BM^{Csf1r-Mer2.Cre-mTmG}) failed to 146 147 identify bone marrow derived CLECs, although the numbers of tumour macrophage were 148 similar to controls (Fig. 3a). Conversely, chimeras of wild-type donor bone marrow transplanted into Csflr- $Mer2.Cre^{mTmG}$ recipients (Csflr- $Mer2.Cre^{mTmG}$::BM^{WT}), 149 150 displayed similar CLECs numbers in glioma vessels as observed earlier in non-chimeric *Csf1r-Mer2.Cre^{mTmG}* mice (Fig. 3b). These results were confirmed by intravital imaging in the glioma of *Csf1r-Mer2.Cre^{mTmG}*::BM^{WT} (Fig. 3c, Supplemental Movie 2). Thus, the bone marrow chimera and lineage tracing experiments strongly suggest that CLECs do not originate from bone marrow-derived cells, nor do they represent previously suggested transdifferentiation events from bone marrow-derived macrophages. Surgical removal of the spleen also did not affect CLECs numbers, further ruling out spleen-derived macrophages as source of CLECs (Extended Data Fig. 4).

158 *CLECs regulate vascular patterning and support glioma growth.*

159 In late-stage gliomas, CLECs preferentially localized to the peripheral tumor area, 160 correlating with the zone of most active angiogenesis (Extended Data Fig. 5). Real-time 161 intravital imaging of mouse gliomas also revealed that CLECs incorporation into vessels 162 is very dynamic. This implied that CLECs might have unique and potentially transient 163 functions during tumour vessel formation. To gain a better understanding of CLECs 164 behaviour during tumour growth, we developed dual-recombination combinatorial 165 genetics as a tool for CLECs tracing ^{23,24}. To achieve selective recombination in CLECs, 166 we used the *Cdh5* endothelial driver for expression of the Dre recombinase, and the same 167 transgenic mouse line as above expressing the tamoxifen inducible form of the Cre 168 recombinase under control of the Csf1r transgene. The combined action of these distinct 169 recombinases, Dre and Cre, allowed for deletion of two stop cassettes flanked by RoxP 170 and LoxP sites, respectively, located upstream of a tdTomato expression cassette (Fig. 171 4a). Using this strategy, we could selectively trace tdTomato-positive CLECs in mouse 172 gliomas following tamoxifen exposure (Fig. 4b). Note that the appearance of labelled 173 CLECs and blood vessel labelling differs from the images of the mTmG line, as the dual 174 Dre/Cre tdTomato reporter used lacks a general label of unrecombined cells. Fluorescent 175 dextran injection instead was used to label vascular lumen, highlighting dtTomato

176 positive CLECs that line the vessel lumen. As expected, counterstaining on fixed samples 177 demonstrated that CLECs express the prototypical endothelial marker, ERG (ETS-related 178 gene) (Fig. 4c). In order to assess whether CLECs contribute functionally to tumour 179 growth and vascular patterning in glioma, we used the same dual recombination strategy 180 to selectively ablate CLECs (Fig. 4d). In this model, CLECs expressed both GFP as well 181 as the diphtheria toxin receptor after tamoxifen induction enabling depletion of GFP-182 positive CLECs by administration of diphtheria toxin (Fig. 4e, h). Strikingly, CLECs 183 ablation significantly decreased glioma growth (Fig. 4g, i). CLECs ablation also 184 decreased the vascular network length and the number of bifurcations in tumour vessels 185 (Fig. 4j, k, l), indicating that CLECs may promote tumour growth by supporting vessel 186 branching.

187 CLECs express a unique set of markers

188 To further understand the nature of tumour CLECs (T CLECs), we performed single-cell 189 RNA sequencing (scRNA-seq) on isolated T_CLECs in the mouse glioma model. Single-190 cell transcriptome analysis using Smart-seq2²⁵ identified that the gene expression pattern 191 of T CLECs was similar to lineage-negative tumour endothelial cells (T ECs), but not 192 to tumour macrophages (T_MACs), as determined by UMAP (Uniform Manifold 193 Approximation and Projection) analysis (Fig. 5a). Single-cell differential expression 194 (SCDE) analysis ²⁶ confirmed that T CLECs expressed typical endothelial markers, but 195 lacked macrophage and myeloid cell markers (Fig. 5b). Unexpectedly, however, 196 T_CLECs showed no *Csf1r* expression, despite carrying the *Csf1r*-lineage marker (Fig. 197 5c). Beyond the overlapping gene expression pattern between T_CLECs and T_ECs, 198 SCDE analysis identified a unique signature that can distinctly define the T_CLECs 199 population; RNAs coding for 24 genes including three types of cell surface proteins 200 (Aqp1: Aquaporin 1, Fabp4: Fatty acid binding protein 4, Kcnj8: Potassium voltage*gated channel subfamily j member 8*) were significantly enriched in T_CLECs (Fig. 5d;
Extended Data Fig. 6; Supplementary Table 1). *Aqp1* is highly expressed in vascular
endothelial cells and was reported to contribute to tumour growth and angiogenesis ²⁷.
Staining of lineage traced T_CLECs on fixed mouse glioma samples confirmed AQP1
expression (Extended Data Fig. 7).

206 The T_CLEC state is enriched for "Tip-like" and a lymphatic gene signature

207 Gene set enrichment analysis (GSEA) identified that the T CLEC state shows significant 208 enrichment for a Tip-cell-like associated gene expression program (Fig. 5e), thus 209 supporting the morphological observations and their functional importance for branching 210 and tumour growth. Intriguingly, GSEA identified that T_CLECs were also enriched for lymphatic gene signatures (Fig. 5e). AUCell analysis ²⁸ of scRNA-seq data from multiple 211 212 human primary tumours identified subpopulations of endothelial cells that express the 213 T CLECs marker gene set (Fig. 5f-k, 442 out of 7314 ECs are candidates: 6.0%; 214 Supplementary Table 2). These bona-fide T_CLECs were highly enriched in tip cell, 215 neophalanx cell, lymphatic endothelial cell subpopulations and particularly abundant in 216 human pancreatic cancer (Fig. 5k). Neophalanx endothelial cells, which express two pro-217 angiogenesis markers, Glul (Glutamine synthetase) and Cxcl12 (C-X-C motif chemokine *ligand 12*) ^{29,30} have recently been identified in lung tumour-associated endothelial cell 218 219 studies ^{31,32}, and were proposed to line neo-vessels after termination of vessel sprouting. 220 Together this data suggests that based on the transcriptional state, T_CLECs might have 221 a supportive role in tumor vessel sprouting and remodeling. Detailed comparison with 222 recent single cell RNA seq data on tumour endothelial cells in mouse and human tumours 223 ³¹ confirmed the Tip-like gene signature of T_CLECs, but also identified significant 224 similarities with lymphatics (Fig. 5k). The latter is particularly surprising given that our

perfusion experiments with dextran clearly demonstrated that T_CLECs in the gliomamodel line blood vessels, not lymphatics.

227 Acute recombination identifies a CLEC lineage in lymph nodes

228 The transcriptional similarities between T_CLECs and lymphatic endothelial cells are 229 noteworthy as blood endothelial cells, in particular in the brain environment, and 230 lymphatic EC represent highly differentiated cell types. During vertebrate embryogenesis, 231 lymphovenous specification sees common progenitor cells diverge in their differentiation 232 to establish mature blood endothelium and lymphatic endothelium³³. The apparent 233 similarities could therefore suggest that T_CLECs represent an immature endothelial cell 234 state with partially overlapping profiles of both immature blood and lymphatic cells. In 235 this case, the expression of lymphatic genes would signify a return to an earlier cell state 236 or potentially some level of transdifferentiation between the two endothelial cell states.

The latter would imply a distant source of lymphatic endothelium as possible origin of CLECs. Indeed, when searching for CLECs in all organs, we observed selective endothelial labelling in lymph nodes, notably restricted to the periphery where specialized lymphatic endothelial cells form the floor of the subcapsular sinus (SCS, Fig. 6a). Bonemarrow transplantation to eliminate the abundant Csfr1-lineage macrophage population highlighted the specific labelling of CLECs in lymph nodes (Fig. 6b).

To gain more insight into the emergence of T_CLECs, we asked whether acute recombination could indicate their possible origin. Acute induction experiments (24h after single tamoxifen injection) illustrated that the Csf1r-expressing progenitors of this tumour endothelial cell population do not reside locally in pre-existing brain or the associated brain tumour vasculature (Extended Fig. 8). Close inspection however revealed GFP-positive endothelial cells with significant labelling 24h post-injection in the floor endothelial cells of the SCS of lymph nodes (Fig. 6c). The mTmG Cre-reporter

250 switches from red fluorescent protein expression to mEGFP expression upon 251 recombination. Therefore, cells that acutely recombine, carry both GFP and Tomato 252 expression until the latter is fully degraded. Strikingly, floor SCS endothelial cells 253 exhibited both GFP and simultaneous Tomato expression, demonstrating recent 254 recombination. FACS analysis confirmed the presence of double positive EC in lymph 255 nodes, but not in brain or brain tumours. In fact, double positive cells for GFP and Tomato 256 were not observed in the brain tumours at any stage, excluding the possibility of any acute 257 or continuous local recombination of T_CLECs within the tumour microenvironment. 258 Prox-1 staining confirmed the lymphatic nature of CLECs in the lymphnode, and 259 demonstrated that only a subpopulation of lymphatic endothelial cells in the lymph node 260 carry the Csfr1-lineage trace, the cells in the floor of the SCS (Fig. 6d).

