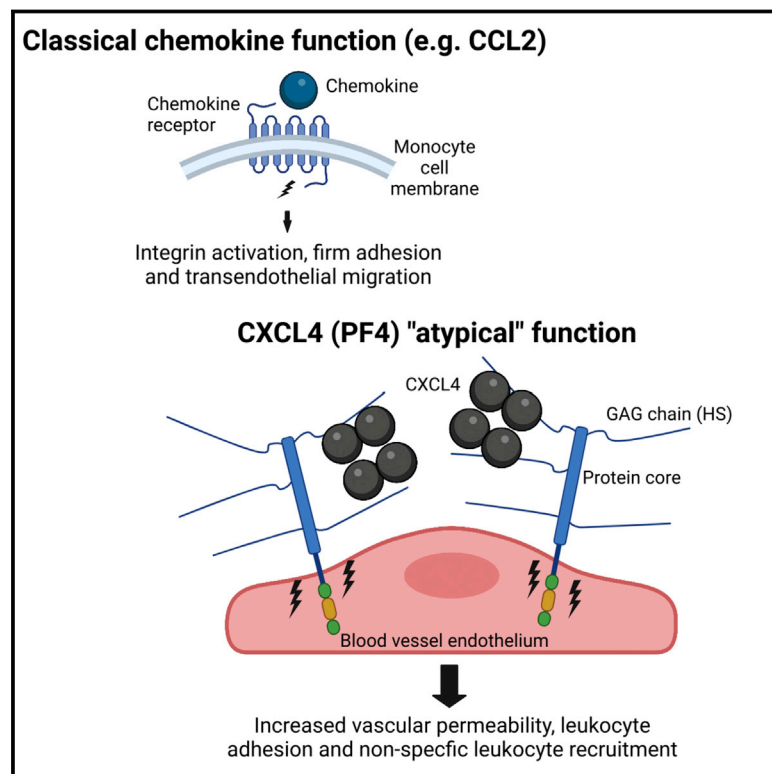


Chemokine CXCL4 interactions with extracellular matrix proteoglycans mediate widespread immune cell recruitment independent of chemokine receptors

Graphical abstract



Authors

Anna L. Gray, Richard Karlsson, Abigail R.E. Roberts, ..., Ralf P. Richter, Rebecca L. Miller, Douglas P. Dyer

Correspondence

douglas.dyer@manchester.ac.uk

In brief

Chemokines are central to immune cell recruitment but have not been therapeutically targeted during inflammation due to lack of full understanding of their action mechanisms. Gray et al. show that CXCL4 binds to endothelial extracellular matrix proteoglycans producing immune cell recruitment.

Highlights

- CXCL4 (PF4) mediates non-specific recruitment of a range of immune cells
- CXCL4 functions by binding to glycosaminoglycans on endothelial proteoglycans
- CXCL4 cross-linking of glycosaminoglycans produces endothelial signaling
- Glycosaminoglycan sulfation mediates specific chemokine interactions



Article

Chemokine CXCL4 interactions with extracellular matrix proteoglycans mediate widespread immune cell recruitment independent of chemokine receptors

Anna L. Gray,^{1,2} Richard Karlsson,³ Abigail R.E. Roberts,⁴ Amanda J.L. Ridley,¹ Nabina Pun,¹ Bakhtbilland Khan,¹ Craig Lawless,¹ Rafael Luís,^{5,6,7} Martyna Szpakowska,⁵ Andy Chevné,⁵ Catherine E. Hughes,⁸ Laura Medina-Ruiz,⁸ Holly L. Birchenough,¹ Iashia Z. Mulholland,¹ Catherina L. Salanga,⁹ Edwin A. Yates,¹⁰ Jeremy E. Turnbull,^{3,10,11} Tracy M. Handel,⁹ Gerard J. Graham,⁸ Thomas A. Jowitt,¹ Ingo Schiessl,^{2,12} Ralf P. Richter,⁴ Rebecca L. Miller,³ and Douglas P. Dyer^{1,2,13,*}

¹Wellcome Centre for Cell-Matrix Research, Lydia Becker Institute of Immunology and Inflammation, Faculty of Biology, Medicine and Health, Manchester Academic Health Science Centre, University of Manchester, Manchester, UK

²Geoffrey Jefferson Brain Research Centre, Manchester Academic Health Science Centre, Northern Care Alliance NHS Group, University of Manchester, Manchester, UK

³Copenhagen Center for Glycomics, Department of Cellular and Molecular Medicine, Faculty of Health Sciences, University of Copenhagen, Blegdamsvej 3, 2200 Copenhagen, Denmark

⁴University of Leeds, School of Biomedical Sciences, Faculty of Biological Sciences, School of Physics and Astronomy, Faculty of Engineering and Physical Sciences, Astbury Centre for Structural Molecular Biology, and Bragg Centre for Materials Research, Leeds LS2 9JT, UK

⁵Immuno-Pharmacology and Interactomics, Department of Infection and Immunity, Luxembourg Institute of Health, 4354 Esch-sur-Alzette, Luxembourg

⁶Faculty of Science, Technology and Medicine, University of Luxembourg, Esch-sur-Alzette, Luxembourg

⁷Tumor Immunotherapy and Microenvironment, Department of Cancer Research, Luxembourg Institute of Health, 2012 Luxembourg, Luxembourg

⁸Chemokine Research Group, School of Infection and Immunity, College of Medical, Veterinary and Life Sciences, University of Glasgow, Glasgow G12 8TA, UK

⁹Skaggs School of Pharmacy and Pharmaceutical Sciences, University of California, San Diego, La Jolla, CA 92093, USA

¹⁰Department of Biochemistry and Systems Biology, Institute of Systems, Molecular and Integrative Biology, University of Liverpool, Liverpool L69 7ZB, UK

¹¹Centre for Glycosciences, Keele University, Keele, Staffordshire ST5 5BG, UK

¹²Division of Neuroscience, School of Biological Sciences, Faculty of Biology, Medicine and Health, University of Manchester, Manchester, UK

¹³Lead contact

*Correspondence: douglas.dyer@manchester.ac.uk

<https://doi.org/10.1016/j.celrep.2022.111930>

SUMMARY

Leukocyte recruitment from the vasculature into tissues is a crucial component of the immune system but is also key to inflammatory disease. Chemokines are central to this process but have yet to be therapeutically targeted during inflammation due to a lack of mechanistic understanding. Specifically, CXCL4 (Platelet Factor 4, PF4) has no established receptor that explains its function. Here, we use biophysical, *in vitro*, and *in vivo* techniques to determine the mechanism underlying CXCL4-mediated leukocyte recruitment. We demonstrate that CXCL4 binds to glycosaminoglycan (GAG) sugars on proteoglycans within the endothelial extracellular matrix, resulting in increased adhesion of leukocytes to the vasculature, increased vascular permeability, and non-specific recruitment of a range of leukocytes. Furthermore, GAG sulfation confers selectivity onto chemokine localization. These findings present mechanistic insights into chemokine biology and provide future therapeutic targets.

INTRODUCTION

Leukocyte recruitment is central to fighting infection, coordinating the immune response to injury, and driving inflammatory disease.¹ Chemokines are critical during leukocyte recruitment, facilitating firm adhesion of leukocytes to the vascular endothelium.² Due to the central role of chemokines in many diseases,

they are prime therapeutic targets,³ but drugs that successfully target them are lacking.⁴ This is, in part, due to an under-developed understanding of the mechanisms underlying chemokine biology.

The dogma of the chemokine field states that they bind to G protein-coupled receptors (GPCRs) on leukocytes, producing integrin activation and firm adhesion to the vascular endothelium.²



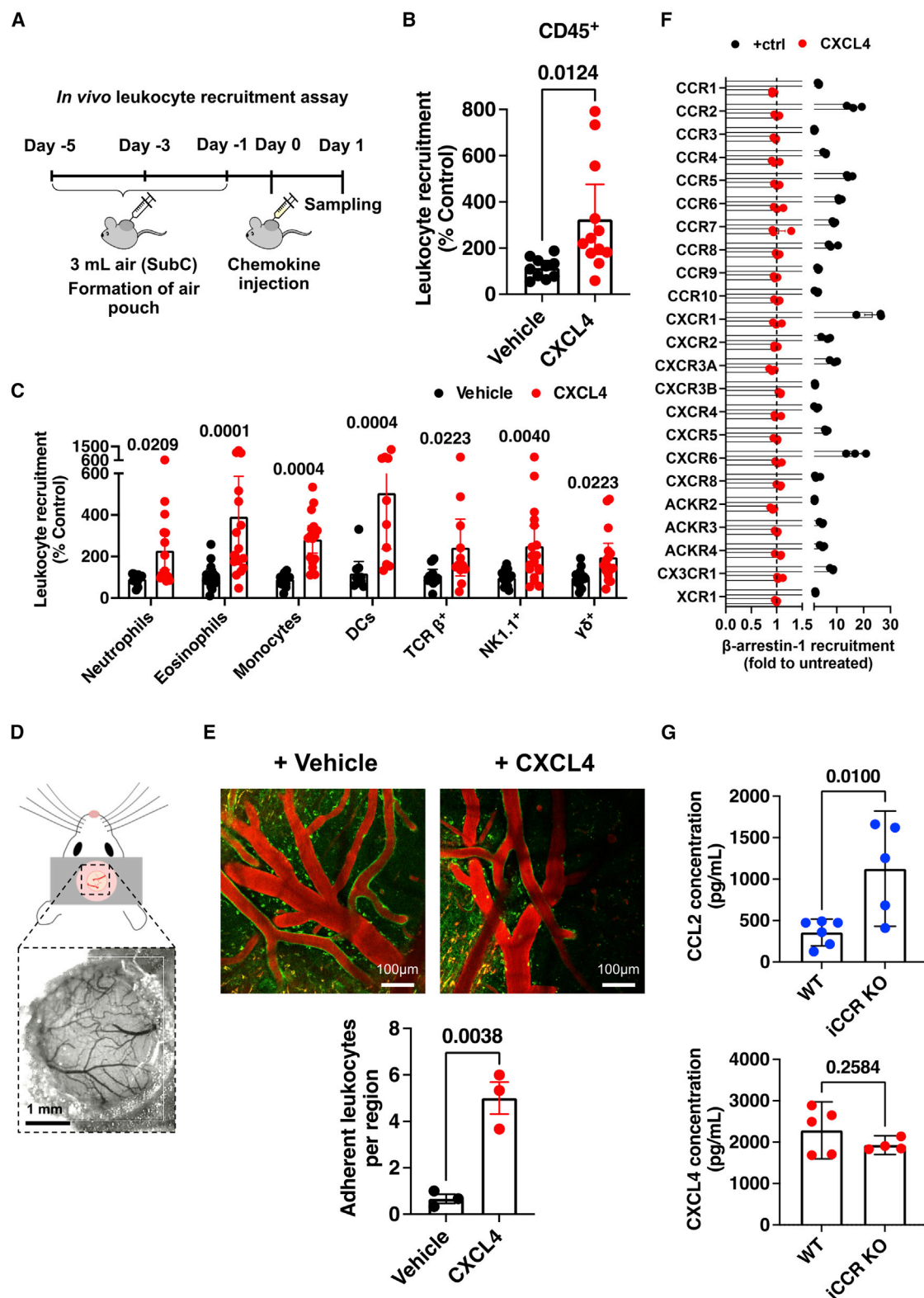


Figure 1. CXCL4 recruits a wide range of different leukocytes *in vivo*

(A) Schematic of the *in vivo* leukocyte recruitment assay.

(B) CD45⁺ cell counts 24 h after CXCL4 injection.

(legend continued on next page)

However, this does not hold for CXCL4 (Platelet Factor 4, PF4).⁵ CXCL4 is produced by platelets and macrophages, regulates leukocyte recruitment, and is implicated in a range of diseases.⁶ It does not have a clearly defined receptor that explains its ability to recruit leukocytes, limiting drug discovery efforts.⁵

Extracellular matrix proteoglycans are present on most cells and are particularly prominent on the vascular endothelium lining blood vessels where they form the endothelial glycocalyx.⁷ This barrier on the apical side of the endothelium regulates vascular permeability, interactions between circulating leukocytes and the vascular endothelium and is thus a central component of leukocyte recruitment into tissues.⁸ Chemokine interactions with key glycocalyx components, glycosaminoglycan (GAG) sugar side chains on proteoglycans, are critical for chemokine function *in vivo* but do not directly affect chemokine:receptor interactions.⁹

The functional role of chemokine:GAG interactions is proposed to be localization of chemokines to the endothelial surface to ensure that chemokine:receptor interactions occur at a specific site.¹⁰ This does not explain why chemokines have a wide range of GAG-binding affinities, where some, like CXCL4, have very high-affinity GAG interactions and also cross-link GAG chains.^{10–12} Thus, the biological consequences of CXCL4:GAG interactions have yet to be understood. GAGs and their larger proteoglycan structures can directly mediate cell signaling in other contexts,¹³ but the relevance of this to chemokine function has been largely overlooked. Currently we do not understand the mechanisms that drive CXCL4-mediated leukocyte recruitment and associated disease.⁶ Interactions with extracellular matrix GAGs may provide an “atypical” mechanistic explanation for CXCL4 function.

Herein, we demonstrate that CXCL4 promotes leukocyte adhesion to the endothelium, increases endothelial permeability, and recruits a wide range of leukocytes driven by the CXCL4:GAG interaction. This demonstrates a chemokine functioning primarily through GAGs and not via a receptor. We also show that GAG fine structure (sulfation pattern) confers specificity to chemokine:GAG interactions, which illustrates how GAGs can impose selectivity onto the supposedly “redundant” chemokine system.

RESULTS

CXCL4 atypically recruits a broad spectrum of different leukocytes

To understand the mechanism underlying CXCL4-mediated leukocyte recruitment, we used an *in vivo* leukocyte recruitment assay to dissect this process (Figure 1A). This involves injecting

air under the mouse dorsum to create a pouch that can be analyzed for immune cell recruitment in response to chemokine injection. CXCL4 increased the number of CD45⁺ (general leukocyte marker) cells, compared with vehicle controls (Figure 1B). Surprisingly, CXCL4 recruited a wide range of different leukocyte types (Figure 1C, gating strategy in Figure S1A), specifically, neutrophils, eosinophils, monocytes, dendritic cells, and T cells (TCR beta⁺, NK1.1⁺, and gamma delta⁺). In contrast, CCL2, which serves as a comparator for CXCL4 given its well established classical recruitment mechanism via a leukocyte chemokine receptor (CCR2),² recruited only monocytes (Figure S1B). Furthermore, using a transcranial window imaging approach (Figure 1D), we also determined that CXCL4 increases adhesion of leukocytes to the vascular endothelium (Figure 1E).

Recruitment of human monocytes by CXCL4 has been reported to be mediated by CCR1.¹⁴ However, CXCL4-mediated recruitment of CD45⁺ cells or monocytes remained unchanged in the CCR1 knockout (KO) mouse compared with wild-type controls (Figure S2A). CXCR3 has also been associated with CXCL4 function, albeit at high concentrations.⁶ Analysis of cells recruited by CXCL4 into the air pouch demonstrated that only T cells expressed CXCR3 (Figure S2B). Furthermore, the Immgen database demonstrates that of the leukocytes recruited by CXCL4, only T cells express CXCR3.¹⁵ Thus, our data suggest that there was no chemokine receptor expression pattern across the leukocyte types recruited by CXCL4 that would explain their recruitment. Further analysis indicates that CXCR3 is also not present on endothelial cells sourced from the air pouch lining (Figure S2C).

In order to comprehensively determine which chemokine receptors may be responsible for CXCL4 function, we examined the ability of CXCL4 to activate all known human chemokine and atypical chemokine receptors in a global screen using β -arrestin-1 recruitment as a functional readout.¹⁶ The resulting data indicate that CXCL4 does not activate any known chemokine receptor (Figure 1F). Therefore, the mechanism underlying widespread CXCL4-mediated recruitment remained unclear.

We next sought to determine whether leukocytes can directly bind and use CXCL4 to facilitate their migration. We exploited previous observations that when chemokine receptors are knocked out, their ligands are increased in the serum due to reduced uptake and degradation.¹⁷ For example, iCCR (CCR1, 2, 3, and 5 combined) KO mice fail to recruit monocytes to the carrageenan-inflamed air pouch.¹⁷ As a result of the ablated monocyte migration, the chemoattractant CCL2 is no longer being used (bound and degraded) during monocyte egress from the blood into the inflamed air pouch of iCCR KO mice, and it is found at elevated levels in the serum (Figure 1G). In contrast, CXCL4 levels remained unchanged under the same conditions.

(C) Quantification of different leukocytes recruited by CXCL4.

(D) Schematic of cranial window implantation for *in vivo* leukocyte adhesion analysis.

(E) *In vivo* analysis of leukocyte (labeled green) adhesion to the walls of the vasculature (labeled red) following injection of CXCL4 or vehicle control.

(F) Agonist activity of CXCL4 (100 nM) toward 19 classical and three atypical chemokine receptors evaluated in a β -arrestin-1 recruitment assay. For all receptors, one known agonist chemokine (100 nM) listed in the IUPHAR repository of chemokine receptor ligands was added as positive control.

(G) CCL2 and CXCL4 quantification in the serum of wild-type or iCCR (CCR1, 2, 3, and 5) KO inflamed mice.

All plots are mean with 95% confidence intervals and represent at least two separate experiments; data have been pooled, and each dot in (B), (C), (E), and (G) represents an individual mouse, and each dot in (F) represents the mean of an individual experiment. (B) and (C) are normalized to vehicle controls. Results in (F) are expressed as fold change relative to untreated controls and are presented as mean of three independent experiments. Individual p values are shown, with (B), (E), and (G) analyzed using an unpaired t test and (C) analyzed using a one-way ANOVA with a post-hoc Sidak analysis of log-transformed data.

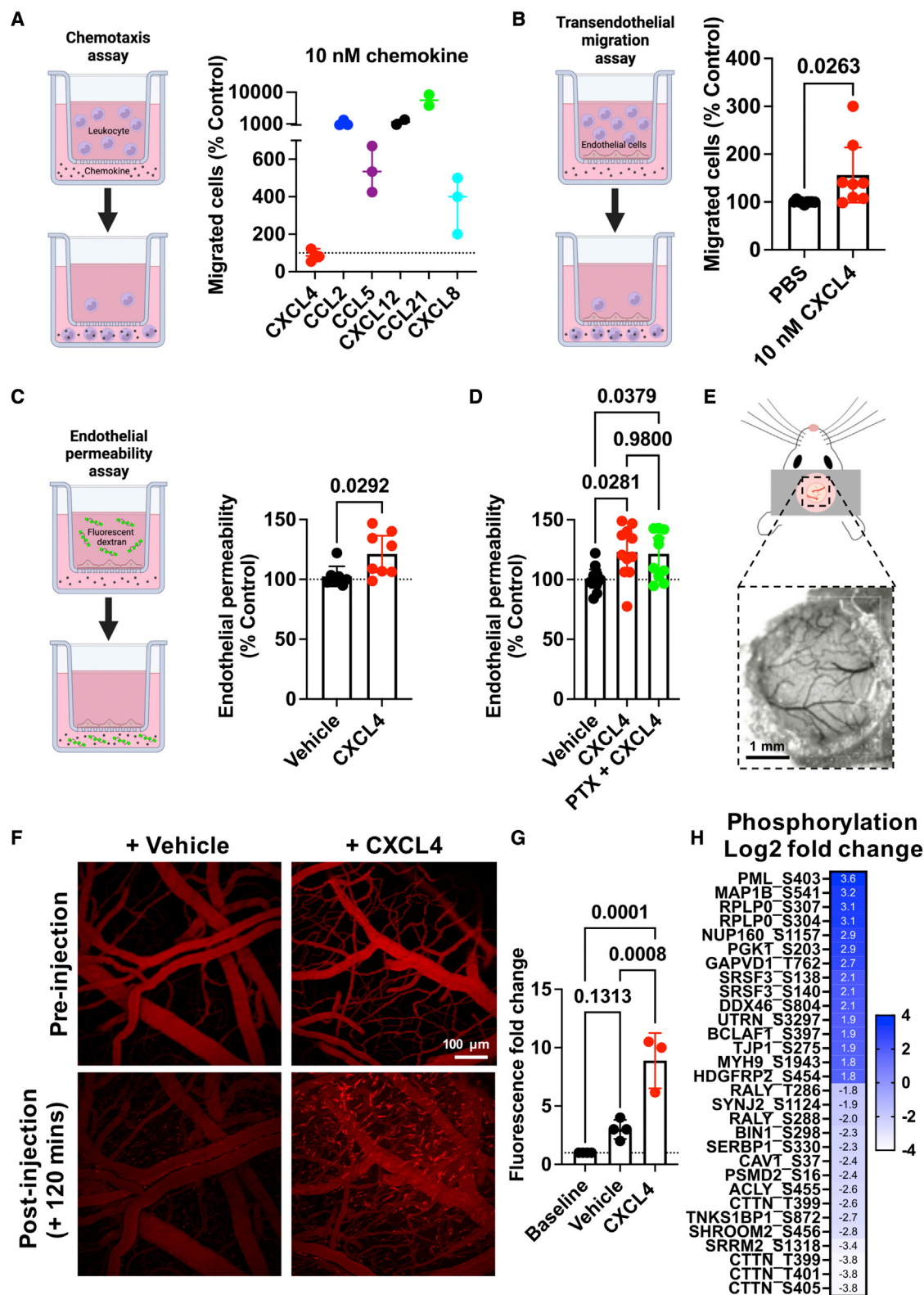


Figure 2. CXCL4 increases endothelial permeability in a receptor-independent manner and mediates endothelial intracellular signaling
(A) Chemokine-mediated chemotaxis of relevant receptor-expressing cells: CXCL4, CCL2, or CCL5 (monocytes), CXCL12 (CXCR4⁺ Jurkat cells), CCL21 (CCR7⁺ L1.2 cells), and CXCL8 (CXCR2⁺ neutrophils).

(legend continued on next page)

This suggests that monocytes do not directly utilize CXCL4 during trafficking to the inflamed air pouch, even though CXCL4 facilitated their recruitment (Figure 1C).

Together these data suggest that CXCL4-mediated recruitment is not driven by binding to chemokine receptors on the leukocyte surface. We hypothesized that CXCL4 would have direct effects on the vascular endothelium regulating processes underlying immune cell recruitment.

CXCL4 increases endothelial permeability

Given that our data suggest a direct effect of CXCL4 on endothelial cells, we hypothesized that CXCL4 may regulate endothelial, and thus vascular, permeability. In a Transwell cell migration system (Figure 2A), purified human monocytes produced a classic chemotaxis response to CCL2 (Figure S3A). Similarly, CCL5 (monocytes), CXCL12 (Jurkats), CCL21 (L1.2 cells), and CXCL8 (neutrophils) mediated recruitment of their concomitant receptor-expressing cells at 10 nM concentrations (Figure 2A). In contrast, no increase in monocyte recruitment was produced by CXCL4, even at a relatively high concentration of 10 nM, and despite the ability of CXCL4 to mediate monocyte recruitment *in vivo* (Figure 1).

To better replicate the *in vivo* environment, we added an endothelial cell monolayer to the upper Transwell chamber to analyze transendothelial migration (Figure 2B). Addition of CXCL4 produced a significant increase in movement of monocytes to the lower Transwell chamber, demonstrating that CXCL4 acts on the endothelium not the leukocyte.

Next, we showed that CXCL4 produced a significant increase in endothelial permeability; CCL5 had a similar effect, whereas CCL2 and CXCL8 did not (Figures 2C, S3B, and S3C). To determine whether this effect of CXCL4 was mediated by a classical chemokine receptor (G α i-coupled), we added pertussis toxin (PTx), but we observed no effect at concentrations where PTx inhibits GPCR-mediated signaling (Figure 2D).¹⁸ Crucially, CXCL4 increased vascular permeability *in vivo* as shown by leakage of fluorescent dextran from the brain meningeal vasculature after CXCL4 administration (Figures 2E–2G). These findings suggest that CXCL4 plays a role in regulating the permeability of the vascular endothelium.

CXCL4 regulates protein phosphorylation within endothelial cells

In order to determine how CXCL4 may facilitate leukocyte recruitment, we firstly analyzed the air pouch for changes in the concentration of a range of different cytokines and chemokines that play a key role in this process following CXCL4 injection (Figure S3D). No changes, relative to vehicle controls, were

observed for any of the proteins analyzed, suggesting no displacement/release of GAG-bound molecules by CXCL4.

To test whether CXCL4 has a direct effect on endothelial cells to facilitate leukocyte trafficking, we utilized phosphorylation-enriched mass spectrometry (Figure 2H). CXCL4 had limited effects on endothelial protein levels after 2 h of stimulation (Figure S3E) but had widespread effects on the phosphorylation state of proteins, relative to vehicle controls, indicative of endothelial signaling responses to CXCL4 (Figure 2H). WebGestalt analysis indicated a number of pathways that were dysregulated following CXCL4 stimulation of endothelial cells.¹⁹ The phosphorylation state of proteins associated with cell adhesion molecule binding was changed following CXCL4 endothelial stimulation. Specifically, PML, GAPVD1, MYH9, and UTRN phosphorylation was increased and CTTN, SERBP1, and TNKS1BP1 phosphorylation was decreased by CXCL4. Dysregulated phosphorylation of proteins associated with adherens junctions and vascular permeability was also observed. Specifically increased phosphorylation of TJP1, MYH9, and RPLP0 and decreased phosphorylation of CAV1, CTTN, SHROOM2, and TNKS1BP1 was detected in response to CXCL4 signaling (Figure 2H). These data suggest that CXCL4 can signal to endothelial cells regulating phosphorylation of proteins that mediate leukocyte adhesion to endothelial cells and also endothelial permeability, supporting our *in vivo* and *in vitro* findings of CXCL4-mediated leukocyte recruitment, adhesion, and vascular permeability (Figures 1 and 2).

CXCL4 function is directly mediated via its interaction with the endothelial GAG heparan sulfate (HS)

Given the independence of CXCL4-mediated leukocyte recruitment from a classic chemokine receptor (Figures 1 and 2), its high affinity for GAG side chains of proteoglycans, and the presence of these proteoglycans within the glycocalyx on the blood-exposed surface of the endothelium (Figure 3A),⁸ we hypothesized that CXCL4 interaction with GAGs on these proteoglycans would directly facilitate leukocyte recruitment. To determine this, we added chemokine to Chinese hamster ovary (CHO) cells with and without genetic KO of the key GAG synthesis gene (*B4galT7*).²⁰ CXCL4 accumulation was GAG dependent and much higher than with CCL2 (Figures 3B and S4A), despite their similar molecular weight and oligomerization propensity.^{11,21} HS is thought to be the dominant GAG within the endothelial glycocalyx.⁸ We, therefore, used a biophysical bio-layer interferometry (BLI) approach (Figure 3C) and determined that the binding of CXCL4 to an HS GAG surrogate, heparin octasaccharide (dp8), was much higher than for CCL2 (Figures 3D and S4B). This difference in cellular accumulation and GAG binding

(B) Transendothelial migration of human monocytes toward CXCL4.

(C and D) (C) Transwell endothelial permeability in the absence and presence of CXCL4; (D) CXCL4 alone or in combination with pertussis toxin.

(E) Schematic of cranial window implantation for *in vivo* vascular permeability analysis.

(F) *In vivo* analysis of leakage of intravenously injected fluorescent dextran from the vasculature into the meninges following intravenous injection of CXCL4 or vehicle control.

(G) Quantification of (F).

(H) Heatmap of the Log2 fold change of indicated protein phosphorylation sites from endothelial cells stimulated with CXCL4, relative to vehicle controls.

All plots are mean with 95% confidence intervals and represent at least two separate experiments where data have been pooled. Each dot in (A)–(D) represents an individual Transwell, and each dot in (G) represents an individual mouse. Data in (A)–(D) and (G) are normalized to vehicle controls. Individual p values are shown, with (C) analyzed using an unpaired t test and (D) and (G) using a one-way ANOVA with a post-hoc Tukey analysis.