261 We decided to perform scRNA seq also on endothelial cells isolated from lymph nodes,

assess gene expression and perform pseudotime analysis to understand differentiation of

263 T_CLECs, T_EC, LN_CLECs (GFP positive) and LN_ECs (GFP negative) (Fig. 7a-f).

We used the same Smart-seq2 platform for comparison to sequence another 384 cells of

lymph node origin. After quality control (Fig. 7a), 89 cells EC, not Csfr1 lineage EC, and

266 104 cells expressing GFP (Fig.7b) in the lymph nodes were selected for further analysis.

267 Comparative analysis with recent mouse lymph node scRNA data³⁴ confirmed that the

268 LN_CLEC co-register with identified SCS floor cells (Extended Data Fig. 9a).

269 Unlike T_CLECs, LN_CLECs also expressed Csf1r, in line with our finding of acute270 recombination in the SCS floor cells.

271 UMAP based dimension reduction placed T_CLECs and T_EC closely together, as before,

and also LN_CLEC and LN_EC co-cluster, although two additional distinct LN_EC

273 clusters appear highly distinct and thus unrelated to any of the other cell types (Fig. 7c).

274 Interestingly, some T_CLEC cluster with the LN_CLEC, suggesting that there is at least

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275 some overlap in gene expression between the Csf1r lineage cells in the brain tumour and 276 in the lymph nodes. Looking at relative expression levels of all genes in a heatmap 277 comparing the lineage positive and negative EC from lymph nodes and tumours 278 highlighted that differential expression is more prevalent than any overlap between 279 tumour EC and LN EC (Fig. 7d). However, there are small number of genes like Pde4d, 280 Robo1 and Itga9 that show higher expression in the two lineage positive populations in 281 LN and tumour, compared to the lineage negative cells (Extended Data Fig. 9b). Itga9 is well known to be important in lymphatic development³⁵, confirming again that T CLECs 282 283 carry some gene expression patterns normally associated with lymphatic EC. 284 Nevertheless, despite the fact that only LN_CLEC appear to acutely recombine, 285 differential gene expression analysis does not provide clear evidence for a lineage 286 relationship between LN_CLEC and T_CLEC.

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288 T EC and T CLECs "co-evolve" during tumour growth, suggesting distinct origin

289 To gain more insight into the differentiation state and possible route of differentiation, we next performed pseudotemporal ordering using Monocle³⁶ (Fig. 7f). The Monocle 290 291 inferred trajectory suggested equally distributed pseudotime values for T_CLECs and 292 T_EC then LN_EC and finally LN_CLECs. When measuring single-cell gene expression 293 diversity as surrogate for differentiation potential with the CytoTRACE algorithm³⁷ 294 (higher score means more undifferentiated, stem-like), we observed again equal scores 295 between T_CLECs and T_EC but lower cytoTRACE values for LN_EC and LN_CLECs 296 (Fig. 7e-f). The equal scores between T_CLECs and T_ECs argue against a role of T_ECs 297 as precursors for T_CLECs, independently confirming that T_CLECs are unlikely 298 emerging through differentiation from the pre-existing brain tumour vasculature. 299 However, the LN CLECs display the lowest cytoTRACE scores (most differentiated).

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300 Based on the cytoTRACE analysis, and its logic of developmental potential, it would 301 seem unlikely that LN_CLEC are the origin of T_CLEC (Fig. 7e-f). Moreover, it appears 302 that both lineage positive and negative EC in the lymph nodes are more differentiated 303 than endothelial cells in the tumour. It is noteworthy that Monocle was designed for 304 developmental questions, which are strictly unidirectional. Interestingly, when plotting 305 CytoTRACE scores onto the Monocle trajectory (Fig. 7f), the right side shows higher 306 scores (T CLEC and T EC) and the left side (darker blue) lower scores (LN CLECs 307 lowest), being in line with pseudotime. This again predicts that LN_CLECs represent an 308 end state rather than the origin. In summary, these data predict that during tumour growth, 309 the T_EC and T_CLECs coevolve from a distinct origin to then share many aspects of 310 their gene expression and biology, yet with a few very distinct gene profiles.

311

312 **DISCUSSION**

313 In this study, using inducible *Csf1r* lineage tracing in adult mice intended to study 314 macrophage recruitment during glioma development, we identify a distinct endothelial 315 cell lineage that forms up to 10% of the tumour vasculature in adult mice. Selective 316 experimental depletion of this Csflr endothelial cell lineage demonstrated a role in 317 promoting vascular branching and tumour growth. Through a combination of genetic 318 lineage tracing, longitudinal live two-photon imaging, and single cell analysis, we find 319 that these cells arise during tumour progression, adopt dynamic endothelial phenotypes 320 including tip cell characteristics, and express common endothelial as well as unique 321 markers that can also be found in patient-derived tumour endothelium.

322 Csflr is a well-known regulator of monocyte/ macrophage differentiation ^{38,39}, but is 323 also important for yolk sac hematopoiesis and progenitor cell differentiation. Klotz and 324 coworkers reported that Csflr-expressing cells in the mouse embryo contribute to Proxl-

325 positive lymphatic endothelial cells in the cardiac lymphatic vasculature ⁴⁰. Moreover, a 326 recent report identified that *Csf1r*-expressing erythro-myeloid progenitors from the early 327 embryonic yolk-sac could give rise to intraembryonic endothelial cells in mouse 328 development⁴¹. Csf1r is also expressed in EPCs, such as colony forming unit-Hill cells 329 and circulating angiogenic cells 42 . In the adult however, *Csf1r* is not expressed in the 330 mature endothelium of blood and lymphatic vessels, but continues to drive recruitment 331 and expansion of myeloid cells in inflammation, ischemia and tumour growth ⁴³. 332 Intriguingly, Csflr itself is no longer expressed in the Csflr lineage cells within the 333 tumour and acute recombination of endothelial cells 24h after tamoxifen injection fails to 334 label cells in the tumour vasculature. These results raise the possibility that a Csflr-335 expressing cell population of endothelial nature or potential is present elsewhere in the 336 body, recombines upon tamoxifen injection and is recruited to the tumour vasculature 337 where it differentiates into blood endothelium to contribute to tumour angiogenesis. 338 Alternatively, the fact that Csf1r expression is not detected in isolated tumour CLECs 339 could also indicate that the Cre-line is leaky and gives rise to labelled cells through 340 stochastic recombination. In that case, one would expect the labelled cells to populate any 341 part of the vasculature and be otherwise indistinguishable from the unlabelled tumour EC. 342 Single cell analysis however identifies a unique gene expression signature beyond the 343 large number of genes that are co-expressed in T_CLEC and T_EC, and demonstrates 344 that T CLEC are distinctly enriched in tip-cell and neo-phalanx cell populations. Thus, 345 labelled cells show distinct phenotypes, location and expression patterns, ruling out 346 purely stochastic recombination events. Furthermore, such stochastic recombination 347 within the tumour vasculature would necessitate that we identify acutely labelled cells 348 marked by co-expression of Tomato and GFP. As we failed to identify such cells at any 349 timepoint during tumour growth, and also could detect the emergence of T CLECs in

350 mice only injected with tamoxifen before tumour implantation, the cumulative evidence 351 would suggest that T_CLECs originate outside of the tumour and also that the progenitor 352 population resides outside of the brain. Despite all our efforts, the present work fails to 353 provide ultimate and definitive proof of the true origin of T_CLECs, but arrives at a clear 354 direction and working hypothesis for future studies. In search of their origin and a 355 potential progenitor population, we imaged and analysed all organs by FACS. We 356 performed bone-marrow transplantation, performed splenectomy and examined the entire 357 mouse by fluorescence stereomicroscopy after clearing to identify regions of 358 recombination. The results of bone-marrow transplantation effectively ruled out EPC or 359 any other known progenitors that may be found in circulation, including 360 transdifferentation from macrophages. Four independent observations placed a spotlight 361 on lymphatics, and the lymph nodes in particular. First, the only *bonafide* Csf1r lineage 362 endothelial cell population that we could identify outside of the tumours, including after 363 complete reconstitution of the bone marrow by wildtype bone marrow, was found in the 364 subcapsular sinus of the lymph nodes. Here the floor cells, specialized lymphatic 365 endothelial cells that we identify as GFP and Csf1r positive line the capsule in lymph nodes in close apposition to the Csf1 expressing ceiling cells. Second, after acute 366 367 tamoxifen injection the cells in the floor of the subcapsular sinus are uniquely double 368 positive for Tomato and GFP, demonstrating acute and likely local recombination. Third, 369 single cell RNAseq of T-CLECs identified lymphatic markers including the lymphatic 370 master regulator Prox1 co-expressed with blood endothelial markers, and SCDE placed 371 T_CLECs close to lymphatic endothelial cells from various mouse tumour models in 372 recently published datasets. And finally, CLECs can also be found in lymphatics in 373 subcutaneous tumour models and in skin. These independent observations clearly make

the SCS floor cells prime candidates for the origin of T_CLECs, but future work will be
necessary for a possible direct demonstration of this link.