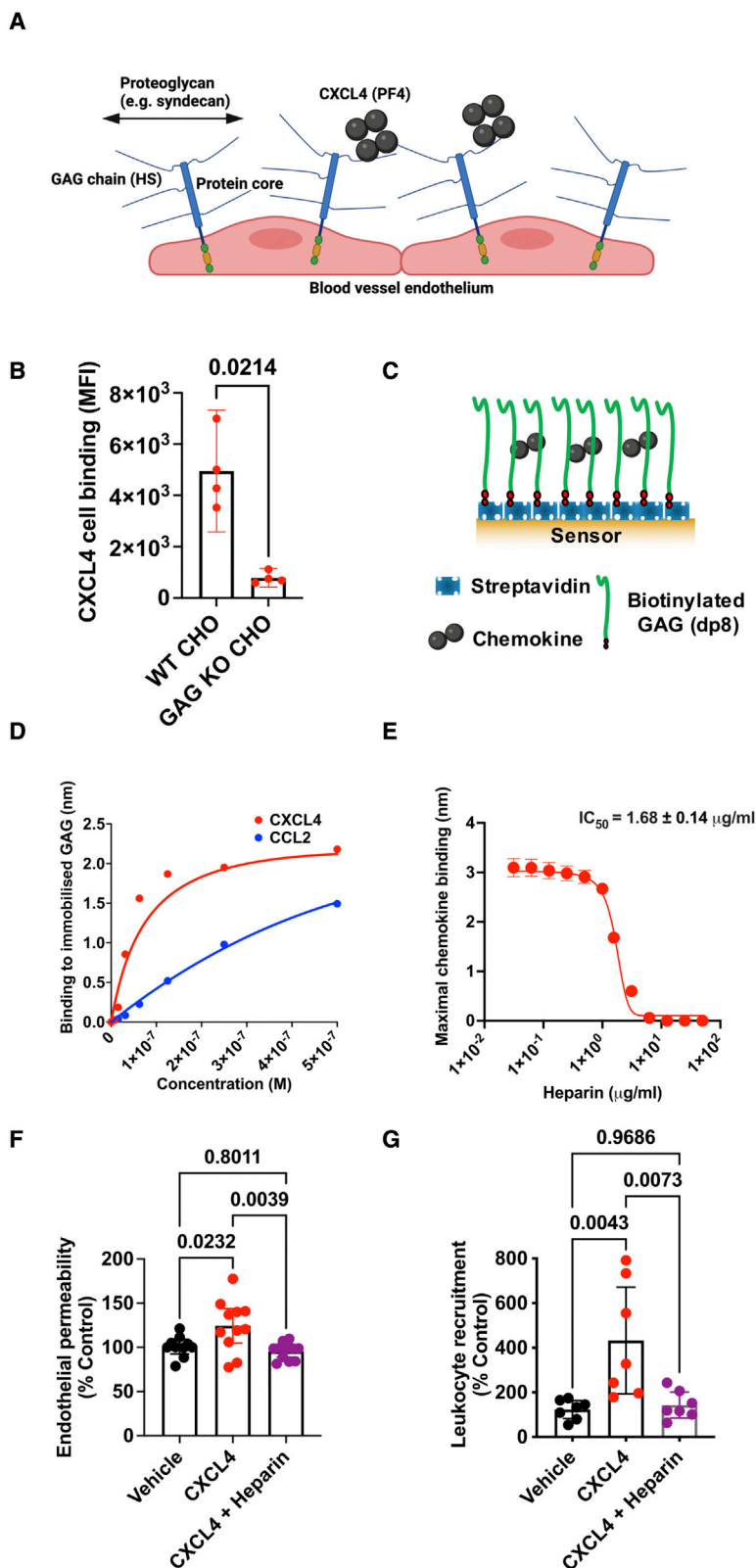


Figure 3. CXCL4 function is mediated by its interaction with glycosaminoglycans (GAGs)

(A) Schematic of proteoglycans forming the luminal endothelial glycocalyx.

(B) CXCL4 binding to CHO cells with and without surface GAGs.

(C) Schematic of the biophysical BLI GAG-binding assay.

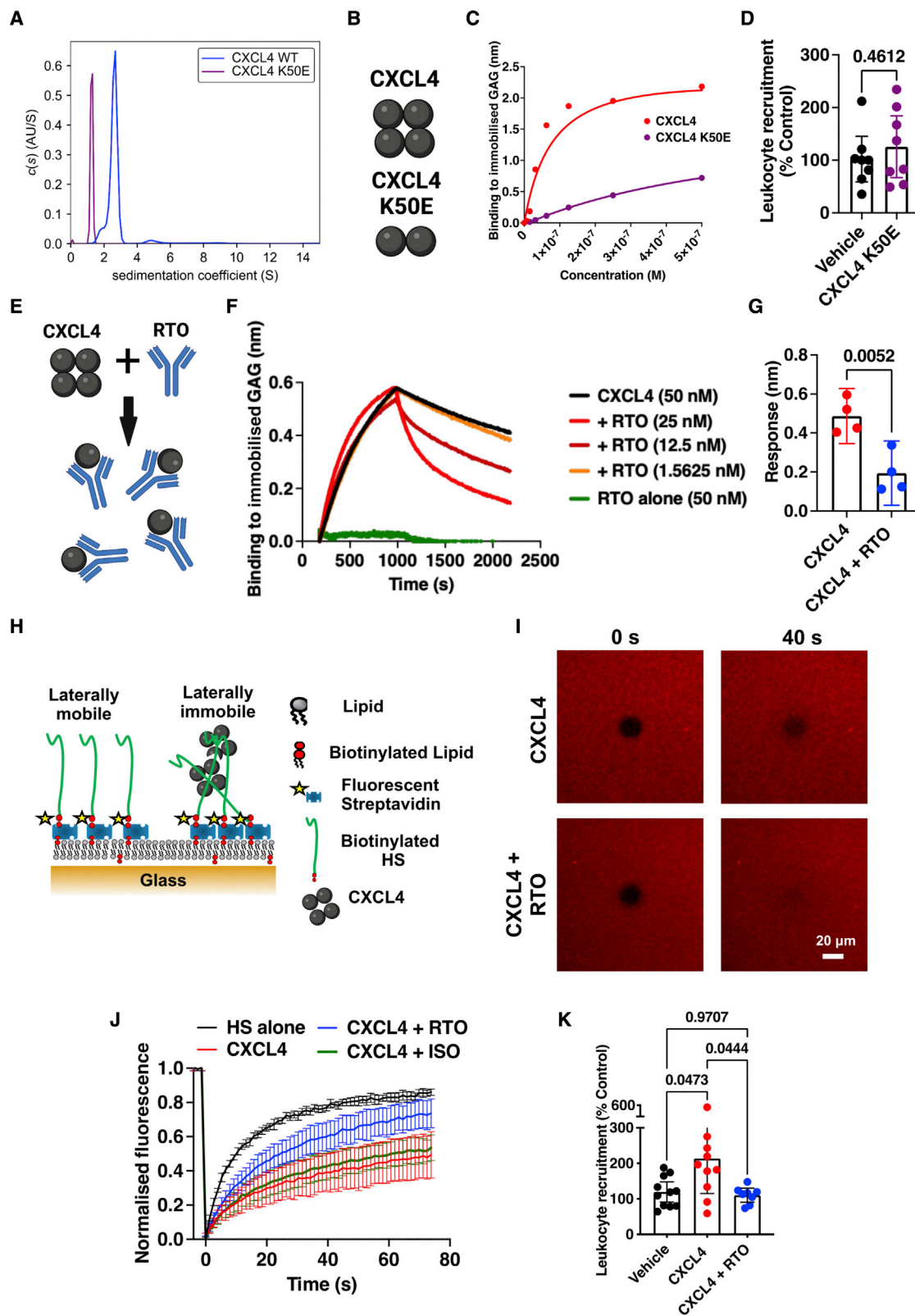
(D) Maximum signal of chemokine:dp8 binding at different chemokine concentrations in the BLI assay.

(E) Maximum CXCL4 (100 nM) binding to dp8 signal following pre-incubation with increasing concentrations of heparin.

(F) Endothelial permeability (Transwell) assay with and without CXCL4 and exogenous heparin.

(G) *In vivo* leukocyte (CD45⁺) recruitment (air pouch) to CXCL4.

(B), (F), and (G) are mean with 95% confidence intervals and represent at least two separate experiments where data have been pooled. (D) and (E) are representative of at least two separate experiments. Each dot in (B) and (F) represents a technical replicate, and each dot in (G) represents an individual mouse. (F) and (G) are normalized to vehicle controls. Individual p values are shown, with (B) analyzed using an unpaired t test and (F) and (G) analyzed using a one-way ANOVA with a post-hoc Tukey analysis.



(legend on next page)

indicated that CXCL4 function could be directly dependent on its GAG interaction, unlike CCL2.

To test this hypothesis, we firstly showed that CXCL4 binding to GAG was inhibited by pre-incubation with exogenous purified GAG (heparin) in the BLI assay (Figures 3E and S4C) and on the cell surface (Figure S4D), demonstrating specificity of the interaction. Subsequently we determined that pre-incubation with heparin reversed CXCL4-mediated increases in endothelial permeability *in vitro* (Figure 3F) and ablated CXCL4-mediated leukocyte recruitment *in vivo* (Figure 3G). In contrast, CCL2-mediated recruitment of monocytes to the air pouch was only reduced by a third following pre-incubation with exogenous heparin (Figure S4E).

These experiments demonstrate that the interaction of CXCL4 with GAGs is responsible for CXCL4-mediated recruitment, likely by modulating endothelial cell functions that regulate leukocyte recruitment. In contrast, the CCL2:GAG interaction has an important but less direct role in facilitating CCL2-mediated leukocyte recruitment, presumably via localization within the endothelial glycocalyx to facilitate interaction with leukocyte receptors.

CXCL4 oligomerization mediates GAG binding and leukocyte recruitment

We have previously shown that high-affinity chemokine:GAG interactions are mediated by chemokine oligomerization and cross-linking of GAG chains.^{11,12} Therefore, we next tested the hypothesis that CXCL4 oligomerization is a critical driver of leukocyte recruitment via cross-linking of GAGs (Figure 4).

To analyze chemokine oligomerization, we used sedimentation velocity analytical ultra-centrifugation (SV-AUC). SV-AUC showed that CXCL4 has a sedimentation coefficient of 2.7S, with an estimated frictional ratio of 1.33 and a mass estimate using the Svedberg equation of 28.9 kDa, consistent with its formation of tetramers in solution (Figure 4A).^{22,23} A mutant of CXCL4 has previously been generated (K50E) to inhibit oligomerization of CXCL4.²⁴ The CXCL4 K50E mutant had a much lower sedimentation coefficient of 1.26S, suggestive of either a compact monomer or an elongated dimer, confirming that the K50E mutation inhibits CXCL4 oligomerization (Figures 4A and 4B).²³ CXCL4 K50E also exhibited a large reduction in binding to GAG in the BLI assay, compared with WT CXCL4 (Figure 4C), and did not recruit leukocytes (CD45⁺) *in vivo* (Figure 4D). We

therefore hypothesized that inhibition of CXCL4 oligomerization is a potential avenue to target CXCL4-driven leukocyte recruitment via inhibition of GAG binding.

Incubation of CXCL4 with an anti-CXCL4 antibody (clone: RTO) that binds monomeric CXCL4 and inhibits CXCL4 oligomerization (Figure 4E)²⁵ produced a concentration-dependent inhibition of the CXCL4:GAG interaction in BLI (Figures 4F, 4G, and S5A), further demonstrating specificity of the CXCL4:GAG interaction. The inhibitory effect of RTO produced an increased off-rate of CXCL4 from GAG, re-creating that observed for CXCL4 K50E (Figure S5B). RTO-mediated inhibition was specific to CXCL4 since it did not affect CCL2 binding to GAG (Figure S5C) or bind to the GAG directly (Figure 4F).

Since the RTO antibody affected the off-rate of the CXCL4:GAG interaction (Figure 4F), we next sought to determine whether RTO could affect CXCL4-mediated cross-linking of GAG chains. To test this, we used a biophysical model of the glycocalyx formed by HS GAG chains anchored on a fluid-supported lipid bilayer, enabling analysis of GAG in-plane mobility and cross-linking using fluorescence recovery after photo bleaching (FRAP) (Figures 4H–4J, S5D, and S5E). CXCL4 inhibited recovery 40 s after bleaching (Figure 4I) and over time (Figure 4J), indicating cross-linking of GAG chains. Pre-incubation of CXCL4 with RTO returned the bleaching recovery close to that seen in the absence of CXCL4 (HS alone) (Figures 4I, 4J, and S5E). These data demonstrate that RTO-mediated inhibition of CXCL4 oligomerization prevents cross-linking of GAGs. Together, the AUC, BLI, and FRAP data demonstrate that chemokine oligomerization and cross-linking of GAGs are related events that drive the interaction with GAGs.

To determine the importance of CXCL4 oligomerization, GAG binding, and cross-linking in leukocyte recruitment, we utilized RTO in our *in vivo* leukocyte recruitment model (Figure 4K). RTO resulted in ablation of CXCL4-mediated CD45⁺ leukocyte recruitment, confirming the importance of oligomerization *in vivo*. This further supports the direct role of GAGs in CXCL4-driven leukocyte recruitment since monomeric chemokines promote chemokine receptor signaling through their GPCRs.^{10,26} Indeed, recent structural studies suggest that chemokine oligomerization and receptor binding are likely to be mutually exclusive due to overlap in binding sites, at least for many receptor:chemokine pairs.⁹

Figure 4. CXCL4 oligomerization drives leukocyte recruitment

- (A) AUC analysis of CXCL4 and the mutant K50E to show sedimentation coefficients (indicative of size).
(B) Schematic of oligomerization states of CXCL4 or the CXCL4 K50E mutant.
(C) BLI analysis of CXCL4 K50E binding to immobilized heparin dp8.
(D) *In vivo* leukocyte (CD45⁺) recruitment (air pouch) to CXCL4 K50E mutant.
(E) Schematic of RTO antibody binding to CXCL4 monomer to inhibit oligomerization.
(F) CXCL4 (50 nM) binding to dp8 was monitored in the absence and presence (at a range of concentrations) of RTO antibody.
(G) Final CXCL4:dp8 signal (after washing) from (F) is plotted with and without pre-incubation with RTO antibody (25 nM).
(H) Schematic of FRAP assay to analyze HS cross-linking.
(I) FRAP analysis of GAG in-plane mobility with CXCL4 alone or in combination with the RTO antibody; images post bleaching at indicated times.
(J) Fluorescence recovery over time.

(K) CXCL4-mediated leukocyte recruitment (CD45⁺) with and without RTO antibody.

(D), (G), (J), and (K) are mean with 95% confidence intervals and represent at least two separate experiments where data have been pooled. Each dot in (D) and (K) represents an individual mouse, and each dot in (G) represents a technical replicate. (A), (C), (F), (I), and (J) are representative of at least two separate experiments. Data in (D) and (K) are normalized to vehicle controls. Individual p values are shown, with (G) analyzed using an unpaired t test and (K) analyzed using a one-way ANOVA with a post-hoc Tukey analysis.