376 In developmental studies, lineage relationships have recently been studied by scRNAseq 377 data sets through pseudotime analysis. In particular the CytoTRACE analysis can be 378 useful to estimate the degree of differentiation of cell populations and thereby also predict 379 their stemness ³⁷. Our comparative analysis of endothelial cells isolated from tumour and 380 lymph nodes assigned the highest differentiation scores to cells in the lymph node, and 381 lower values to the EC in tumour, including T_CLECs. Accordingly, the cells in the floor 382 of the SCS that carry the lineage label would rather be assumed at the end of a 383 differentiation process rather than progenitors for other populations such as the T_CLECs. 384 Whether however such algorithms are suitable to determine the potential of endothelial 385 cells to become activated and possibly transdifferentiate is unclear. In particular given the 386 altered environmental context of EC within a growing tumour compared to a vascular bed in homeostasis. The combined monocle³⁶ and cytoTRACE analysis would fit with a 387 388 general level of dedifferentiation of endothelial cells within the tumour microenvironment, 389 and therefore a co-evolution of the cell states of both T_EC and T_CLEC. It is likely that 390 pseudotemporal ordering of endothelial cells in a reactivation state will always show the 391 reactivated cells, like in the tumour environment as less differentiated. Indeed, recent 392 work studying the effect of chronic hypoxia on brain endothelium illustrated the wide 393 range of reactivation, in particular noting a strong increase in tip cell population amongst 394 the brain endothelial cells⁴⁴. The current assumption states that these new tip cells arise 395 under hypoxia from pre-existing brain endothelial cells, and possibly emerge from 396 particular endothelial subpopulations. In the absence of lineage tracing, an alternative 397 possibility, raised by our current observation in tumour angiogenesis, could be that new 398 tip cells arise at least in part elsewhere, and are recruited to the hypoxic vessel areas.

More work on vascular single cell analysis will be required to establish the fundamental
principles of endothelial activation and their interpretation for the different tissue
challenges that require vascular adaptations.

402 Angiogenesis is essential for tumour growth, and contributes to metastasis ^{45,46}. Early 403 hopes that targeting tumour angiogenesis may prove uniquely selective and effective as 404 anti-tumour treatment have, however, been disappointing. Intense research efforts 405 therefore currently focus on understanding reasons and mechanisms for resistance to anti-406 angiogenic therapy, and identifying new cellular and molecular targets to effectively 407 modify the tumour vasculature ⁴⁷. Intravital imaging shows that T_CLECs arise during 408 tumour progression, adopt endothelial phenotypes with tip cell characteristics and line 409 perfused vascular tubes. Consistent with this observation, AUCell analysis of scRNA-seq 410 data in human samples identifies that T_CLECs signatures are highly enriched in tip cell 411 subpopulation. A recent study using single-cell transcriptome analysis revealed that anti-412 VEGF (vascular endothelial growth factor) treatment reduces the proportion of 413 endothelial tip-like cells in a tumour xenografts ⁴⁸. How CLECs affect the outcome of 414 anti-angiogenic treatment will be important to investigate in the future. Considering that 415 T_CLECs prominently express *Vegfr3*, even more so than regular endothelial cells in the 416 tumour, we can speculate that T_CLECs are resistant to anti-VEGFR2 treatment, and 417 possibly escape treatment through the activity of VEGFR3. If so CLECs may adversely 418 affect anti-angiogenic therapy targeting VEGFR2 selectively. Tumour blood vessels are 419 generally highly tortuous and the branching pattern is greatly different from normal blood 420 vessels. Our results from the selective targeting of CLECs in the mouse glioma model 421 show that CLECs ablation decreases both vessel branching and tumour growth. Thus, in 422 the absence of CLEC, the dysmorphic vascular phenotype in tumours appears exacerbated, 423 suggesting that CLECs support the formation of a highly branched vascular network,

which in turn better promotes tumour growth. Future work will be required to identify the
true origin of T_CLECs and to extend this functional analysis to fully appreciate the
consequence and opportunities to target or utilize CLECs for therapeutic approaches.

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428

- 429 **METHODS**
- 430 *Mice*

Mice expressing both the Csflr-Mer-iCre-Mer transgene (Csflr-Mer2.Cre)¹⁷ and the 431 432 *Rosa^{mTmG}* Cre recombination reporter ¹⁸ were generated by breeding. Mice carrying the. Endothelial specific Cdh5-Dre mice ²⁴ were mated with Csf1r-Mer2.Cre mice and 433 434 Rosa26-RSR-LSL-tdTomato mice²³ to obtain Cdh5-Dre:: Csf1r-Mer2.Cre:: Rosa26-RSR-435 LSL-tdTomato mice (DreMer2.Cre^{tdTomato}) that allows combinatorial genetics as a tool for 436 CLECs tracing. Endothelial specific Cdh5-Dre mice were mated with Csf1r-Mer2.Cre 437 mice and Rosa26-RSR-mTmG-DTR mice to obtain Cdh5-Dre:: Csf1r-Mer2.Cre:: Rosa26-438 RSR-mTmG-DTR mice (DreMer2.Cre^{mTmG-DTR}) that allows for selective tracing and ablation of CLECs. Rosa26-RSR-mTmG-DTR mice were designed by Dr. Fabio Stanchi 439 440 and generated by the team of Dr. Ralf Kühn at the Max Delbrück Center in Germany. All 441 mouse strains were maintained on C57BL/6J background. Animal housing and all 442 experimental procedures were approved by the Institutional Animal Care and Research 443 Advisory Committee of the KU Leuven (085/2016).

444 Cell lines

445 CT2A, GL261 glioma cell lines and B16F1 melanoma cell line were cultured in DMEM
446 (Life Technologies) supplemented with 10% FBS (Life Technologies), 1% penicillin/
447 streptomycin (Life Technologies) and 1% glutamine (Life Technologies). Spheroids were

448 obtained by seeding the glioma cells for 48h on non-adherent culture dishes. Spheroids

449 of 200- 250µm were selected for implantation.

450 Intravital imaging on mouse gliomas

451 Surgery for tumor implantation, installation of cranial windows and in vivo imaging were 452 performed as described previously ¹⁶. Briefly, 8–12-weeks-old mice were anesthetized 453 with ketamine/ xylazine and a craniotomy was performed on the parietal bone. Spheroids 454 of CT2A or GL261 cells modified to express the blue fluorescent protein TagBFP ⁵⁰ were 455 injected in the exposed parietal brain cortex, which was then sealed by cementing a glass 456 coverslip to the bone surrounding the craniotomy. After surgery, mice were injected 457 intraperitoneally with tamoxifen (65 μ g/g body weight) every 5 days. For CLECs 458 depletion, mice were injected intraperitoneally with diphtheria toxin (4 ng/g body weight) 459 the next day after tamoxifen induction. Longitudinal *in vivo* imaging of growing tumors 460 was performed in mice under isoflurane anesthesia using a SP8 upright microscope (Leica 461 Microsystems) equipped with a HCX IRAPO L25x/0.95 water objective and a Titanium: 462 Sapphire laser (Vision II, Coherent Inc.) tuned at 925 nm. Humane end-point of 463 experiments was applied if animals lost 15-20% of their original weight or shown evident 464 signs of distress.

465 *Immunofluorescence imaging*

466 Mice anesthetized with ketamine/ xylazine were perfused through the heart with 15 ml of 467 ice-cold PBS, followed by 10 ml of 2% PFA in PBS. Tumour tissues or organs were 468 harvested and fixed overnight in 4% PFA. For tumour tissues, sections (200 µm-thick) 469 were prepared with a vibratome 650V (Thermo Scientific). For organ samples, frozen 470 sections (10 µm-thick) were prepared with a cryostat NX70 (Thermo Scientific). For 471 immunofluorescence imaging studies, the PFA-fixed sections were blocked and 472 permeabilized in TNBT (0.1 M Tris, pH 7.4, 150 mM NaCl, 0.5% blocking reagent from 473 Perkin Elmer, 0.5% Triton X-100) for 4h at room temperature. Sections were incubated

474 with antibodies against CDH5 (1:25, AF1002, R&D), CD31 (1:100, ab28364, Abcam), 475 F4/80 (1:100, MF48000, Invitrogen), CD45 (1:100, 550539, BD Pharmingen), CD169 476 (1:100, 142401, Biolegend), PROX1 (1:200, 11-002, Angiobio), AQP1 (1:4000, AB2219, 477 Millipore), LYVE1 (1:100, 50-0433-80, eBioscience) diluted in TNBT buffer overnight 478 at 4°C, washed in TNT buffer (0.1 M Tris pH 7.4; 150 mM NaCl, 0.5% Triton X-100) 479 and incubated with an Alexa Fluor conjugated antibodies (1:200, ThermoFisher 480 Scientific). Sections were washed and mounted in fluorescent mounting medium (Dako). 481 Images were obtained with a Leica TCS SP8 confocal microscope.

482 Flow cytometric analysis

483 Mice were anesthetized with ketamine/ xylazine, then tumour tissues or organs were 484 harvested and incubated with PBS containing 1mg/ml collagenase I (Gibco), 2 mg/ ml 485 Dispase I (Gibco), 100ug/ml DNase I (Roche) and 2 mM CaCl₂ for 30 min at 37 °C. After 486 incubation, the digested tissue was passed through a cell strainer and then washed by PBS 487 including 2% FBS. For red cell exclusion, we incubated samples with red cell lysis buffer 488 (Sigma) for 5 min at 37 °C and then washed by PBS including 2% FBS and 2 mM EDTA. 489 Cells were stained with the following monoclonal antibodies: PE/ Cv7 anti-CD45 490 (552848, BD), APC anti-CD31 (561814, BD). Data acquisition was performed with BD 491 FACSVerse and analysis was performed with BD FACSuite software, and FlowJo 492 software.

493 Mouse melanoma model

494 Mice (8–12-weeks-old) were anesthetized with ketamine/ xylazine and then 1×10^{6} B16F1 495 melanoma cells in PBS were implanted into the right flank region of the mice. B16F1 496 melanomas were removed from mice 12 days after implantation for immunofluorescent 497 imaging studies and flow cytometric analysis. Mice were injected intraperitoneally with 498 tamoxifen (65 µg/g body weight) 4 time/ 2 weeks before tumour implantation and then

499 intraperitoneally with tamoxifen (65 μ g/g body weight) every 5 days after tumour

500 implantation.

501 Single cell isolation and RNA-sequencing

502 Single cells were sorted (BD FACSAriaIII) in 96 well plates (VWR) containing 2 µL of 503 PBS including 0.2% Triton X-100 and 4U of RNase inhibitor (Takara) per well. Plates 504 were properly sealed and spun down at 2000 g for 1 min before storing at -80°C. Whole 505 transcriptome amplification was performed with a modified SMART-seq2 protocol as 506 described previously ²⁵, using 20 instead of 18 cycles of cDNA amplification. PCR 507 purification was realized with a 0.8:1 ratio (ampureXP beads:DNA). Amplified cDNA 508 quality was monitored with a high sensitivity DNA chip (Agilent) using the Bioanalyzer 509 (Agilent). Sequencing libraries were performed using the Nextera XT kit (Illumina) as 510 described previously ²⁵, using 1/4th of the recommended reagent volumes and 1/5th of 511 input DNA with a tagmentation time of 9 min. Library quality was monitored with a high 512 sensitivity DNA chip (Agilent) using the Bioanalyzer (Agilent). Indexing was performed 513 with the Nextera XT index Kit V2 (A-D). Up to 4 x 96 single cells were pooled per 514 sequencing lane. Samples were sequenced on the Illumina NextSeq 500 platform using 515 75bp single-end reads.

516 scRNA-sequencing data analysis

517 BAM files were converted to merged, demultiplexed FASTQ files, cleaned using fastq-518 mcf (ea-utils r819), and QC checked with FastQC (0.11.4). Reads were then mapped to 519 the mouse genome (mm18) using STAR (2.4.1b) and quantified with Subread (1.4.6-p2). 520 Cells with less than 100,000 reads and/or 500 genes expressed, more than 20% 521 mitochondrial reads, and less than an average expression level of 3.0 of about 80 522 housekeeping genes ⁵¹ were discarded. 469 cells passed these stringent quality criteria. 523 Winsorized Highly Variable genes (HVGs) were identified for tumor derived

524 macrophages (T MACs), endothelial cell (T ECs) and *Csf1r* lineage endothelial cells 525 (T_CLECs)²⁶. These cells were then clustered based on HVG expression using non-526 negative matrix factorization as dimension reduction approach (run=40, rank=10, in MeV 527 4.8.1). The "best fit" (numbers of clusters) was chosen based on the highest cophenetic 528 correlation coefficient. Next, Single-cell Differential Expression analysis (SCDE) was 529 performed between the different NMF-clusters using the global gene expression matrix 530 ²⁶. Expression values were library-size-factor-normalized (DESeq) and log2 transformed 531 (log2+1). T_MAC, T_EC, T_CLEC and LN_CLECS, LN_EC, T_CLEC, T_EC cells 532 were projected into a two-dimensional space using Uniform Manifold Approximation and 533 Projection (dims=15, resolution = 0.4) based on 5k most variable features (Seurat 534 pipeline)⁵². Monocle (2.14) based trajectory analysis was performed on T CLEC, T EC, 535 LN_CELC and LN_EC (total=454 cells) taking into account n=790 m3Drop genes. The 536 same data set was subjected to CytoTRACE analysis using the online tool 537 (https://cytotrace.stanford.edu/) ³⁷.

538

539 AUCell analysis of human scRNA-seq data in multiple tumours

540 The droplet-based scRNA-seq data of 4 human tumour types were downloaded from 541 publicly available sources including lung cancer (ArrayExpress: E-MTAB-6149 and E-542 MTAB-6653)⁵³, PDAC (GSA: CRA001160)⁵⁴, liver cancer (GEO: GSE125449)⁵⁵, and BCC (GEO: GSE123814)⁵⁶. These datasets were processed and clustered per tumour type 543 544 using Seurat (v3.0.4) package, and the endothelial cell clusters were identified using 545 marker genes (CLDN5, VWF, PECAM1). The resulting endothelial cell data were pooled 546 together and aligned using anchor-based canonical correlation analysis (CCA) to mitigate 547 the difference related to distinct tumor types and scRNA-seq technologies. Initial graph-548 based clustering identified 10 clusters including 3 potential doublet clusters as predicted

549 by DoubleFinder (v2), which were subsequently removed before second round of 550 clustering. The resulting 8 clusters consisting of 7314 endothelial cells were annotated 551 based on known marker genes, and their CLECs signature were calculated using AUCell 552 package (v1.6.1). The threshold of AUC score was chosen at the peak of the right modal, 553 and the percentage of putative CLECs were calculated for different subtypes and their 554 tumour type of origin.

555 Generation of bone marrow chimeras

Recipient 8-10-weeks-old mice were lethally irradiated (9.5 Gy) and then intravenously injected with 1x10⁷ bone marrow cells from donor mice 16 h later. Tumour implantation experiments were initiated 8 weeks after bone marrow reconstitution. Reconstitution rate was determined using flow cytometric analysis in the peripheral blood.

560 Surgical removal of the spleen

Mice (8–12-weeks-old) were anesthetized with ketamine/ xylazine and then the spleen was gently taken out from the connective tissue, while cauterizing associated vessels. The incision was thereafter sutured. These procedures were performed as described previously 564 ⁵⁷.

565 Vessel morphology analysis

566 Confocal images of the tumour vasculature were analyzed for two-dimensional vessel 567 area, network length and number of network bifurcation points. The tdTomato and GFP 568 channels were merged and a Gaussian blur filter as well as Otsu thresholding was applied 569 to obtain binary vessel mask for sectional area estimation. To obtain the vessel network, 570 a semi-automatic multicut workflow from ilastik ⁵⁸ was applied to all images. The 571 segmented vessels were skeletonized and converted into a network graph with nodes and 572 edges by using the Python package ImagePy ⁵⁹. The lengths of network edges were 573 summed up to obtain the total network length for each image, all nodes with a degree of

- 574 three or more were counted as bifurcations.
- 575 Statistics
- 576 Statistical analyses of data were performed with GraphPad Prism 7.0 software. All data
- 577 are shown as the means \pm standard deviations. Differences were assessed using a two-
- 578 tailed unpaired Mann-Whitney's U test or two-tailed unpaired Welch's correction.

579

580 AUTHOR CONTRIBUTIONS

581 KM led and conceived the project, designed and performed experiments, analysed and

582 interpreted data. FR, FS, TM, WG, LH, JQ performed experiments and analyzed data.

- 583 DL, BZ, CB, JM and HG analyzed data. HG conceived and supervised the project. KM
- and HG wrote and edited the manuscript, with contribution by all authors.

585

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597

598 CODE AVAILABILITY

24

- 599 The computational code (R) used for scRNA-seq analysis is based on already published
- 600 pipelines (see Material and Methods) and available upon request.

601

602 DATA AVAILABILITY

- 603 All scRNA-seq data are available at Gene Expression Omnibus under accession number
- 604 GSE157507.

605

606 STATISTICS

- 607 Differences were assessed using either a two-tailed unpaired Mann-Whitney's or two-
- tailed unpaired t-test with Welch correction. All planned comparisons were considered

stand-alone and were not corrected for multiple testing.