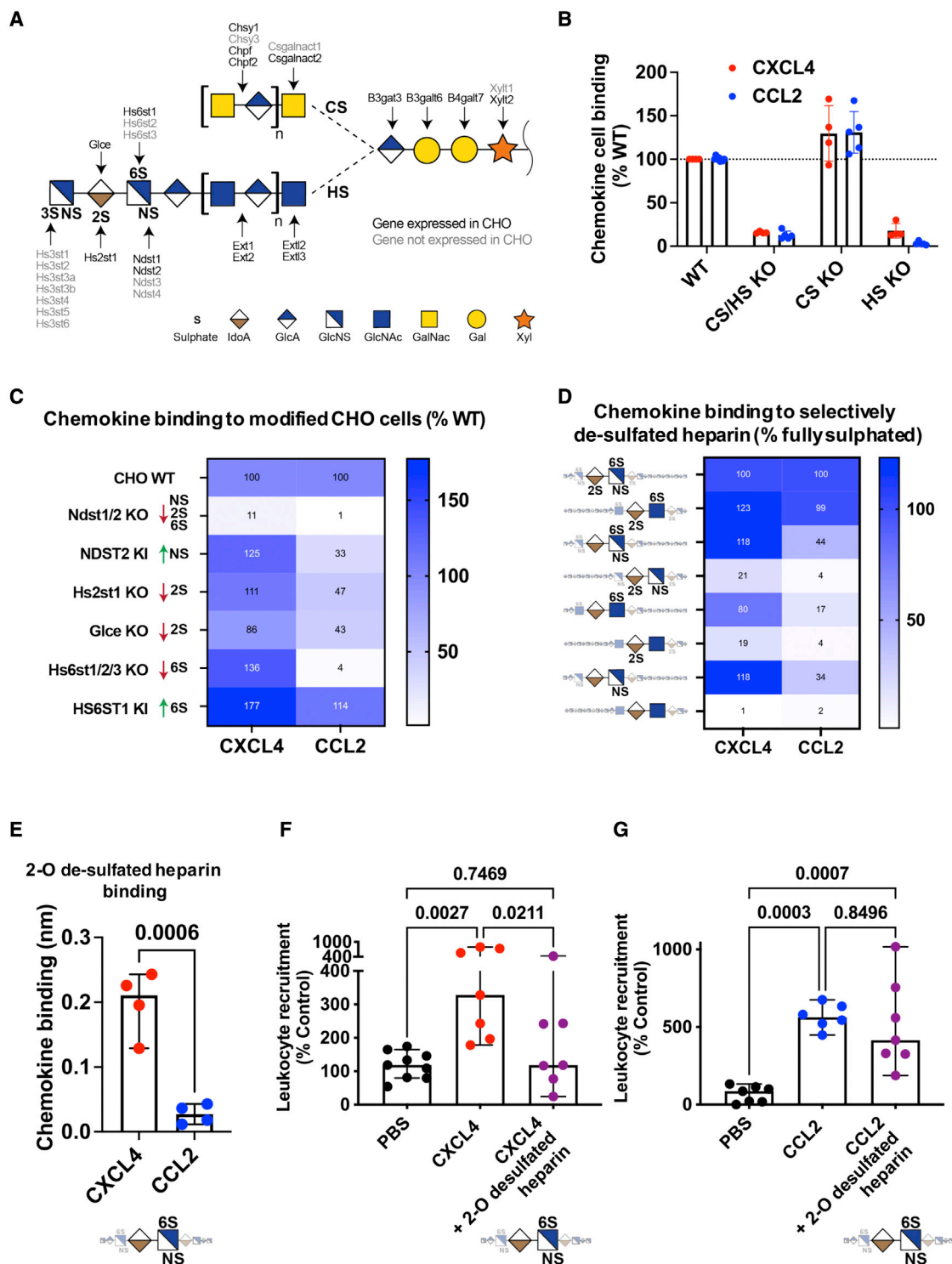


Figure 5. GAG sulfation mediates chemokine interaction selectivity and cellular localization

(A) Overview of enzymes involved in the biosynthesis of HS and CS GAGs. Elongation proceeds from right to left; a tetrasaccharide linker is elongated with either HS or CS disaccharide repeats into long linear polysaccharides. HS is further modified by addition of sulfate groups at the N-, 2-O, 6-O, or 3-O (rarely) positions, and this process is catalyzed by the indicated sulfotransferases.

(B) Chemokine binding to WT CHO cells or CHO cells with no GAGs (KO B4gal7), no CS (KO Csgalnact1/2/Chsy1), or no HS (KO Extl3).

(C) Heatmap analysis of chemokine binding to CHO cells with KO/KI of sulfotransferases acting on HS, where data are normalized to WT cells.

(legend continued on next page)

GAG sulfation confers selectivity on chemokine localization

The above data show that CXCL4 function is driven by its interaction with GAG side chains of proteoglycans on the endothelial cell surface. To further characterize chemokine:GAG interactions, we probed the role of specific GAG sulfation positions (Figure 5A). We hypothesized that changes in GAG sulfation would alter which chemokines are bound and presented on cells.

Earlier experiments (Figure 3B) demonstrated that GAG deficiency in CHO cells resulted in reduced binding to CXCL4. To determine which GAG types are responsible for chemokine binding, we used CHO and human embryonic kidney (HEK) cells with gene KOs to remove, individually or in combination, HS and/or chondroitin sulfate (CS) (Figures 5A and S6).^{20,27} We determined that HS bound to CCL2 and CXCL4 (Figure 5B), while CS GAG only bound to CXCL4 on HEK cells (Figure S6A).

HS GAG chains are 20–200 repeating disaccharides in length and are selectively sulfated on the N-, 2-O, 6-O, and 3-O positions, which mediate chemokine:GAG binding (Figure 5A).²⁸ The enzymes that produce these different sulfation points were systematically knocked out in CHO cells, facilitating analysis of their contribution to chemokine:HS interactions (Figure 5C). KO of NDST1/2, which reduces N-, 2-O, and 6-O sulfation on HS,²⁰ largely ablated CHO cell binding to CXCL4 and CCL2. CXCL4 binding was not reduced in any of the other cell lines. CCL2 binding was reduced in the absence of HS2ST1 (2-O sulfation), GLCE (required for 2-O sulfation), and combined HS6ST1, 2, and 3 (6-O sulfation). Since CHO cells do not make 3-O sulfated HS, the genes that produce it were knocked in.²⁹ 3-O sulfation generally increased CHO cell interaction with CXCL4 and CCL2 (Figure S6B), but HS3ST1 KI significantly enhanced binding to CCL2 but not CXCL4. These data demonstrate that differential sulfation of GAG chains provides considerable selectivity in binding to chemokines and presenting them on the cell surface.

Preferential targeting of CXCL4 function using GAG mimetics

Disruption of chemokine:GAG interactions is a possible avenue for therapeutic intervention in disease.³⁰ Heparin GAG and its derivatives are used as anti-coagulants and are well tolerated in the clinic.³¹ Given the selective interactions mediated by GAG sulfation (Figures 5A–5C), we hypothesized that modified heparin derivatives, with specific sulfation sites removed, may allow preferential inhibition of chemokine-driven leukocyte recruitment.

To determine binding selectivity of differentially sulfated heparins, we used BLI to analyze binding to CXCL4 and CCL2 (Figure 5D). This confirmed higher accumulation of CXCL4,

over CCL2, on fully sulfated (2-O, 6-O, and N- sulfated) heparin (Figure S6C). Again CCL2 was more sensitive to selective removal of specific sulfate groups than CXCL4 (Figure 5D). The CXCL4 K50E mutant demonstrated that oligomerization is important for overall GAG binding and not sulfation specificity (Figures S6C and S6D). These data further demonstrated chemokine:GAG selectivity.³²

This BLI approach identified a heparin form (2-O de-sulfated) with full binding activity for CXCL4 and reduced binding to CCL2 (Figure 5E). Pre-incubation with the 2-O de-sulfated fragment abolished CXCL4- but not CCL2-mediated leukocyte recruitment *in vivo* (Figures 5F and 5G). These data demonstrate sulfation-driven selectivity of chemokine:GAG interactions and suggest that this can be exploited to use GAG mimetics to selectively inhibit certain chemokines.

DISCUSSION

Here, we show that the chemokine CXCL4 mediates recruitment of a range of different leukocytes by binding to the GAG chains of proteoglycans within the endothelial extracellular matrix and not chemokine receptors on leukocytes or the endothelium (Figure 6). We hypothesize that CXCL4 binding and cross-linking of GAG chains on endothelial proteoglycans trigger signaling mechanisms within the endothelial cells that mediate the effects observed here (Figure 6)¹³ and elsewhere.³³ while we present extensive data demonstrating the importance of GAG binding for CXCL4 function, it remains difficult to exclude a role for a non-defined alternative receptor in this system. Future studies will be needed to fully dissect the underlying signaling pathways and integration with known endothelial signaling mechanisms that regulate leukocyte recruitment.

This generalized recruitment of different leukocytes is supported by previous observations that CXCL4 recruits neutrophils and monocytes in response to ischemia reperfusion injury,³⁴ neutrophils to influenza- or *Pseudomonas aeruginosa*-infected lungs,^{35,36} monocytes to atherosclerotic plaques,³⁷ and T cells into the malaria-infected brain.³⁸ Our study may also provide a mechanism explaining previous observations that CXCL4 increases vascular permeability in the brain during cerebral malaria and in the lung following acute injury.^{38,39} CXCL4-mediated increase of vascular permeability and leukocyte adhesion also fits with the role of platelets in facilitating rapid recruitment of leukocytes in response to inflammation.⁴⁰ Thus we propose that CXCL4 production/release is induced during inflammation, enabling subsequent CXCL4-mediated vascular permeability, leukocyte adhesion to the endothelium, and leukocyte recruitment via binding to GAG chains on endothelial proteoglycans.

(D) Heatmap analysis of maximum BLI signal of chemokine (500 nM) binding to differentially de-sulfated heparin fragments; data normalized to binding on fully sulfated heparin.

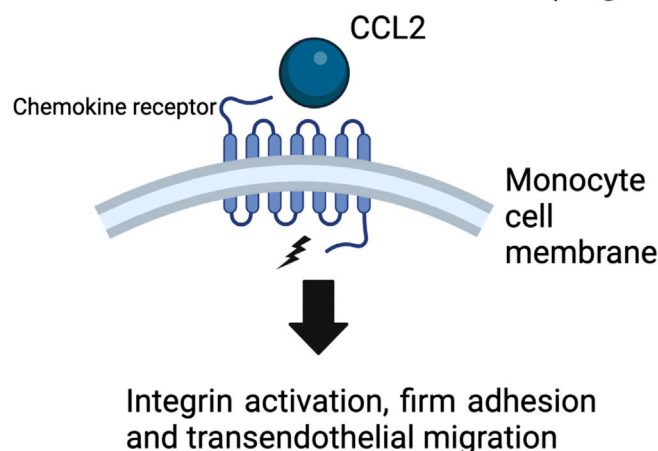
(E) Raw signal of chemokine binding to 2-O de-sulfated heparin in the BLI assay. Chemokine mediated *in vivo* leukocyte recruitment with and without 2-O de-sulfated heparin.

(F) CD45⁺ cell counts (CXCL4) or (G) monocyte (Ly6C⁺) cell counts (CCL2).

(B), (E), (F), and (G) are mean with 95% confidence intervals and represent at least two separate experiments where data have been pooled; each dot in (B) and (E) represents a technical replicate, and each dot in (F) and (G) represents an individual mouse. Data in (F) and (G) are normalized to vehicle controls. Individual p values are shown, with (E) analyzed using an unpaired t test and (F) and (G) analyzed using a one-way ANOVA with a post-hoc Tukey analysis.

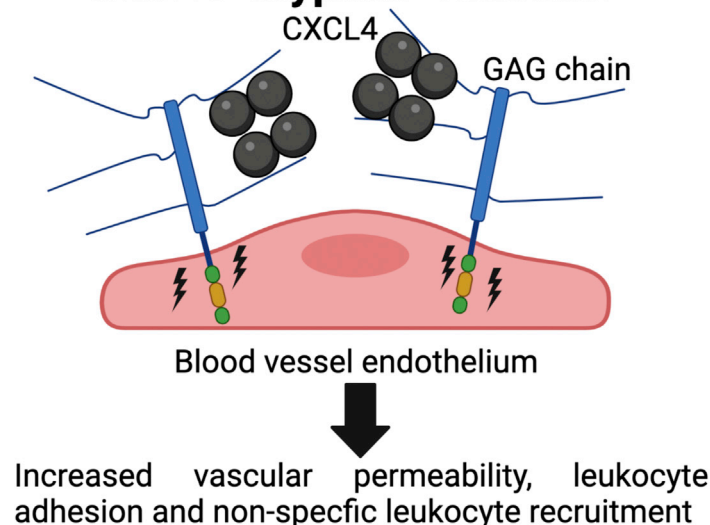
A

Classical chemokine function (e.g. CCL2)



B

CXCL4 "atypical" function



CCL5, which can also cross-link GAGs,¹² increased endothelial permeability *in vitro*, whereas CCL2 and CXCL8, which cannot cross-link GAGs, did not. These data further associate GAG cross-linking with proteoglycan-mediated effects and suggest other chemokines, cytokines, and proteins may function, at least in part, through proteoglycans similarly to CXCL4.

The chemokine system has long been described as redundant, whereby multiple ligands can bind to the same receptor and multiple receptors can bind to the same ligand. However, studies designed to address whether chemokine redundancy exists have found limited evidence for it and in fact suggest specificity.^{17,41–43} Specificity of receptor function is largely explained by coordinated cellular expression.⁴⁴ Contrastingly, there are many examples where chemokine ligands that recruit the same cell types are present at comparable concentrations

Figure 6. CXCL4 binds to endothelial GAG sugars, resulting in an increase of vascular permeability and non-specific leukocyte recruitment

(A) Classical chemokines, e.g., CCL2, facilitate leukocyte recruitment by binding to seven trans-membrane receptors on circulating leukocytes, leading to signaling, integrin activation, and firm adhesion of the leukocyte to the endothelium.

(B) We propose that CXCL4 binds and re-models (as a tetramer) endothelial GAG sugars within the glycocalyx. This produces increased vascular permeability and leukocyte adhesion to endothelial cells possibly via signaling through the proteoglycan, facilitating "non-specific" recruitment of a range of different leukocytes from the vasculature and into inflamed tissues.

in the same context.^{32,45} Our data suggest that chemokine:GAG interactions, coordinated by the differential GAG sulfation patterns found across cells and tissues,⁴⁶ will produce specific localization of chemokines at the cellular and tissue levels.

This study demonstrates ways to target CXCL4 in inflammatory disease by inhibiting oligomerization and GAG binding using GAG mimetics or specific antibodies. Blocking CXCL4-mediated recruitment of a range of leukocytes and increased vascular permeability is particularly promising in acute hyper-inflamed disease, e.g., sepsis.³⁹ Such approaches may also be relevant to targeting CXCL4 in the rare side effects of adenovirus vaccines against SARS-CoV-2 infection.⁴⁷ Future work will be important to address the potential role of CXCL4 as a master regulator of leukocyte recruitment in health and disease.

Limitations of the study

Although we present extensive data demonstrating the importance of GAG binding for CXCL4 function, it remains difficult to

exclude a role for a non-defined alternative receptor in this system. In addition, future studies will be needed to fully dissect the underlying signaling pathways and integration with known endothelial signaling mechanisms that regulate leukocyte recruitment. We also demonstrate that CXCL4 mediates signaling in endothelial cells; however, it remains possible that CXCL4-driven cross-linking of GAG chains within the endothelial glycocalyx may play a more "physical" role in facilitating leukocyte recruitment. Furthermore, we present data demonstrating that specific sulfation of GAG chains mediates specificity of binding to chemokines. It remains to be seen whether this is a key mechanism during tissue specific leukocyte recruitment *in vivo*. It is also possible that the mechanism we describe may be important in the function of other chemokines and related proteins. These issues need to be addressed in future studies.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

- **KEY RESOURCES TABLE**
- **RESOURCE AVAILABILITY**
 - Lead contact
 - Materials availability
 - Data and code availability
- **EXPERIMENTAL MODEL AND SUBJECT DETAILS**
 - Mice
 - Cell lines
- **METHOD DETAILS**
 - Materials
 - *In vivo* leukocyte recruitment
 - Chemokine activity analysis (β -arrestin recruitment)
 - ELISA and Luminex analysis of air pouch fluid
 - *In vitro* chemotaxis, endothelial transmigration and endothelial permeability
 - *In vivo* vascular permeability and leukocyte adhesion
 - Phosphorylation enriched mass spectrometry analysis of CXCL4 stimulated endothelial cells
 - Analysis of chemokine binding to cell surface GAGs
 - Biophysical chemokine:GAG interaction analysis (Bio-layer interferometry)
 - Analytical ultra-centrifugation
 - Fluorescence recovery after photo bleaching (FRAP) of films of in-plane mobile GAGs
 - Immgen
- **QUANTIFICATION AND STATISTICAL ANALYSIS**

SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.celrep.2022.111930>.

ACKNOWLEDGMENTS

The authors would like to acknowledge numerous colleagues for reading and helping in drafting the manuscript. The Bioimaging Facility microscopes used in this study were purchased with grants from BBSRC, Wellcome Trust, and the University of Manchester Strategic Fund. Special thanks to Peter March, Roger Meadows, and Steven Marsden for their help with the microscopy. FRAP analyses was performed on a microscope within the Bioimaging Facilities at University of Leeds, supported by the Wellcome Trust (WT104818MA). Schematics have been drawn in BioRender. D.P.D. was supported by a Sir Henry Dale fellowship jointly funded by the Wellcome Trust and Royal Society (Grant Number 218570/Z/19/Z) and a Wellcome Trust center grant (Grant Number 203128/A/16/Z). R.L.M. was supported by The Carlsberg Foundation CF20-0412. J.T. and R.L.M. were supported by The European Union's Horizon 2020 Research and Innovation Program 899687, HS-SEQ. R.L., M.S. and A.C. were supported by the Luxembourg Institute of Health (LIH), Luxembourg National Research Fund (INTER/FNRS grants 20/15084569), F.R.S.-FNRS-Tél vie (grants 7.4529.19, 7.8504.20 and 7.8508.22). R.L. is a Luxembourg National Research Fund PhD fellow (PRIDE-14254520 "I2TRON"). T.M.H. was supported by NIH grants 1R01AI161880 and 1R01CA254402.

AUTHOR CONTRIBUTIONS

Conceptualization: A.L.G., R.L.M., and D.P.D. Methodology: A.L.G., R.K., A.R.E.R., A.J.L.R., N.P., B.K., C.H., L.M.R., H.L.B., C.L.S., E.A.Y., R.P.R.,

J.E.T., T.M.H., G.J.G., T.A.J., I.S., R.P.R., D.P.D., R.L., M.S., A.C., and I.M. Investigation: A.L.G., R.K., A.R.E.R., A.J.L.R., N.P., B.K., C.H., L.M.R., H.L.B., C.L.S., E.A.Y., R.P.R., T.A.J., I.S., D.P.D., R.L., M.S., A.C., and I.M. Funding acquisition: R.L.M., J.E.T., and D.P.D. Project administration: D.P.D. Supervision: T.M.H., G.J.G., R.P.R., R.L.M., and D.P.D. Writing – original draft: R.L.M., D.P.D.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: May 23, 2022

Revised: November 18, 2022

Accepted: December 14, 2022

REFERENCES

1. Nourshargh, S., and Alon, R. (2014). Leukocyte migration into inflamed tissues. *Immunity* 41, 694–707. <https://doi.org/10.1016/j.immuni.2014.10.008>.
2. Bachelierie, F., Ben-Baruch, A., Burkhardt, A.M., Combadi re, C., Farber, J.M., Graham, G.J., Horuk, R., Sparre-Ulrich, A.H., Locati, M., Luster, A.D., et al. (2014). International Union of Basic and Clinical Pharmacology. [corrected]. LXXXIX. Update on the extended family of chemokine receptors and introducing a new nomenclature for atypical chemokine receptors. *Pharmacol. Rev.* 66, 1–79. <https://doi.org/10.1124/pr.113.007724>.
3. Schall, T.J., and Proudfoot, A.E.I. (2011). Overcoming hurdles in developing successful drugs targeting chemokine receptors. *Nat. Rev. Immunol.* 11, 355–363. <https://doi.org/10.1038/nri2972>.
4. Proudfoot, A.E.I., Bonvin, P., and Power, C.A. (2015). Targeting chemokines: pathogens can, why can't we? *Cytokine* 74, 259–267. <https://doi.org/10.1016/j.cyto.2015.02.011>.
5. von Hundelshausen, P., and Schmitt, M.M.N. (2014). Platelets and their chemokines in atherosclerosis-clinical applications. *Front. Physiol.* 5, 294. <https://doi.org/10.3389/fphys.2014.00294>.
6. Vandercappellen, J., Van Damme, J., and Struyf, S. (2011). The role of the CXC chemokines platelet factor-4 (CXCL4/PF-4) and its variant (CXCL4L1/PF-4var) in inflammation, angiogenesis and cancer. *Cytokine Growth Factor Rev.* 22, 1–18. <https://doi.org/10.1016/j.cytogr.2010.10.011>.
7. Gray, A.L., Pun, N., Ridley, A.J.L., and Dyer, D.P. (2022). Role of extracellular matrix proteoglycans in immune cell recruitment. *Int. J. Exp. Pathol.* 103, 34–43. <https://doi.org/10.1111/iep.12428>.
8. Marki, A., Esko, J.D., Pries, A.R., and Ley, K. (2015). Role of the endothelial surface layer in neutrophil recruitment. *J. Leukoc. Biol.* 98, 503–515. <https://doi.org/10.1189/jlb.3mr0115-011r>.
9. Graham, G.J., Handel, T.M., and Proudfoot, A.E.I. (2019). Leukocyte adhesion: reconceptualizing chemokine presentation by glycosaminoglycans. *Trends Immunol.* 40, 472–481. <https://doi.org/10.1016/j.it.2019.03.009>.
10. Handel, T.M., and Dyer, D.P. (2021). Perspectives on the biological role of chemokine:glycosaminoglycan interactions. *J. Histochem. Cytochem.* 69, 87–91. <https://doi.org/10.1369/0022155420977971>.
11. Dyer, D.P., Salanga, C.L., Volkman, B.F., Kawamura, T., and Handel, T.M. (2016). The dependence of chemokine-glycosaminoglycan interactions on chemokine oligomerization. *Glycobiology* 26, 312–326. <https://doi.org/10.1093/glycob/cwv100>.
12. Dyer, D.P., Migliorini, E., Salanga, C.L., Thakar, D., Handel, T.M., and Richter, R.P. (2017). Differential structural remodelling of heparan sulfate by chemokines: the role of chemokine oligomerization. *Open Biol.* 7, 160286. <https://doi.org/10.1098/rsob.160286>.
13. Lambaerts, K., Wilcox-Adelman, S.A., and Zimmermann, P. (2009). The signaling mechanisms of syndecan heparan sulfate proteoglycans. *Curr. Opin. Cell Biol.* 21, 662–669. <https://doi.org/10.1016/j.ceb.2009.05.002>.
14. Fox, J.M., Kausar, F., Day, A., Osborne, M., Hussain, K., Mueller, A., Lin, J., Tsuchiya, T., Kanegasaki, S., and Pease, J.E. (2018). CXCL4/Platelet

- Factor 4 is an agonist of CCR1 and drives human monocyte migration. *Sci. Rep.* 8, 9466. <https://doi.org/10.1038/s41598-018-27710-9>.
15. Heng, T.S.P., and Painter, M.W.; Immunological Genome Project Consortium (2008). The Immunological Genome Project: networks of gene expression in immune cells. *Nat. Immunol.* 9, 1091–1094. <https://doi.org/10.1038/ni1008-1091>.
 16. Kouzeli, A., Collins, P.J., Metzemaekers, M., Meyrath, M., Szpakowska, M., Artinger, M., Struyf, S., Proost, P., Chevigne, A., Legler, D.F., et al. (2020). CXCL14 preferentially synergizes with homeostatic chemokine receptor systems. *Front. Immunol.* 11, 561404. <https://doi.org/10.3389/fimmu.2020.561404>.
 17. Dyer, D.P., Medina-Ruiz, L., Bartolini, R., Schuette, F., Hughes, C.E., Pallas, K., Vidler, F., Macleod, M.K.L., Kelly, C.J., Lee, K.M., et al. (2019). Chemokine receptor redundancy and specificity are context dependent. *Immunity* 50, 378–389.e5. <https://doi.org/10.1016/j.immuni.2019.01.009>.
 18. Gilliland, C.T., Salanga, C.L., Kawamura, T., Trejo, J., and Handel, T.M. (2013). The chemokine receptor CCR1 is constitutively active, which leads to G protein-independent, β -arrestin-mediated internalization. *J. Biol. Chem.* 288, 32194–32210. <https://doi.org/10.1074/jbc.m113.503797>.
 19. Liao, Y., Wang, J., Jaehnig, E.J., Shi, Z., and Zhang, B. (2019). WebGestalt 2019: gene set analysis toolkit with revamped UIs and APIs. *Nucleic Acids Res.* 47, W199–W205. <https://doi.org/10.1093/nar/gkz401>.
 20. Chen, Y.-H., Narimatsu, Y., Clausen, T.M., Gomes, C., Karlsson, R., Steentoft, C., Sliid, C.B., Gustavsson, T., Salanti, A., Persson, A., et al. (2018). The GAGome: a cell-based library of displayed glycosaminoglycans. *Nat. Methods* 15, 881–888. <https://doi.org/10.1038/s41592-018-0086-z>.
 21. Salanga, C.L., Dyer, D.P., Kiselar, J.G., Gupta, S., Chance, M.R., and Handel, T.M. (2014). Multiple glycosaminoglycan-binding epitopes of monocyte chemoattractant protein-3/CCL7 enable it to function as a non-oligomerizing chemokine. *J. Biol. Chem.* 289, 14896–14912. <https://doi.org/10.1074/jbc.m114.547737>.
 22. Zhang, X., Chen, L., Bancroft, D.P., Lai, C.K., and Maione, T.E. (1994). Crystal structure of recombinant human platelet factor 4. *Biochemistry* 33, 8361–8366. <https://doi.org/10.1021/bi00193a025>.
 23. Rauova, L., Zhai, L., Kowalska, M.A., Arepally, G.M., Cines, D.B., and Poncz, M. (2006). Role of platelet surface PF4 antigenic complexes in heparin-induced thrombocytopenia pathogenesis: diagnostic and therapeutic implications. *Blood* 107, 2346–2353. <https://doi.org/10.1182/blood-2005-08-3122>.
 24. Rauova, L., Poncz, M., McKenzie, S.E., Reilly, M.P., Arepally, G., Weisel, J.W., Nagaswami, C., Cines, D.B., and Sachais, B.S. (2005). Ultralarge complexes of PF4 and heparin are central to the pathogenesis of heparin-induced thrombocytopenia. *Blood* 105, 131–138. <https://doi.org/10.1182/blood-2004-04-1544>.
 25. Cai, Z., Yarovoi, S.V., Zhu, Z., Rauova, L., Hayes, V., Lebedeva, T., Liu, Q., Poncz, M., Arepally, G., Cines, D.B., et al. (2015). Atomic description of the immune complex involved in heparin-induced thrombocytopenia. *Nat. Commun.* 6, 8277. <https://doi.org/10.1038/ncomms9277>.
 26. Salanga, C.L., and Handel, T.M. (2011). Chemokine oligomerization and interactions with receptors and glycosaminoglycans: the role of structural dynamics in function. *Exp. Cell Res.* 317, 590–601. <https://doi.org/10.1016/j.yexcr.2011.01.004>.
 27. Narimatsu, Y., Joshi, H.J., Nason, R., Van Coillie, J., Karlsson, R., Sun, L., Ye, Z., Chen, Y.-H., Schjoldager, K.T., Steentoft, C., et al. (2019). An atlas of human glycosylation pathways enables display of the human glycome by gene engineered cells. *Mol. Cell* 75, 394–407.e5. <https://doi.org/10.1016/j.molcel.2019.05.017>.
 28. Xu, D., and Esko, J.D. (2014). Demystifying heparan sulfate-protein interactions. *Annu. Rev. Biochem.* 83, 129–157. <https://doi.org/10.1146/annurev-biochem-060713-035314>.
 29. Karlsson, R., Chopra, P., Joshi, A., Yang, Z., Vakhrushev, S.Y., Clausen, T.M., Painter, C.D., Szekeres, G.P., Chen, Y.-H., Sandoval, D.R., et al. (2021). Dissecting structure-function of 3-O-sulfated heparin and engineered heparan sulfates. *Sci. Adv.* 7, eabl6026. <https://doi.org/10.1126/sciadv.abl6026>.
 30. Crijns, H., Vanheule, V., and Proost, P. (2020). Targeting chemokine–glycosaminoglycan interactions to inhibit inflammation. *Front. Immunol.* 11, 483. <https://doi.org/10.3389/fimmu.2020.00483>.
 31. Glass, C.A. (2018). Recombinant heparin-new opportunities. *Front. Med.* 5, 341. <https://doi.org/10.3389/fmed.2018.00341>.
 32. Dyer, D.P. (2020). Understanding the mechanisms that facilitate specificity, not redundancy, of chemokine mediated leukocyte recruitment. *Immunology* 160, 336–344. <https://doi.org/10.1111/imm.13200>.
 33. Affandi, A.J., Carvalho, T., Ottria, A., de Haan, J.J., Brans, M.A.D., Brandt, M.M., Tieland, R.G., Lopes, A.P., Fernández, B.M., Bekker, C.P.J., et al. (2022). CXCL4 drives fibrosis by promoting several key cellular and molecular processes. *Cell Rep.* 38, 110189. <https://doi.org/10.1016/j.celrep.2021.110189>.
 34. Lapchak, P.H., Ioannou, A., Rani, P., Lieberman, L.A., Yoshiya, K., Kannan, L., Dalle Lucca, J.J., Kowalska, M.A., and Tsokos, G.C. (2012). The role of platelet factor 4 in local and remote tissue damage in a mouse model of mesenteric ischemia/reperfusion injury. *PLoS One* 7, e39934. <https://doi.org/10.1371/journal.pone.0039934>.
 35. Guo, L., Feng, K., Wang, Y.C., Mei, J.J., Ning, R.T., Zheng, H.W., Wang, J.J., Worthen, G.S., Wang, X., Song, J., et al. (2017). Critical role of CXCL4 in the lung pathogenesis of influenza (H1N1) respiratory infection. *Mucosal Immunol.* 10, 1529–1541. <https://doi.org/10.1038/mi.2017.1>.
 36. Yue, L., Pang, Z., Li, H., Yang, T., Guo, L., Liu, L., Mei, J., Song, X., Xie, T., Zhang, Y., et al. (2018). CXCL4 contributes to host defense against acute *Pseudomonas aeruginosa* lung infection. *PLoS One* 13, e0205521. <https://doi.org/10.1371/journal.pone.0205521>.
 37. Sachais, B.S., Turrentine, T., Dawicki McKenna, J.M., Rux, A.H., Rader, D., and Kowalska, M.A. (2007). Elimination of platelet factor 4 (PF4) from platelets reduces atherosclerosis in C57Bl/6 and apoE^{-/-} mice. *Thromb. Haemost.* 98, 1108–1113. <https://doi.org/10.1160/th07-04-0271>.
 38. Srivastava, K., Cockburn, I.A., Swaim, A., Thompson, L.E., Tripathi, A., Fletcher, C.A., Shirk, E.M., Sun, H., Kowalska, M.A., Fox-Talbot, K., et al. (2008). Platelet factor 4 mediates inflammation in experimental cerebral malaria. *Cell Host Microbe* 4, 179–187. <https://doi.org/10.1016/j.chom.2008.07.003>.
 39. Bdeir, K., Gollomp, K., Stasiak, M., Mei, J., Papiewska-Pajak, I., Zhao, G., Worthen, G.S., Cines, D.B., Poncz, M., and Kowalska, M.A. (2017). Platelet-specific chemokines contribute to the pathogenesis of acute lung injury. *Am. J. Respir. Cell Mol. Biol.* 56, 261–270. <https://doi.org/10.1165/rcmb.2015-0245oc>.
 40. Ed Rainger, G., Chimen, M., Harrison, M.J., Yates, C.M., Harrison, P., Watson, S.P., Lordkipanidzé, M., and Nash, G.B. (2015). The role of platelets in the recruitment of leukocytes during vascular disease. *Platelets* 26, 507–520. <https://doi.org/10.3109/09537104.2015.1064881>.
 41. Groom, J.R., Richmond, J., Murooka, T.T., Sorensen, E.W., Sung, J.H., Bankert, K., von Andrian, U.H., Moon, J.J., Mempel, T.R., and Luster, A.D. (2012). CXCR3 chemokine receptor-ligand interactions in the lymph node optimize CD4⁺ T helper 1 cell differentiation. *Immunity* 37, 1091–1103. <https://doi.org/10.1016/j.immuni.2012.08.016>.
 42. Girbl, T., Lenn, T., Perez, L., Rolas, L., Barkaway, A., Thiriot, A., Del Fresno, C., Lynam, E., Hub, E., Thelen, M., et al. (2018). Distinct compartmentalization of the chemokines CXCL1 and CXCL2 and the atypical receptor ACKR1 determine discrete stages of neutrophil diapedesis. *Immunity* 49, 1062–1076.e6. <https://doi.org/10.1016/j.immuni.2018.09.018>.
 43. Coombs, C., Georgantzoglou, A., Walker, H.A., Patt, J., Merten, N., Poplimont, H., Busch-Nentwich, E.M., Williams, S., Kotsi, C., Kostenis, E., et al. (2019). Chemokine receptor trafficking coordinates neutrophil clustering and dispersal at wounds in zebrafish. *Nat. Commun.* 10, 5166. <https://doi.org/10.1038/s41467-019-13107-3>.