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775 FIGURE LEGENDS

776 Fig. 1: Csf1r lineage endothelial cells (CLECs) contribute to perfused, dynamically 777 sprouting and remodeling blood vessels in mouse glioma. a, Csf1r-Mer2.CremTmG: 778 a mouse model for Csf1r-lineage tracing combining the tamoxifen-inducible Cre-driver 779 under control of the Csf1r-promoter and the mTmG Cre-reporter. b, Timing schedule of 780 glioma implantation, tamoxifen induction and in vivo imaging. c-f, Longitudinal 781 intravital imaging of the tumour vasculature in CT2A mouse glioma of Csf1r-782 Mer2.CremTmG by cranial window. Scale bar: 50 µm. c, In vivo imaging in 3 weeks 783 glioma. Endothelial cells in red (tdTomato), Csf1r-labeled cells in green (GFP) and 784 glioma cells in blue (BFP). d, Detail image of outlined square in Fig. 1c illustrating tracing 785 of GFP-positive cells. White arrowheads, macrophages; blue arrowhead, CLECs at tip 786 position. e, In vivo imaging of CLECs in 2 weeks glioma. Blue arrowheads, tip of 787 endothelial cells. f, Visualization of blood vessels in 3 weeks glioma by Intravenous 788 injection of dextran with texas red dye. Yellow arrowhead, bridging CLECs; orange 789 arrowhead, tube formation by CLECs; blue arrowhead, CLECs at tip position. g-h, 790 Counterstaining on PFA-fixed 4 weeks CT2A glioma sections of Csf1r-Mer2.CremTmG 791 with indicated antibodies. Yellow arrows, CLECs. Scale bar: 25 µm.

Fig. 2: Up to 10% of intra-tumour endothelium is comprised of CLECs not only in glioma, but also in other tumour types and locations. a-i, Quantification of the population of CLECs in glioma of Csf1r-Mer2.CremTmG mice. a, Gross appearance of sham control and 4 weeks CT2A glioma (Upper panel). Purple arrowhead, the area of capillary injection; blue arrowhead, glioma. Scale bar: 5 mm. Flow cytometric analysis of CLECs in sham control and 4 weeks CT2A glioma (Lower panel). b, In 3 weeks CT2A

798 glioma (n= 9), CLECs constitute $2.1 \pm 1.0\%$ of total endothelial cells (CD45-CD31+); in 799 4 weeks CT2A glioma (n= 5), they make up $7.5 \pm 1.6\%$ of total endothelial cells. c, In 3 800 weeks GL261 glioma (n= 5), CLECs constitute $1.5 \pm 1.6\%$ of total endothelial cells. d, 801 Gross appearance of normal brain, sham control and 3 weeks CT2A glioma. Purple 802 arrowhead, area of capillary injection. Scale bar: 5 mm. e-f, Quantification of the 803 population of CLECs by flow cytometric analysis in brain and tumour samples. Bars 804 represent mean ± s.d. *p<0.05, **p<0.01, ***p<0.001, ****P<0.0001. Two-tailed 805 unpaired Mann-Whitney's U test. i-j, Analysis of CLECs in B16F1 melanoma in Csf1r-806 Mer2.CremTmG mice. g, Counterstaining on PFA-fixed section of day12 melanoma with 807 CDH5 antibody. Yellow arrows, CLECs. Scale bar: 50 μ m. h, CLECs constitute 2.8 \pm 808 0.75% of total endothelial cells in day 12 melanoma (n=4).

809 Fig. 3: CLECs do not originate from the bone marrow. a-b, Tamoxifen induction 810 timing. b, Quantification of the population of tumour macrophagens (T_MACs: 811 CD45+GFP+) and tumour CLECs (T_CLECs: CD45-CD31+GFP+) by flow cytometric 812 analysis in tumour samples (n=4). Bars represent mean \pm s.d. c, Fluorescent images of 813 PFA-fixed 3 weeks CT2A glioma de of Csf1r-Mer2.CremTmG mice within 24 hours after 814 tamoxifen induction. Scale bar: 100 µm. d-e, Flow cytometric analysis of CT2A-glioma 815 in bone marrow chimeras. d, BMCsf1r-mer2.CremTmG: C57BL/6J mice are used as 816 recipients and Csf1r-Mer2.CremTmG mice used as donors. Within the sorted myeloid 817 cell population (BFP-CD45+), $32.3 \pm 12.1\%$ are GFP-positive in Csf1r-Mer2.CremTmG 818 mice (n= 10), and $33.8 \pm 6.9\%$ are GFP-positive in bone-marrow transplanted BMCsf1r-819 Mer2.CremTmG mice (n= 12). Among endothelial cell population (BFP-CD45-CD31+), 820 $4.3 \pm 1.9\%$ are GFP-positive in Csf1r-Mer2.CremTmG mice (n= 10), whereas no GFP-821 positive endothelial cells are found in BMCsf1r-Mer2.CremTmG mice (n= 12). e, Csf1r-

822 Mer2.CremTmG:: BMWT: Csf1r-Mer2.CremTmG are used as recipients and C57BL/6J 823 as donors. Of the sorted myeloid cell population, $43.4 \pm 11.3\%$ GFP-positive cells are 824 found in Csf1r-Mer2.CremTmG mice (n= 6), whereas $0.1 \pm 0.1\%$ GFP-positive cells are 825 found in Csf1r-Mer2.CremTmG:: BMWT mice (n=4). Of the endothelial cell population, 826 $10.5 \pm 9.7\%$ GFP-positive cells are found in Csf1r-Mer2.CremTmG mice (n= 6), and 9.3 827 \pm 3.7% GFP-positive cells are found in Csf1r-Mer2.CremTmG:: BMWT mice (n= 4). 828 Bars represent mean ± s.d. *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001. Two-tailed 829 unpaired Mann-Whitney's U test. e, Longitudinal intravial imaging of 2 weeks CT2A 830 glioma of Csf1r-Mer2.CremTmG and Csf1r-Mer2.CremTmG :: BMWT mice. White 831 arrows, CLECs. Scale bar: 50 µm.

832 Fig. 4: Dual recombination lineage tracing and inducible depletion model identifies 833 a role for CLECs in tumour blood vessel patterning supporting glioma growth. a, 834 Cdh5-Dre: Csf1r-Mer2.Cre: RSR-LSL-tdTomato (DreMer2.CretdTomato): a genetic 835 mouse model for CLEC selective tracing by dual recombination. b-c, Analysis in 836 DreMer2.CretdTomato. b, Longitudinal intravital imaging of the tumour vasculature in 3 837 weeks CT2A mouse glioma by Intravenous injection of FITC dextran. Yellow 838 arrowheads, CLECs. Scale bar: 50 µm. c, ERG antibody counterstaining on PFA-fixed 839 sections of 3 weeks CT2A glioma. Yellow arrows, CLECs. Scale bar: 50 µm. d, Cdh5-Dre: Csf1r-Mer2.Cre: RSR-mTmG-DTR (DreMer2.CremTmG-DTR): a genetic mouse 840 841 model for CLEC selective tracing and inducible ablation. DTR, diphtheria toxin receptor. 842 e-m, Analysis in DreMer2.CremTmG-DTR. e, Longitudinal intravital imaging of the tumour vasculature in 3 weeks CT2A mouse glioma. Mer2.CremTmG, Csf1r-843 844 Mer2.CremTmG. Yellow arrowheads, CLECs. f, Timing schedule of glioma implantation, 845 tamoxifen induction and diphtheria toxin (DT) treatment. g, Gross appearance of 3 weeks

846 CT2A gliomas. Scale bar: 5 mm. h, Confocal images of PFA-fixed 3 weeks CT2A glioma 847 sections. Yellow arrows, CLECs. Scale bar: 50 μ m. i, Tumour weight at 3 weeks CT2A 848 glioma. j-l, Analysis of PFA-fixed 3 weeks CT2A glioma sections. j, Tumour blood 849 vessels area. k, Network length of tumour blood vessels. l, Number of bifurcations of 850 tumour blood vessels. Bars represent mean \pm s.d. *p<0.05, **p<0.01. Two-tailed unpaired 851 Mann-Whitney' s U test.

852 Fig. 5: Single-cell analysis identifies a unique signature that defines the tumour 853 CLECs. a-d, Single-cell RNA sequencing (scRNA-seq) analysis of tumour macrophages 854 (T_MACs: CD45+GFP+, 79 cells), tumour endothelial cell (T_ECs: CD45-CD31+GFP-, 855 78 cells), tumour CLECs (T_CLECs: CD45-CD31+GFP+, 141 cells) in 4 weeks CT2A 856 glioma of Csf1r-Mer2.CremTmG mice. b, Gene expression of typical endothelial cell 857 markers, macrophages and myeloid cell markers. c, Csf1r gene expression. d, Three 858 characteristic genes expressed in T_CLECs. e, Enrichment of Tip-cell signature in 859 T_CLECs compared to T_ECs as shown by GSEA. f-k, Analysis of scRNA-seq data of 860 human endothelial cells from multiple primary tumours. f, UMAP plot of human 861 endothelial cell subtypes. g, UMAP plot of human endothelial cells colour-coded for the 862 tumour type of origins. h, UMAP plots with marker gene expression for each EC subtype. 863 i, Heatmap of marker gene expression for each EC phenotype (each cell as a column and 864 each gene as a row). j-k, AUCell analysis of 24 T_CLECs markers. j, T_CLECs 865 subpopulation indicated (AUC>0.165) by red colored dots. k, Cell type-specific (left) and 866 tumour type-specific (right) populations of T_CLECs. BCC, basal cell carninoma: PDAC, 867 pancreatic ductal adenocarcinoma.