44. Medina-Ruiz, L., Bartolini, R., Wilson, G.J., Dyer, D.P., Vidler, F., Hughes, C.E., Schuette, F., Love, S., Pingen, M., Hayes, A.J., et al. (2022). Analysis of combinatorial chemokine receptor expression dynamics using multi-receptor reporter mice. *Elife* 11, e72418. <https://doi.org/10.7554/elif.72418>.
45. Ridley, A.J., Ou, Y., Karlsson, R., Pun, N., Birchenough, H.L., Jowitt, T.A., Lawless, C., Miller, R.L., and Dyer, D.P. (2022). Chemokines form complex signals during inflammation and disease that can be decoded by extracellular matrix proteoglycans. Preprint at bioRxiv. <https://doi.org/10.1101/2022.09.20.508420>.
46. Warda, M., Toida, T., Zhang, F., Sun, P., Munoz, E., Xie, J., and Linhardt, R.J. (2006). Isolation and characterization of heparan sulfate from various murine tissues. *Glycoconj. J.* 23, 555–563. <https://doi.org/10.1007/s10719-006-7668-1>.
47. Baker, A.T., Boyd, R.J., Sarkar, D., Teixeira-Crespo, A., Chan, C.K., Bates, E., Waraich, K., Vant, J., Wilson, E., Truong, C.D., et al. (2021). ChAdOx1 interacts with CAR and PF4 with implications for thrombosis with thrombocytopenia syndrome. *Sci. Adv.* 7, eabl8213. <https://doi.org/10.1126/sciadv.abl8213>.
48. Thakar, D., Migliorini, E., Coche-Guerente, L., Sadir, R., Lortat-Jacob, H., Boturyn, D., Renaudet, O., Labbe, P., and Richter, R.P. (2014). A quartz crystal microbalance method to study the terminal functionalization of glycosaminoglycans. *Chem. Commun.* 50, 15148–15151. <https://doi.org/10.1039/c4cc06905f>.
49. Brautigam, C.A. (2015). Calculations and publication-quality illustrations for analytical ultracentrifugation data. *Methods Enzymol.* 562, 109–133. <https://doi.org/10.1016/bs.mie.2015.05.001>.
50. Yates, E.A., Santini, F., Guerrini, M., Naggi, A., Torri, G., and Casu, B. (1996). ¹H and ¹³C NMR spectral assignments of the major sequences of twelve systematically modified heparin derivatives. *Carbohydr. Res.* 294, 15–27. [https://doi.org/10.1016/s0008-6215\(96\)90611-4](https://doi.org/10.1016/s0008-6215(96)90611-4).
51. Dixon, A.S., Schwinn, M.K., Hall, M.P., Zimmerman, K., Otto, P., Lubben, T.H., Butler, B.L., Binkowski, B.F., Machleidt, T., Kirkland, T.A., et al. (2016). NanoLuc complementation reporter optimized for accurate measurement of protein interactions in cells. *ACS Chem. Biol.* 11, 400–408. <https://doi.org/10.1021/acscchembio.5b00753>.
52. D'Uonno, G., Reynders, N., Meyrath, M., Abboud, D., Uchański, T., Laeremans, T., Volkman, B.F., Janji, B., Hanson, J., Szpakowska, M., et al. (2022). The extended N-terminal domain confers atypical chemokine receptor properties to CXCR3-B. *Front. Immunol.* 13, 868579. <https://doi.org/10.3389/fimmu.2022.868579>.
53. Chevnigné, A., Janji, B., Meyrath, M., Reynders, N., D'Uonno, G., Uchański, T., Xiao, M., Berchem, G., Ollert, M., Kwon, Y.-J., et al. (2021). CXCL10 is an agonist of the CC family chemokine scavenger receptor ACKR2/D6. *Cancers* 13, 1054. <https://doi.org/10.3390/cancers13051054>.
54. Goldey, G.J., Roumis, D.K., Glickfeld, L.L., Kerlin, A.M., Reid, R.C., Bonin, V., Schafer, D.P., and Andermann, M.L. (2014). Removable cranial windows for long-term imaging in awake mice. *Nat. Protoc.* 9, 2515–2538. <https://doi.org/10.1038/nprot.2014.165>.
55. Yoon, J.-H., Shin, P., Joo, J., Kim, G.S., Oh, W.-Y., and Jeong, Y. (2022). Increased capillary stalling is associated with endothelial glycocalyx loss in subcortical vascular dementia. *J. Cereb. Blood Flow Metab.* 42, 1383–1397. <https://doi.org/10.1177/0271678x221076568>.
56. Pai, S., Qin, J., Cavanagh, L., Mitchell, A., El-Assaad, F., Jain, R., Combes, V., Hunt, N.H., Grau, G.E.R., and Weninger, W. (2014). Real-time imaging reveals the dynamics of leukocyte behaviour during experimental cerebral malaria pathogenesis. *PLoS Pathog.* 10, e1004236. <https://doi.org/10.1371/journal.ppat.1004236>.
57. Tape, C.J., Worboys, J.D., Sinclair, J., Gourlay, R., Vogt, J., McMahon, K.M., Trost, M., Lauffenburger, D.A., Lamont, D.J., and Jørgensen, C. (2014). Reproducible automated phosphopeptide enrichment using magnetic TiO₂ and Ti-IMAC. *Anal. Chem.* 86, 10296–10302. <https://doi.org/10.1021/ac5025842>.
58. Tyanova, S., Temu, T., and Cox, J. (2016). The MaxQuant computational platform for mass spectrometry-based shotgun proteomics. *Nat. Protoc.* 11, 2301–2319. <https://doi.org/10.1038/nprot.2016.136>.
59. UniProt Consortium; Bateman, A., Martin, M.-J., Orchard, S., Magrane, M., Agivetova, R., Ahmad, S., Alpi, E., Bowler-Barnett, E.H., Britto, R., et al. (2021). UniProt: the universal protein knowledgebase in 2021. *Nucleic Acids Res.* 49, D480–D489. <https://doi.org/10.1093/nar/gkaa1100>.
60. Tyanova, S., Temu, T., Sinitcyn, P., Carlson, A., Hein, M.Y., Geiger, T., Mann, M., and Cox, J. (2016). The Perseus computational platform for comprehensive analysis of (prote)omics data. *Nat. Methods* 13, 731–740. <https://doi.org/10.1038/nmeth.3901>.
61. Schuck, P. (2000). Size-distribution analysis of macromolecules by sedimentation velocity ultracentrifugation and lamm equation modeling. *Biophys. J.* 78, 1606–1619. [https://doi.org/10.1016/s0006-3495\(00\)76713-0](https://doi.org/10.1016/s0006-3495(00)76713-0).
62. Jasnin, M., van Eijck, L., Koza, M.M., Peters, J., Laguri, C., Lortat-Jacob, H., and Zaccari, G. (2010). Dynamics of heparan sulfate explored by neutron scattering. *Phys. Chem. Chem. Phys.* 12, 3360–3362. <https://doi.org/10.1039/b923878f>.
63. Migliorini, E., Thakar, D., Sadir, R., Pleiner, T., Baleux, F., Lortat-Jacob, H., Coche-Guerente, L., and Richter, R.P. (2014). Well-defined biomimetic surfaces to characterize glycosaminoglycan-mediated interactions on the molecular, supramolecular and cellular levels. *Biomaterials* 35, 8903–8915. <https://doi.org/10.1016/j.biomaterials.2014.07.017>.
64. Migliorini, E., Thakar, D., Kühnle, J., Sadir, R., Dyer, D.P., Li, Y., Sun, C., Volkman, B.F., Handel, T.M., Coche-Guerente, L., et al. (2015). Cytokines and growth factors cross-link heparan sulfate. *Open Biol.* 5, 150046. <https://doi.org/10.1098/rsob.150046>.
65. Richter, R.P., Béat, R., and Brisson, A.R. (2006). Formation of solid-supported lipid bilayers: an integrated view. *Langmuir* 22, 3497–3505. <https://doi.org/10.1021/la052687c>.
66. Schindelin, J., Arganda-Carreras, I., Frise, E., Kaynig, V., Longair, M., Pietzsch, T., Preibisch, S., Rueden, C., Saalfeld, S., Schmid, B., et al. (2012). Fiji: an open-source platform for biological-image analysis. *Nat. Methods* 9, 676–682. <https://doi.org/10.1038/nmeth.2019>.

STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Anti-mouse CXCR3 (Clone 173)	BioLegend	Cat# 126522; RRID:AB_2562205
Anti-mouse F4/80 (Clone BM8)	eBioscience	Cat#25-4801-82; RRID: AB_469653
Anti-mouse Ly6C (Clone HK1.4)	BioLegend	Cat#128024; RRID: AB_10643270
Anti-mouse TCR β (Clone H57-597)	BioLegend	Cat#47-5961-82; RRID: AB_1272173
Anti-mouse Ly6G (Clone 1A8)	BioLegend	Cat#127606; RRID: AB_1236494
Anti-mouse CD11b (Clone M1/70)	eBioscience	Cat#47-0112-82; RRID: AB_1603193
Anti-mouse Siglec-F (Clone E50-2440)	BD Biosciences	Cat#552126; RRID: AB_394341
Anti-mouse MHCII (Clone M5/114.15.2)	BioLegend	Cat#107639; RRID: AB_2565894
Anti-mouse NK1.1 (Clone PK136)	BD Biosciences	Cat#108732; RRID: AB_2562218
Anti-mouse TCR $\gamma\delta$ (Clone GL3)	BioLegend	Cat#25-5711-82; RRID: AB_2573464
Anti-mouse CD31 (Clone 390)	eBioscience	Cat#17-0311-82; RRID: AB_657735
Anti-mouse CD31 (Clone 390)	BD Biosciences	Cat#740690; RRID: AB_2740374
Anti CXCL4 (Clone RTO)	ThermoFisher Scientific	Cat#MA5-17639; RRID:AB_2539029
Isotype control for anti-CXCL4	ThermoFisher Scientific	Cat#14-4732-82; RRID: AB_470117
Chemicals, peptides, and recombinant proteins		
Recombinant CXCL4 (PF4)	Protein Foundry	Cat#PFP014
Recombinant CXCL4 K50E	Protein Foundry	NA
Recombinant CCL2	Protein Foundry	Cat#PFP008
Recombinant CCL5	Protein Foundry	Cat#PFP020
Recombinant CXCL12	Protein Foundry	Cat#PFP001
Recombinant CCL21	Protein Foundry	Cat#PFP010
Recombinant CXCL8	Protein Foundry	Cat#PFP036
High Grade Heparin Mw 15,700	Iduron	Cat#HEP-HG 100
Heparin Oligosaccharide dp8	Iduron	Cat#HO08
Differentially de-sulphated heparins	Prof. Turnbull and Dr Yates	NA
Sav Atto 565	Sigma	Cat#74784
DOPC	Avanti Polar Lipids	Cat#850375
DOPE-CAP-B	Avanti Polar Lipids	Cat#870273
Biotinylated hapran sulfate	D. Thakar et al., <i>Chem. Commun. (Camb.)</i> , 50, 15148–15151 (2014). ⁴⁸	NA
What Germ Agglutinin Alex Fluor 488 Conjugate	ThermoFisher Scientific	Cat#W11261
Dextran, Texas Red, 70,000 MW	ThermoFisher Scientific	Cat#D1830
Super-Bond Universal Kit dental cement	Prestige Dental	Cat#K058E
Critical commercial assays		
Mouse CXCL4/PF4 DuoSet ELISA	R&D Systems	Cat#DY595
Mouse CCL2/MCP-1 DuoSet ELISA	R&D Systems	Cat#DY479-05
Bio-Plex Pro Mouse Chemokine Panel, 31-Plex Assay	Bio-Rad	Cat#12002231
Experimental models: Cell lines		
HEK-293T	ATCC	Cat#CRL-3216
Immortalised human umbilical vein endothelial (Ea.Hy 926 cells)	ATCC	Cat#CRL-2922
Jurkat	ATCC	Cat#TIB-152
L1.2 murine pre-B lymphoma cells	Dr Brian Zabel	NA

(Continued on next page)

Continued

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Experimental models: Organisms/strains		
Mouse: CCR1def: B6.129S4- <i>Ccr1</i> ^{tm1Gao}	Gift from Dr. Takanori Kitamura	Taconic 4087; RRID: MGI:3614571 "title="http://www.informatics.jax.org/accession/MGI:3614571">3614571
Mouse: iCCRdef: C57BL/6 (<i>iCCR</i>) ^{KO}	Prof. Graham	NA
C57BL6/JCrl	Charles River	MGI:3775640
Software and algorithms		
FloJo v10.4	BD Biosciences	https://www.flowjo.com/
Prism 9	GraphPad	https://www.graphpad.com/scientific-software/prism/
MaxQuant (v1.6.17.0)	MQ	https://www.maxquant.org/maxquant/
Perseus software (1.6.15.0)	MQ	https://maxquant.net/perseus/
FIJI	ImageJ	https://imagej.net/software/fiji/
Sedfit	Sedfit	https://sedfitsedphat.github.io/
Gussi	C. A. Brautigam, <i>Meth Enzymol.</i> 562, 109–133 (2015). ⁴⁹	C. A. Brautigam, <i>Meth Enzymol.</i> 562, 109–133 (2015).

RESOURCE AVAILABILITY

Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Dr Douglas Dyer, (douglas.dyer@manchester.ac.uk).

Materials availability

This study did not generate new unique reagents.

Data and code availability

- All data reported in this paper will be shared by the [lead contact](#) upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the [lead contact](#) upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Mice

Adult male C57/Bl6 mice (8–12 weeks) were housed in cages of up to four on a 12-h light/dark cycle, with ad libitum access to food and water. All experiments were carried out following ethical approval from The University of Manchester and University of Glasgow and under licence from the UK Home Office (Scientific Procedures Act 1986).

Cell lines

HEK-293T (ATCC), immortalised human umbilical vein endothelial (Ea.Hy 926 cells) (ATCC), Jurkat and L1.2⁺ (kind gift from Brian Zabel) cell lines were grown in Dulbecco's Modified Eagle Medium (DMEM) supplemented with 10% FBS, penicillin (100 Units/mL) and streptomycin (100 µg/mL).

METHOD DETAILS

Materials

All chemokines were purchased from Protein Foundry unless otherwise specified. The dp8 and heparin GAGs were purchased (Iduron) and the de-sulphated heparin fractions were generated as detailed in.⁵⁰

In vivo leukocyte recruitment

The air pouch was created by injecting 3 mL of sterile air subcutaneously under the dorsal skin on 3 occasions 48 h apart as described previously.¹⁷ The indicated quantity of chemokine was re-suspended in 100 µL of PBS (Sigma) and injected into the air pouch. Mice

were culled 24 h later and the air pouch flushed twice with 3 mL of air pouch buffer (PBS containing 1 mM EDTA (Sigma) and 1% (w/v) fetal bovine serum (FBS, Sigma)) to retrieve recruited cells. Retrieved cells were washed in air pouch buffer and stained with fixable cell viability dye (eBioscience, 1:1000 in PBS) for 20 min at 4°C before being washed in flow cytometry buffer and re-suspended in 100 μ L antibody staining cocktail (antibodies and FcR blocking reagent (Miltenyi Biotec) diluted 1:200). Cells were incubated for 30 min at 4°C to stain before being washed in flow cytometry buffer and fixed for 10 min using 100 μ L fixation buffer (BioLegend) at room temperature. CXCR3 staining was performed at 37°C for 15 min prior to surface antibody staining. Cells were then re-suspended in flow cytometry buffer before addition of counting beads (ThermoFisher Scientific) and analyzed using a Fortessa flow cytometer (BD Biosciences).

Flow cytometry data was analyzed to quantify absolute cell counts or as the percentage of live cells before being normalised relative to vehicle controls to facilitate comparison across experiments.

Chemokine activity analysis (β -arrestin recruitment)

Cells and proteins

HEK-293T cell line was purchased from ATCC and were grown in Dulbecco's Modified Eagle Medium (DMEM) supplemented with 10% FBS, penicillin (100 Units/mL) and streptomycin (100 μ g/mL). Chemokines were purchased from PeproTech with the exception of CCL1 and CXCL17 that were purchased from R&D Systems.

β -arrestin recruitment assay

β -arrestin recruitment to chemokine receptors in response to CXCL4 or positive control chemokines (listed in the IUPHAR repository of chemokine receptors) was monitored by NanoLuc complementation assay [NanoBiT, Promega; Madison, USA, as previously described.^{16,51–53} In brief, 8×10^5 HEK293T cells were plated per well of a 6-well dish and 24 h later co-transfected with pNBe vectors encoding chemokine receptors C-terminally tagged with SmBiT and human β -arrestin-1 N-terminally tagged with LgBiT. 24 h after transfection cells were harvested, incubated for 15 min at 37°C with Nano-Glo Live Cell substrate diluted 200-fold and distributed into white 96-well plates (1×10^5 cells per well). CXCL4 or one known agonist chemokine listed in the IUPHAR repository of chemokine receptor ligands was added as positive control to each receptor¹⁶ at a final concentration of 100 nM. Ligand-induced β -arrestin recruitment to chemokine receptors was monitored with a GloMax Discover plate reader (Promega, Madison, USA) for 20 min.

ELISA and Luminex analysis of air pouch fluid

Specific concentrations of CXCL4 and CCL2 in mouse serum samples were measured by enzyme-linked immunosorbent assay (ELISA), using Mouse CXCL4/PF4 and Mouse CCL2/JE/MCP-1 DuoSet ELISA kits (R&D Systems, Minneapolis, MN) as per the manufacturer's instruction. Optical densities were read using a Versamax Microplate Reader (Marshall Scientific, Hampton, NH) at 450 nm wavelength. Air pouch fluid samples were analyzed using a Bio-Plex Pro Mouse Chemokine Panel, 31- Plex Assay (Bio-Rad, UK), and the data was acquired on a Bio-Plex Manager (Software 6.2).

In vitro chemotaxis, endothelial transmigration and endothelial permeability

Chemotaxis

Monocyte and neutrophil purification was performed as described previously,²¹ briefly blood was obtained from the San Diego Blood Banks and peripheral blood mononuclear cells (PBMCs) were separated by using a Ficoll-Paque gradient and centrifugation. Monocytes were then purified from freshly isolated PBMCs using a CD14⁺ selection MACS purification (Miltenyi Biotec) followed by final re-suspension in chemotaxis buffer (DMEM (Sigma) + 10% (v/v) FBS (Sigma)). Neutrophils were removed from the bottom of the Ficoll-Paque gradient and red blood cells were removed using red blood cell lysis buffer followed by washing and final re-suspension in chemotaxis buffer.

Chemokine or vehicle control was added in the bottom well of endothelial transwells (5 μ m pores (Corning)) at the indicated concentration in 600 μ L of chemotaxis buffer. 100 μ L of 2×10^6 (2×10^4 total) cell suspension monocytes (CCL2 and CCL5), Jurkats (CXCL12), CCR7⁺ L1.2 cells (CCL21) and neutrophils (CXCL8) were added to the top well for the indicated chemokine. Transwells were then incubated for 2 h at 37°C/5% CO₂, removed and faux adherent cells from the lower membrane washed into the bottom well before migrated cells were counted using a Guava EasyCyte 8HT flow cytometer (Merck Millipore). Data was then presented as the total number of migrated cells or relative to vehicle controls (% control).

Transendothelial migration and permeability

Transendothelial migration experiments were undertaken as above but prior to the addition of migrating cells, endothelial cells were coated onto the upper side of the transwell insert as follows. Transwells with 5 μ m pores (Corning) were coated with 50 μ L per well of 10 μ g/mL of fibronectin (Sigma) and incubated for 1 h at 37°C/5% CO₂ and washed three times with 50 μ L per well of PBS (Sigma). Immortalised human umbilical vein endothelial cells (Ea.Hy 926 cells) (ATCC) were seeded into each well, 100,000 cells per well in 50 μ L of culture medium (DMEM (Sigma) containing 1% (v/v) penicillin-streptomycin solution (Thermo Fisher Scientific), 10% (v/v) FBS (Sigma) and 1% (v/v) L-Glutamine (Sigma)) and incubated for 2 days at 37°C/5% CO₂. Transwells were then washed with PBS (three times with 50 μ L) and finally chemotaxis buffer prior to endothelial transmigration experiments. In the case of endothelial permeability experiments, 100 μ L of chemotaxis buffer containing 25 μ M 70,000 MW Texas Red dextran (Thermo Fisher Scientific)

was added to the upper side of the endothelial monolayer, following incubation (2 h 37°C/5% CO₂) fluorescence of the bottom well was determined using a FlexStation 3 fluorescent plate reader (Molecular Devices). Data were expressed and plotted relative to vehicle controls.

***In vivo* vascular permeability and leukocyte adhesion**

Animals

Cranial window implantation surgery was performed on male C57BL/6J mice at 20–25 g and multiphoton imaging was conducted two weeks later.

Cranial window implantation

Cranial window implantation was conducted as previously described.⁵⁴ In brief, animals were anesthetized with 2.5% isoflurane in room air and positioned within a stereotactic frame with a heating blanket and temperature probe to ensure maintenance of body temperature at 37.5°C. The cranium was exposed by removal of the scalp on top of both brain hemispheres. A metal head plate (Narishige CP-2, Japan) was mounted using dental cement (Sun Dental, Japan), to allow later fixation within the multiphoton microscope setup. A 3 mm diameter circular piece of bone was then carefully removed using a dental drill and a circular coverslip (Warner Instruments, USA) was secured in place of the removed bone using dental cement.

Multiphoton imaging

Depth of anesthesia was maintained throughout the experiment using 1.5–2% isoflurane in 100% oxygen. Animals were fixed into the setup via the head plate and a heating blanket and temperature probe were used to maintain body temperature at 37.5°C. Images were collected on a Leica SP8 Upright Multiphoton microscope using a Leica 25x/0.95 L HC Fluotar dipping objective. All images equated to a physical size of 620 × 620 μm but image format and thus pixel sizes were varied between different experiments.

Permeability studies

Animals underwent intravenous injection of 50 μL of 10 mg/mL neutral 70,000 MW Texas Red dextran (Thermo Fisher Scientific) in sterile saline following induction of anesthesia. Images were captured at 1024 × 1024 pixel resolution and the z planes were spaced 2 μm apart. Samples were imaged using two-photon excitation from a MaiTai MP laser (Spectra Physics) tuned to 880 nm and signal collected using the Leica external non-descanned hybrid detectors (NDHyD) through a BP624/40 filter for the Texas Red dextran signal and simultaneously onto a second NDHyD through a BP440/20 filter to image the dura via second harmonic generation (SHG) imaging. This approach ensured that the equivalent cranial depth was recorded in each experiment. Baseline images were taken for all regions of interest (ROIs). Animals were then intravenously injected with 5 μg of unlabelled CXCL4 diluted in saline or vehicle control (saline alone). Two or three ROIs were acquired for each animal and three or four animals were used per group. Texas Red channel maximum-intensity projection images of the first 60 μm of tissue including the dura were generated and the fluorescence intensities (mode) for each ROI were calculated. These values were taken for each image at 120 min following CXCL4/vehicle injection and plotted as fold change from the baseline maximum-intensity projection images.

Leukocyte adhesion studies

Intravenous injection of 100 μL of Wheat Germ Agglutinin Alexa Fluor 488 conjugate (Thermo Fisher Scientific) was performed on awake animals 2 h prior to imaging, to label the leukocytes, as previously described.^{55,56} Following induction of anesthesia 5 μg of unlabelled CXCL4 diluted in saline or vehicle control (saline alone) mixed with 50 μL of 10 mg/mL 70,000 MW Texas Red dextran (Thermo Fisher Scientific) was injected intravenously. Two-photon excitation was used with the MaiTai MP laser tuned to 800 nm and imaged simultaneously on to external NDHyD's through BP525/50 (for Alexa Fluor 488) and BP624/40 (for Texas Red) filters. To monitor leukocyte activity in each ROI, a 1-min recording was acquired as a single plane over time (xyt mode) at 256 × 256 pixel resolution at 4 frames per second. Furthermore, images were also acquired at 1024 × 1024 pixel resolution for high resolution representations of each ROI. This imaging regime was repeated 24 h after CXCL4/vehicle injection under terminal anesthesia following an additional injection of 50 μL of 10 mg/mL Texas Red dextran to re-label the bloodstream. Leukocyte adhesion was assessed within the venular system 24 h following CXCL4 or vehicle injection via manual counting. Leukocytes were classed as adhered if they remained in position for the 1 min duration of the xyt recording and data was recorded as average number of adhered leukocytes across the 3 ROIs for each animal. Three ROIs were recorded per animal and 3 animals were used per group.

Phosphorylation enriched mass spectrometry analysis of CXCL4 stimulated endothelial cells

Endothelial stimulation

Wells of a 12-well plate (Sigma-Aldrich) were treated with 1 mL fibronectin bovine plasma (10 μg/mL in PBS, Sigma-Aldrich) and incubated for 1 h at 37°C. Fibronectin was removed and wells were washed with 1 mL PBS. EA.HY 926 endothelial cells were seeded (200,000 cells per well) in culture medium and left to incubate at 37°C for 24 h. Non-adherent cells were removed by washing with 1 mL DPBS twice. Endothelial cells were then stimulated with vehicle control or CXCL4 (7 μg per well) in culture medium for 2 h. Media was then removed from wells, cells washed with PBS and were then lysed with RIPA buffer (Merckmillipore, 1:10 in double distilled H₂O) supplemented with protease inhibitor (cOmplete tablets, Sigma-Aldrich) and phosphatase inhibitor (Sodium orthovanadate, 5 mM, Sigma-Aldrich) and incubated for 30 min at 4°C. Lysates were then frozen at –20°C until needed.