868 Fig. 6: CLECs are linked to Prox1-positive lymphatic endothelium of the
869 subcapsular sinus floor in lymph nodes. a, Fluorescent images of the indicated organs

35

870 of Csf1r-Mer2.CremTmG mice after multiple tamoxifen induction. Scale bar: 50 µm. b, 871 Quantification of the population of CLECs in Csf1r-Mer2.CremTmG mice by flow 872 cytmetric analysis. Of the endothelial cell population (CD45-CD31+), $9.8 \pm 2.1\%$ GFP-873 positive cells are found in the axillary lymph node of Csf1r-Mer2.CremTmG mice (n= 874 10). c, Counterstaining on PFA-fixed Inguinal lymph node of Csf1r-Mer2.CremTmG:: 875 BMWT mice using PROX1 antibody within 24 hours after tamoxifen induction. Yellow 876 arrowheads, GFP+tdTomato+PROX1+ cells in the floor of the subcapsular sinus. Scale 877 bar 25 µm. d, Fluorescent images of PROX1 antibody counterstaining on PFA-fixed 878 inguinal lymph node of Csf1r-Mer2.CremTmG::BMWT mice after multiple rounds of 879 tamoxifen induction. Scale bar: 100 µm. Detailed image of square outlined in d. Yellow 880 arrowheads, GFP+Prox1+ cells; blue arrowheads, GFP+Prox1- cells. h-k, Analysis of 881 CT2A glioma in Prox1-CreERT2mTmG mice.

882 Fig. 7: CLEC and EC heterogeneity in tumor and lymph node. a, Quality measures 883 of 4 SMARTseq2 scRNA-seq. libraries: Csf1r-lineage endothelial cells (CLECs) and 884 endothelial cells (ECs) isolated from tumor (T_) and lymph node (LN_). b, Violin plots 885 show GFP expression per single cell. c, Uniform manifold approximation and projection 886 (UMAP) of T CLECs, T ECs and LN CLECs, LN ECs. d, Heatmap shows the 20 most 887 characteristic markers per cell population. e, UMAP colored by gene expression diversity 888 (CytoTRACE score) and predicted ordering by CytoTRACE. f, Monocle based trajectory 889 analysis of four cell populations, colored by pseudotime and CytoTRACE score.

Extended Data Fig. 1: CLECs in mouse glioma express endothelial cell markers, but lack macrophage and myeloid cell markers. Counterstaining on PFA-fixed 4 weeks CT2A glioma sections of Csf1r-Mer2.CremTmG with indicated antibodies. Lower panels

show the detail images of outlined squares in upper panels. Yellow arrowheads, CLECs;
blue arrowheads, macrophages. Scale bar: 25 μm.

895 Extended Data Fig. 2: CLECs contribute to lymphatic endothelium in the tumour

896 environment. Fluorescent images of counterstaining on PFA-fixed section of day 12

897 B16F1 melanoma under ear skin of Csf1r-Mer2.CremTmG mice using LYVE1 antibody.

898 Lower panel indicates detail image of yellow outlined square in upper panel. Yellow

arrow heads, CLECs. Scale bar: 100 μm.

Extended Data Fig. 3: Reconstitution rate in bone marrow chimeras. Reconstitution
rate was determined in peripheral blood 8 weeks after bone marrow transplantation.
TdTomato-positive cells in blood of: C57BL/6J (0.3 ± 0.1%, n= 6), Csf1rMer2.CremTmG (99.2 ± 0.5%, n= 7), BMCsf1r-Mer2.CremTmG (97.3 ± 1.6%, n= 13).
Bars represent mean ± s.d. ****p<0.0001. Two-tailed unpaired Mann-Whitney' s U test.

905 Extended Data Fig. 4: CLECs do not originate from spleen. Surgical removal of 906 spleen (splenectomy) was performed 3 weeks before glioma injection. CLECs isolated 907 from CT2A glioma of sham control Csf1r-Mer2.CremTmG mice (n= 5) constitute $1.5 \pm$ 908 0.8 % of total tumour endothelial cell population, compared to $1.5 \pm 0.5\%$ found in Csf1r-909 Mer2.CremTmG with splenectomy (n= 4). Bars represent mean \pm s.d. Two-tailed 910 unpaired Mann-Whitney' s U test.

⁹¹¹ Extended Data Fig. 5: CLECs locate at peripheral area of mouse glioma. a,
912 Vibratome section (200 µm-thick) of 4 weeks CT2A glioma of Csf1r-Mer2.CremTmG::
913 BMWT mice. Scale bar: 1 mm. b, Fluorescent images of area outlined in a. Yellow
914 arrowheads, CLECs. Scale bar: 100 µm.

915 Extended Data Fig. 6: 24 tumour CLECs markers in mouse glioma. SCDE analysis

- 916 of scRNA-seq data of tumour macrophages (T_MACs: CD45+GFP+, n= 79), tumour
- 917 endothelial cells (T_ECs: CD45-CD31+GFP-, n=78), tumour CLECs (T_CLECs: CD45-
- 918 CD31+GFP+, n= 141) in 4 weeks CT2A glioma in Csf1r-Mer2.CremTmG mice. Bars
- 919 represent mean ± s.d. *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001. Two-tailed
- 920 unpaired t-test with Welch correction.

921 Extended Data Fig. 7: CLECs in glioma express AQP1. Counterstaining on PFA-fixed

922 CT2A glioma of Csf1r-Mer2.CremTmG::BMWT mice using AQP1 antibody. Yellow

923 arrowheads, T_CLECs; blue arrowheads, T_ECs. Scale bar: 25 μm.

924 Extended Data Fig. 8: CLECs in glioma cannot be labelled by GFP within 24 hours

925 after tamoxifen induction. a-b, Tamoxifen induction timing. b, Quantification of the

926 population of tumour macrophagens (T_MACs: CD45+GFP+) and tumour CLECs

927 (T_CLECs: CD45-CD31+GFP+) by flow cytometric analysis in tumour samples (n=4).

Bars represent mean \pm s.d. c, Fluorescent images of PFA-fixed 3 weeks CT2A glioma de

929 of Csf1r-Mer2.CremTmG mice within 24 hours after tamoxifen induction. Scale bar: 100

930 μm.

931 Extended Data Fig. 9: CLEC and EC heterogeneity in tumor and lymph node.

932 a, Heatmap shows the expression of n=91 fLECs markers in LN_CLEC, LN_EC,

933 T_CLEC and T_EC single cells. b, Violin plots show the expression of 3 CLEC markers:

934 Pde4d, Robo1 and Itga9.

935 Supplemental Movie 1: Time-lapse *in vivo* imaging of Csf1r lineage endothelial tip

936 cell in CT2A glioma of *Csf1r-Mer2.Cre^{mTmG}* mice. Imaging was performed over 4 hours
937 (1 frame/ 10 min).

938

939 Supplemental Movie 2: Time-lapse *in vivo* imaging of *Csf1r* lineage endothelial tip
940 cell in CT2A glioma of *Csf1r-Mer2.Cre^{mTmG}*::BM^{WT}. Imaging was performed over 50
941 minutes (1 frame/ 2.5 min).

942

943 Supplemental Table 1: Details of gene expression analysis. Seurat based differential 944 analysis (non-parameteric Wilcoxon rank sum test) was performed by comparing cell 945 types amongst each other (green columns) and the different Seurat clusters (yellow 946 columns). p_val: p_val (unadjusted). avg_logFC: log fold-change of the average 947 expression between the two groups. Positive values indicate that the feature is more 948 highly expressed in the first group. pct.1: The percentage of cells where the feature is 949 detected in the first group. pct.2: The percentage of cells where the feature is detected in 950 the second group. p val adj: Adjusted p-value, based on Bonferroni correction using all 951 features in the dataset.

952

953 Supplemental Table 2: Marker genes for EC subtypes of multiple tumour types.

954 p_val: p_val (unadjusted). avg_logFC: Average log (fold change). pct.1: Percentage of

955 expressed cells in current cluster. pct.2: Percentage of expressed cells in other cluster.

- 956 p_val_adj: Adjusted p-value. cluster: Cluster name. gene: Gene name. log2fc: log2 (fold
- 957 change). Log2pct: log2 ((pct+0.005)/(pct2+0.005)).

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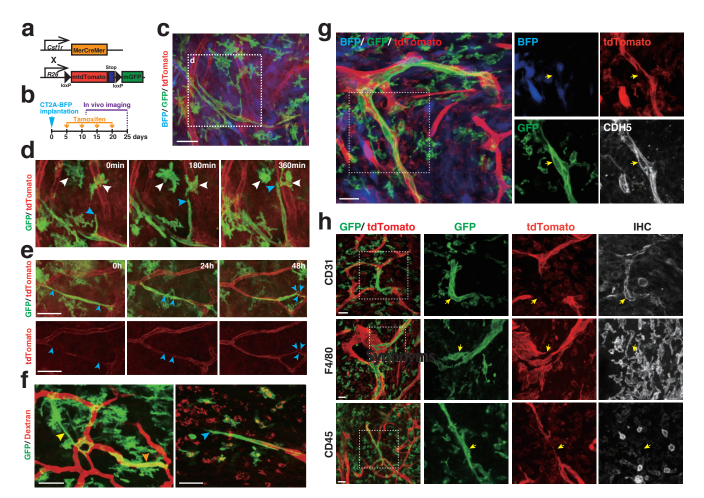


Fig. 1: *Csf1r* lineage endothelial cells (CLECs) contribute to perfused, dynamically sprouting and remodeling blood vessels in **mouse glioma.** a, *Csf1r-Mer2.Cre^{mTmG}*: a mouse model for *Csf1r*-lineage tracing combining the tamoxifen-inducible Cre-driver under control of the *Csf1r*-promoter and the *mTmG* Cre-reporter. b, Timing schedule of glioma implantation, tamoxifen induction and in vivo imaging. c-f, Longitudinal intravital imaging of the tumour vasculature in CT2A mouse glioma of *Csf1r-Mer2.Cre^{mTmG}* by cranial window. Scale bar: 50 μm. c, In vivo imaging in 3 weeks glioma. Endothelial cells in red (tdTomato), *Csf1r*-labeled cells in green (GFP) and glioma cells in blue (BFP). d, Detail image of outlined square in Fig. 1c illustrating tracing of GFP-positive cells. White arrowheads, macrophages; blue arrowhead, CLECs at tip position. e, In vivo imaging of CLECs in 2 weeks glioma. Blue arrowheads, tip of endothelial cells. f, Visualization of blood vessels in 3 weeks glioma by Intravenous injection of dextran with texas red dye. Yellow arrowhead, bridging CLECs; orange arrowhead, tube formation by CLECs; blue arrowhead, CLECs at tip position. g-h, Counterstaining on PFA-fixed 4 weeks CT2A glioma sections of *Csf1r-Mer2.Cre^{mTmG}* with indicated antibodies. Yellow arrows, CLECs. Scale bar: 25 μm.

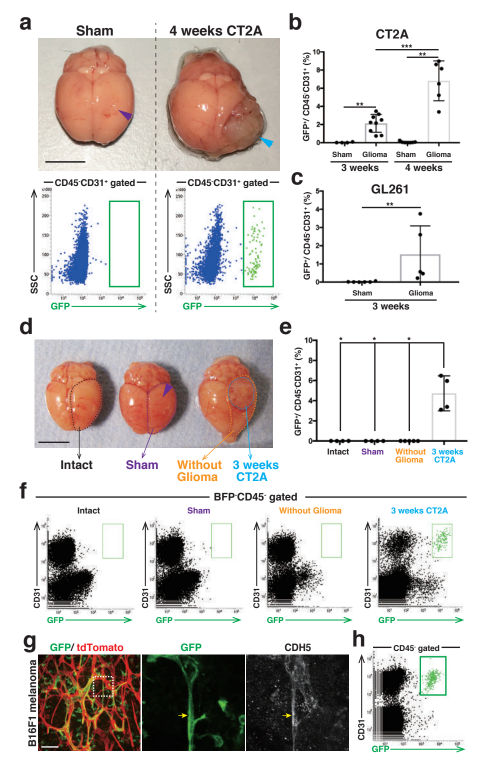


Fig. 2: Up to 10% of intra-tumour endothelium is comprised of CLECs not only in glioma, but also in other tumour types and **locations.** a-f, Quantification of the population of CLECs in glioma of Csf1r-Mer2.Cre^{mTmG} mice. a, Gross appearance of sham control and 4 weeks CT2A glioma (Upper panel). Purple arrowhead, the area of capillary injection; blue arrowhead, glioma. Scale bar: 5 mm. Flow cytometric analysis of CLECs in sham control and 4 weeks CT2A glioma (Lower panel). b, In 3 weeks CT2A glioma (n= 9), CLECs constitute 2.1 ± 1.0% of total endothelial cells (CD45-CD31+); in 4 weeks CT2A glioma (n= 5), they make up 7.5 ± 1.6% of total endothelial cells. c, In 3 weeks GL261 glioma (n= 5), CLECs constitute 1.5 ± 1.6% of total endothelial cells. d, Gross appearance of normal brain, sham control and 3 weeks CT2A glioma. Purple arrowhead, area of capillary injection. Scale bar: 5 mm. e-f, Quantification of the population of CLECs by flow cytometric analysis in brain and tumour samples. Bars represent mean ± s.d. *P<0.05, **P<0.01, ***P<0.001, ****P<0.001. Two-tailed unpaired Mann-Whitney' s U test. g-h, Analysis of CLECs in B16F1 melanoma in Csf1r-Mer2.CremTmG mice. g, Counterstaining on PFA-fixed section of day12 melanoma with CDH5 antibody. Yellow arrows, CLECs. Scale bar: 50 μm. h, CLECs constitute 2.8 ± 0.75% of total endothelial cells in day 12 melanoma (n=4).

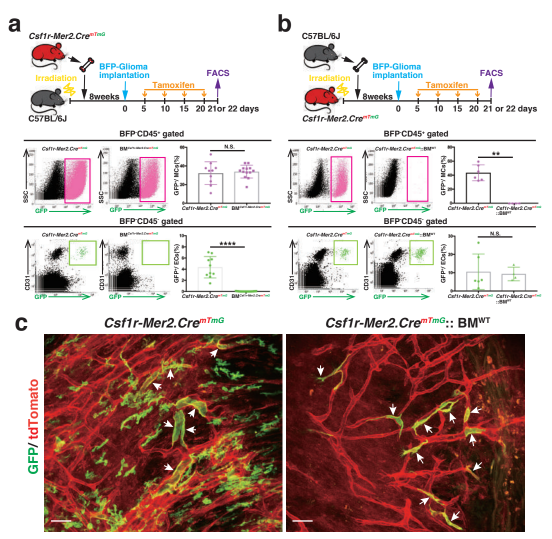


Fig. 3: CLECs do not originate from bone marrow. a-b, Flow cytometric analysis of CT2A-glioma in bone marrow chimeras. a, BM^{Csf1r-mer2.CremTmG}: C57BL/6J mice are used as recipients and *Csf1r-Mer2.Cre^{mTmG}* mice used as donors. Within the sorted myeloid cell population (BFP⁻CD45⁺), 32.3 ± 12.1% are GFP-positive in *Csf1r-Mer2.Cre^{mTmG}* mice (n= 10), and 33.8 ± 6.9% are GFP-positive in bone-marrow transplanted BM^{Csf1r-Mer2.CremTmG} mice (n= 12). Among endothelial cell population (BFP⁻CD45⁺CD31⁺), 4.3 ± 1.9% are GFP-positive in *Csf1r-Mer2.Cre^{mTmG}* mice (n= 10), whereas no GFP-positive endothelial cells are found in BM^{Csf1r-Mer2.CremTmG} mice (n= 12). b, *Csf1r-Mer2.Cre^{mTmG}*:: BM^{WT}: *Csf1r-Mer2.Cre^{mTmG}* are used as recipients and C57BL/6J as donors. Of the sorted myeloid cell population, 43.4 ± 11.3% GFP-positive cells are found in *Csf1r-Mer2.Cre^{mTmG}* mice (n= 6), whereas 0.1 ± 0.1% GFP-positive cells are found in *Csf1r-Mer2.Cre^{mTmG}*:: BM^{WT} mice (n= 4). Of the endothelial cell population, 10.5 ± 9.7% GFP-positive cells are found in *Csf1r-Mer2.Cre^{mTmG}* mice (n= 4). Bars represent mean ± s.d. *P<0.05,**P<0.01,***P<0.001, ****P<0.0001. Two-tailed unpaired Mann-Whitney' s U test. c, Longitudinal intravial imaging of 2 weeks CT2A glioma of *Csf1r-Mer2.Cre^{mTmG}* and *Csf1r-Mer2.Cre^{mTmG}* :: BM^{WT} mice. White arrows, CLECs. Scale bar: 50 μm.

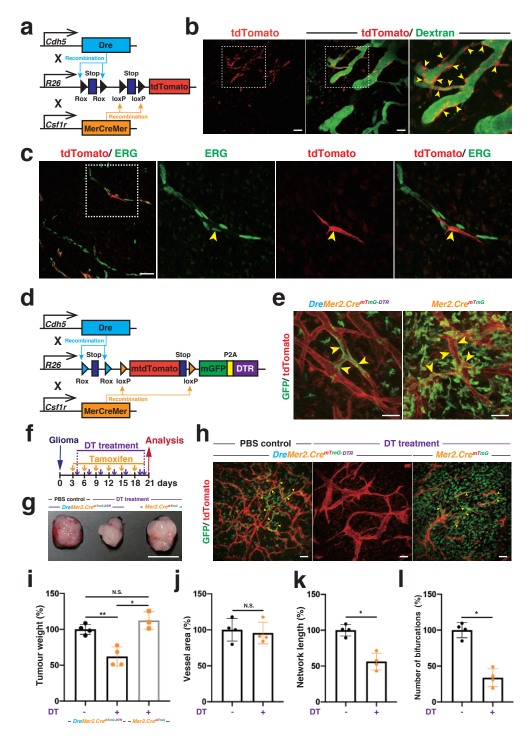
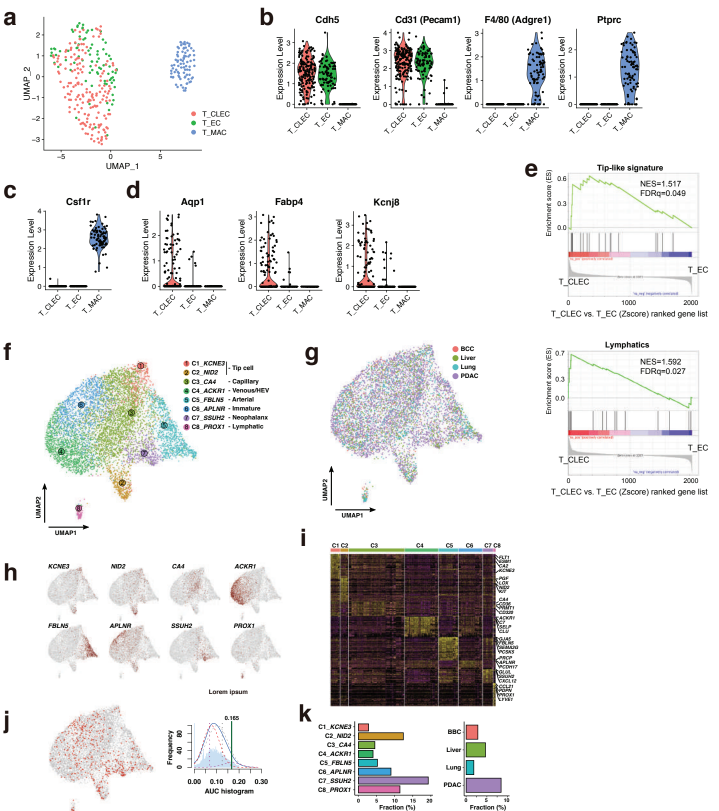
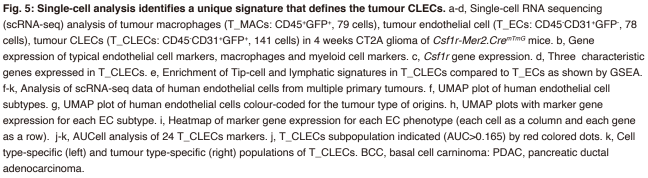


Fig. 4: Dual recombination lineage tracing and inducible depletion model identifies a role for CLECs in tumour blood vessel patterning supporting glioma growth. a, *Cdh5-Dre: Csf1r-Mer2.Cre: RSR-LSL-tdTomato* (*DreMer2.Cre^{ldTomato}*): a genetic mouse model for CLEC selective tracing by dual recombination. b-c, Analysis in *DreMer2.Cre^{ldTomato}*. b, Longitudinal intravital imaging of the tumour vasculature in 3 weeks CT2A mouse glioma by Intravenous injection of FITC dextran. Yellow arrowheads, CLECs. Scale bar: 50 μm. c, ERG antibody counterstaining on PFA-fixed sections of 3 weeks CT2A glioma. Yellow arrows, CLECs. Scale bar: 50 μm. d, *Cdh5-Dre: Csf1r-Mer2.Cre: RSR-mTmG-DTR* (*DreMer2.Cre^{mTmG-DTR}*): a genetic mouse model for CLEC selective tracing and inducible ablation. DTR, diphtheria toxin receptor. e-I, Analysis in *DreMer2.Cre^{mTmG-DTR}*. e, Longitudinal intravital imaging of the tumour vasculature in 3 weeks CT2A mouse glioma. *Mer2.Cre^{mTmG}*, *Csf1r-Mer2.Cre^{mTmG-DTR}*. Yellow arrowheads, CLECs. f, Timing schedule of glioma implantation, tamoxifen induction and diphtheria toxin (DT) treatment. g, Gross appearance of 3 weeks CT2A gliomas. Scale bar: 5 mm. h, Confocal images of PFA-fixed 3 weeks CT2A glioma sections. Yellow arrows, CLECs. Scale bar: 50 μm. i, Tumour weight at 3 weeks CT2A glioma. j-I, Analysis of PFA-fixed 3 weeks CT2A glioma sections. j, Tumour blood vessels area. k, Network length of tumour blood vessels. I, Number of bifurcations of tumour blood vessels. Bars represent mean ± s.d. *P<0.05, **P<0.01. Two-tailed unpaired Mann-Whitney' s U test.

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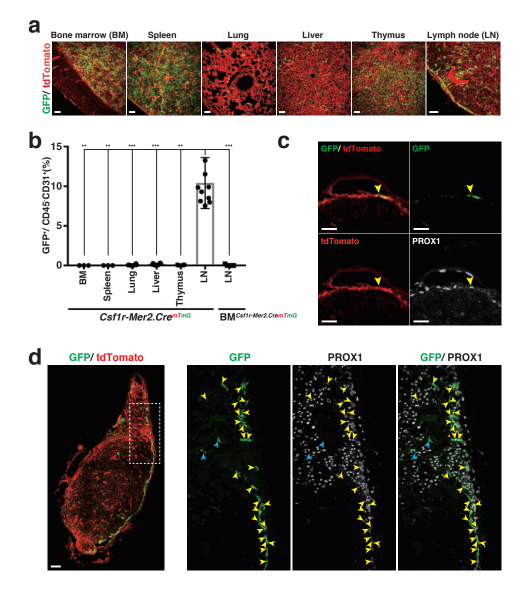
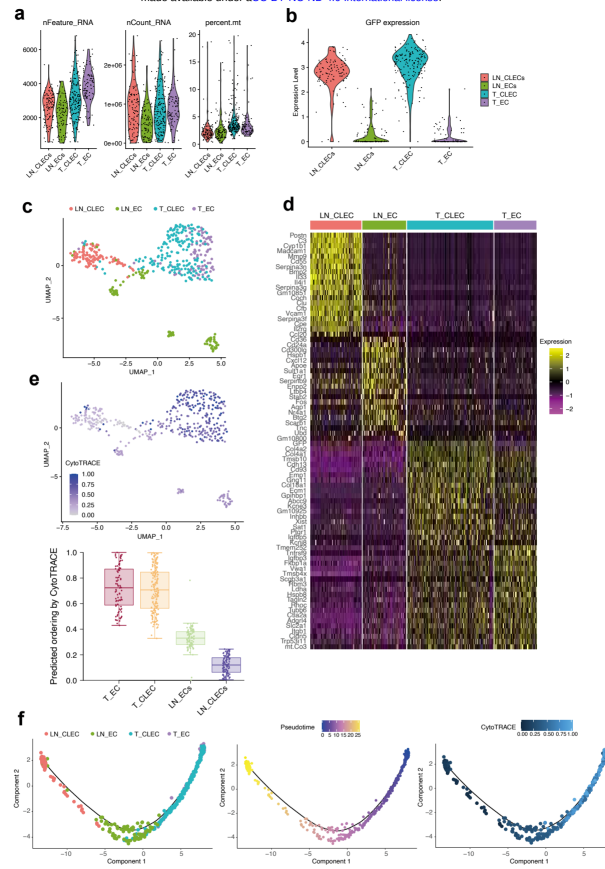


Fig. 6: CLECs are linked to *Prox1*-positive lymphatic endothelium of the subcapsular sinus floor in lymph nodes. a, Fluorescent images of the indicated organs of *Csf1r-Mer2.Cre^{mTmG}* mice after multiple tamoxifen induction. Scale bar: $50 \ \mu$ m. b, Quantification of the population of CLECs in *Csf1r-Mer2.Cre^{mTmG}* mice by flow cytmetric analysis. Of the endothelial cell population (CD45⁻CD31⁺), $9.8 \pm 2.1\%$ GFP-positive cells are found in the axillary lymph node of *Csf1r-Mer2.Cre^{mTmG}* mice (n= 10). c, Counterstaining on PFA-fixed Inguinal lymph node of *Csf1r-Mer2.Cre^{mTmG}*:: BM^{WT} mice using PROX1 antibody within 24 hours after tamoxifen induction. Yellow arrowheads, GFP⁺tdTomato⁺PROX1⁺ cells in the floor of the subcapsular sinus. Scale bar $25 \ \mu$ m. d, Fluorescent images of PROX1 antibody counterstaining on PFA-fixed inguinal lymph node of *Csf1r-Mer2.Cre^{mTmG}*::BM^{WT} mice after multiple rounds of tamoxifen induction. Scale bar: $100 \ \mu$ m. Detailed image of square outlined in d. Yellow arrowheads, GFP⁺Prox1⁺ cells; blue arrowheads, GFP⁺Prox1⁻ cells.





a, Quality measures of 4 SMARTseq2 scRNA-seq. libraries: Csf1r-lineage endothelial cells (CLECs) and endothelial cells (ECs) isolated from tumor (T_) and lymph node (LN_). b, Violin plots show GFP expression per single cell. c, Uniform manifold approximation and projection (UMAP) of T_CLECs, T_ECs and LN_CLECs, LN_ECs. d, Heatmap shows the 20 most characteristic markers per cell population. e, UMAP colored by gene expression diversity (CytoTRACE score) and predicted ordering by CytoTRACE. f, Monocle based trajectory analysis of four cell populations, colored by pseudotime and CytoTRACE score.