Proteolytic digestion and phosphopeptide enrichment

Protein samples were reduced with 15 mM dithiothreitol for 45 min at 56°C. Samples were then cooled and alkylated with 50 mM iodoacetamide (IAM) (45 min, RT, in the dark). –20°C acetone was added at 4 times volume of sample and incubated at –20°C

for 1 h to precipitate. Following centrifugation (16,000 x g, 20 min, 4°C) the pellet was washed with –20°C acetone and centrifuged again as stated. The pellet was left to dry in a fume hood for 20 min. Pellets were resuspended in 50 mM ammonium bicarbonate supplemented with 0.2% (w/v) Rapigest Surfactant (Waters). Samples were digested with sequencing-grade trypsin (Promega) at an enzyme-to-protein ratio of 1:50 and incubated overnight at 37°C whilst shaking (450 rpm).

Digested samples were acidified with trifluoroacetic acid (TFA) to a final concentration of 1% (v/v), pH 3–5. Samples were incubated at 37°C for 45 min, then centrifuged (20,000 x g, 10 min) to precipitate out Rapigest. Desalting was performed with Oasis HLB Cartridges (Waters) which were wet with 50% acetonitrile (ACN) and equilibrated with 0.1% formic acid (FA). Samples were eluted in binding solution (80% v/v ACN, 5% v/v TFA, 1M glycolic acid). Enrichment of phosphopeptides was performed via incubation with MagReSyn titanium dioxide beads (resynbio). A fraction of digested samples was retained for proteomic analysis. Phosphoproteomic analysis preparation involved washing of beads with 1% ammonium hydroxide and binding solution. Automated phosphopeptide enrichment was performed on the KingFisher Flex robot (ThermoFisherScientific) with a homologous set up to previously described.⁵⁷

LC-MS/MS analysis

Digested samples were made up in 12 µL (phospho samples) and 20 µL (protein samples) of 3% ACN and 0.1% FA. Samples were analyzed via tandem mass spectrometry (LC-MS/MS) using an UltiMate 3000 Rapid Separation LC (RSLC, Dionex Corporation, Sunnyvale, CA) coupled to a QE HF (Thermo Fisher Scientific, Waltham, MA) mass spectrometer. Mobile phase A was 0.1% FA in water and B was 0.1% FA in ACN. The column used was 75 mm x 250 µm i.d. 1.7 mM CSH C18, analytical column (Waters). For proteomic analysis 1 µL aliquot of sample was transferred to 5 µL loop and loaded on the column at a flow of 300 nL/min for 5 min at 5% B. For phospho samples, 3 µL was injected and loaded for 13 min. The loop was taken out of line and flow reduced from 300 nL/min to 200 nL/min (in 30 s). Peptides were separated on a gradient starting from 5% to 18% B (in 63 min 30 s), then from 18% to 27% B (in 8 min) and from 27% B to 60% B (in 1 min). At 85 min flow is increased back to 300 nL/min until 90 min (end of run).

Data was acquired in data directed manner for 90 min in positive mode. Peptides were automatically selected for fragmentation by data dependent analysis of the top 12 peptides with mass/charge (m/z) between 300 and 1750Th, a charge state of 2/3/4 and dynamic exclusion set at 15 s. MS resolution was set at 120,000, ACG target of 3e6 and maximum fill time at 20ms. MS2 resolution was set to 30,000, ACG target of 2e5, maximum fill time of 45 ms, isolation window of 1.3Th and collision energy of 28.

Data processing and analysis

Raw data for both proteome and phosphoproteome were processed together using MaxQuant (v1.6.17.0)⁵⁸ against the human proteome obtained from Uniprot (May 2021).⁵⁹ Fixed modification of carbamidomethylation of cysteine and variable modifications of methionine oxidation, protein N-terminal acetylation were set for both proteome and phosphoproteome data with phosphorylation of serine, tyrosine and threonine set for the phosphoproteome data only. Precursor tolerance was set at 20ppm and 4.5pm for the first and main searches, with MS/MS tolerance set at 20ppm. A false discovery rate (FDR) of 0.01 was set for PSM and protein level, up to two missed cleavages were permitted and “match-between-runs” was selected.

Processed data were analyzed using the Perseus software (1.6.15.0).⁶⁰ Proteome data were filtered for reverse hits, potential contaminants and proteins only identified by modified peptides. Proteins were retained that were present in 50% of samples and ratios between stimulated/un-stimulated were calculated. Phosphorylation site data were filtered for reverse hits, potential contaminants and localisation probability greater than 0.75 (class I). Phosphorylation sites were retained that were present in 50% of samples and ratios between stimulated/un-stimulated were calculated.

Analysis of chemokine binding to cell surface GAGs

This analysis was adapted from a previous approach.²⁰ Briefly, 1×10^5 genetically engineered CHO GS^{−/−} or HEK293 6e cells were harvested from suspension culture and washed in 1 x PBS before being incubated with biotinylated recombinant human CXCL4 or CCL2 (Protein Foundry LLC) at 10 µg/mL in 1 x PBS with 1% FBS for 30 min at 4°C. Cells were washed with 1 x PBS with 1% FBS and incubated with 1:2000 Alexa Fluor 488-streptavidin (S32354, Invitrogen) in 1 x PBS with 1% FBS for 30 min at 4°C. After washing with 1 x PBS with 1% FBS, cells were resuspended in 1 x PBS with 1% FBS and fluorescence intensity was analyzed on an SA3800 spectral cell analyzer (SONY). For the heparin inhibition assays, 10 µg/mL CXCL4 or CCL2 was pre-incubated with varying concentrations of porcine mucosal heparin in 1 x PBS with 1% FBS for 30 min at RT prior to incubation with cells. All experiments were performed a minimum of three times using triplicate samples.

Biophysical chemokine:GAG interaction analysis (Bio-layer interferometry)

Bio-layer interferometry was performed on an Octet Red96 system (Sartorius AG, Goettingen, Germany) using a methodology adapted from.¹¹ Firstly, GAGs were biotinylated at their reducing end using an approach described elsewhere⁴⁸ and immobilised onto High Precision Streptavidin (SAX) biosensors (Sartorius AG, Goettingen, Germany). To immobilise, SAX biosensors were hydrated for 10 min in BLI buffer (10 mM HEPES, 150 mM NaCl, 3 mM EDTA, 0.05% Tween 20, pH 7.4). For the heparin dp8 GAG immobilisation was undertaken in BLI buffer at 0.078 µg/mL to achieve an immobilisation level of approx. 1 nm. For the differentially de-sulphated heparin fractions immobilisation was undertaken at 5 µg/mL in BLI buffer until all surfaces were saturated. Sensors were then washed in regeneration buffer (0.1 M Glycine, 1 M NaCl, 0.1% Tween, pH 9.5) and re-equilibrated in assay buffer. Blank reference or GAG coated sensors were then submerged into wells of black-walled 96 well plates containing 200 µL of BLI buffer containing chemokines

at the indicated concentrations for the indicated time (association) before being transferred to assay buffer containing wells (dissociation) before finally being washed in regeneration buffer. The binding signal was recorded throughout and signal from binding of chemokine to blank (no immobilised GAG) sensors and by GAG immobilised sensors in assay buffer alone was subtracted from that produced by chemokine binding to GAG immobilised sensors. As well as signal over time the maximum signal during the association phase of the interaction was recorded and is plotted. Data were acquired at 5 Hz and analyzed using the Octet HT 10.0 analysis program. For inhibition analysis signal was plotted against inhibitor concentration and fitted for IC_{50} values using non-linear regression in the Prism software package (GraphPad).

Analytical ultra-centrifugation

CXCL4 and CXCL4 K50E were re-suspended in water to a concentration of 1 mg/mL and further diluted 1:10 in PBS. Samples were loaded into 2-sector cells with PBS as a reference and centrifuged at 50,000 rpm in a 4-hole An60Ti rotor monitoring the absorbance at 230 nm until sedimentation was reached. The time-resolved sedimenting boundaries were analyzed using Sedfit.⁶¹ The resulting profiles are shown in Gussi.⁴⁹

Fluorescence recovery after photo bleaching (FRAP) of films of in-plane mobile GAGs

Heparan sulfate (HS) used in FRAP assays was biotinylated at the reducing end (HS-b), as described previously.⁴⁸ HS was sourced from porcine intestinal mucosa, had a molecular weight of 12 kDa, a polydispersity of 1.6 and an average of 1.4 sulfates per disaccharide.⁶² Dioleoylphosphatidylcholine (DOPC) and dioleoylphosphatidylethanolamine-CAP-biotin (DOPE-CAP-b) lipids were purchased from Avanti Polar Lipids (Alabaster, AL, USA). Streptavidin was labeled with Atto 565 fluorophores (Sav Atto 565), and all chemicals for buffers were supplied by Sigma. Working buffer used for all experimental steps consisted of 10 mM HEPES, pH 7.4, and 150 mM NaCl in ultrapure water.

Preparation of model glycocalyxes (films of in-plane mobile GAGs)

Films of one-end anchored HS polysaccharides on supported lipid bilayers were formed as previously described.^{63,64} Briefly, pre-conditioned glass coverslips (24 × 24 mm², type 1.5; Menzel Gläser, Braunschweig, Germany) were attached, using a bicomponent glue (Picodent, Wipperfurth, Germany), to a custom-built Teflon holder, thus forming the bottom of four identical wells with a volume of 50 μ L each. Supported lipid bilayers (SLBs) were formed by the method of vesicle spreading,⁶⁵ through exposure of the glass surface to small unilamellar vesicles (100 μ g/mL for 30 min) made from DOPE-CAP-b in a DOPC background (molar ratio 0.5:99.5). SAV Atto 565 (40 μ g/mL for 20 min) and HS-b (10 μ g/mL for 20 min) were then sequentially applied to the SLBs, to anchor HS via SAV to the biotin-presenting SLB. After each incubation step, the surface was washed with a working buffer to remove excess molecules from the solution phase.

Fluorescence recovery after photo bleaching (FRAP) assays

With HS-b anchored to SAV Atto 565, FRAP of the fluorescently labeled SAV was used as a reporter of GAG in-plane mobility. Fluorescence images were taken with a confocal laser scanning microscope (Zeiss LSM880, Zeiss, Germany) using ZEN software. The objective was a Plan-Apochromat 40×/1.4 Oil DIC M27, and the light source a 561 nm laser. The pinhole size was 198 μ m (5 Airy units). In each FRAP experiment, 3 pre-bleach images of the surface (177.12 × 177.12 μ m²; 256 × 256 pixels) were acquired before a circle in the center of the image (10 μ m radius) was bleached, and 57 images were acquired post bleach to monitor the fluorescence recovery.

To quantify fluorescence recovery, all images were analyzed in Fiji.⁶⁶ Mean fluorescence intensities were extracted from two regions of equal radius (6.92 μ m) in each image in the sequence: the bleach region was located at the image center and the reference region at the periphery. Fluorescence intensities were corrected for background fluorescence and intensity fluctuations, and normalised using the following equation: $I_{\text{norm}}(t) = \frac{I(t) - I_{\text{bg}}}{I_{\text{init}} - I(0)} \cdot \frac{I_{\text{ref}}(t) - I_{\text{bg}}}{I_{\text{ref,init}} - I_{\text{bg}}} - \frac{I(0) - I_{\text{bg}}}{I_{\text{init}} - I(0)}$. Here, I_{bg} is the background intensity (measured in the absence of fluorophores, and constant across the image), I_{init} and $I(t)$ are the intensities in the bleach region prior to bleaching ($t < 0$) and at post-bleach time t ($I(0) = I(t = 0)$), respectively, and $I_{\text{ref,init}}$ and $I_{\text{ref}}(t)$ are the corresponding intensities in the reference region. With this normalisation, the unbleached (or fully recovered) intensity is 1, and the intensity in the bleach region immediately after bleaching is 0. Recovery curves were plotted as normalised intensity against time, and the area under each curve from $t = 0$ to $t = 40$ s was calculated to represent an effective measure of the in-plane mobility of SAV (and the attached HS) on the SLB.

To assess the impact of CXCL4 on HS mobility, the HS film was incubated with CXCL4 alone (250 nM), with a mix of CXCL4 and RTO (at 2:1 M ratio), or with a mix of CXCL4 and a control IgG (at 2:1 M ratio) for 1.5 h; the HS film was then analyzed by FRAP (within 10 min, and with excess proteins in the solution phase). Prior to use, the mixtures of CXCL4 and RTO (or control IgG) were incubated for 1 h to allow complexes to form. Three independent experiments were performed for each condition, and in each experiment three FRAP series were acquired at different positions on the surface. Data shown in Figures 4J and S5E represent the mean $\pm$ 95% confidence intervals across all 9 datasets per condition.

Immgen

The Immgen database was probed using the MyGeneSet data browser.¹⁵

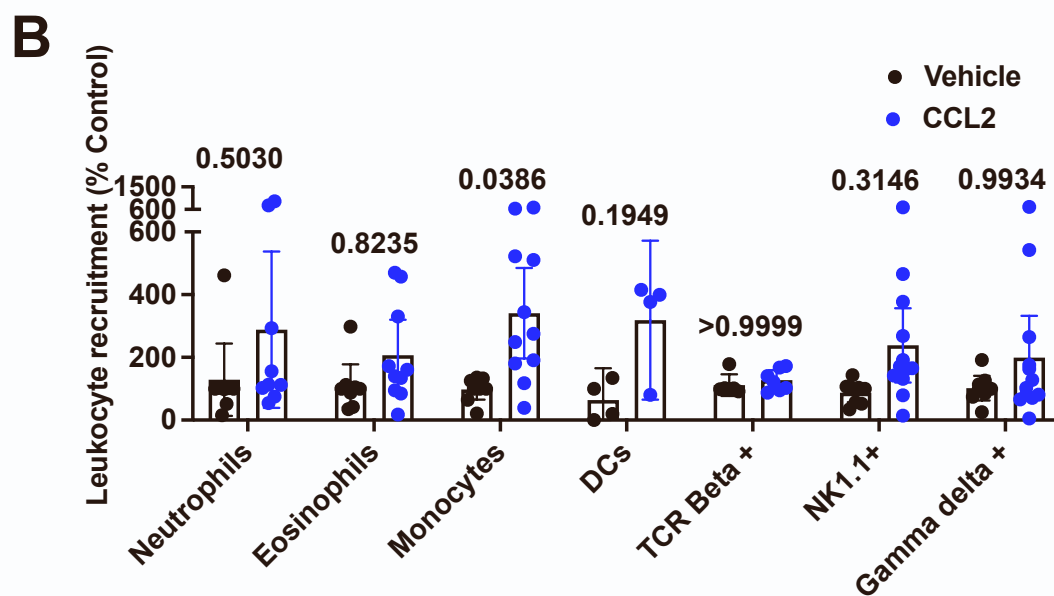
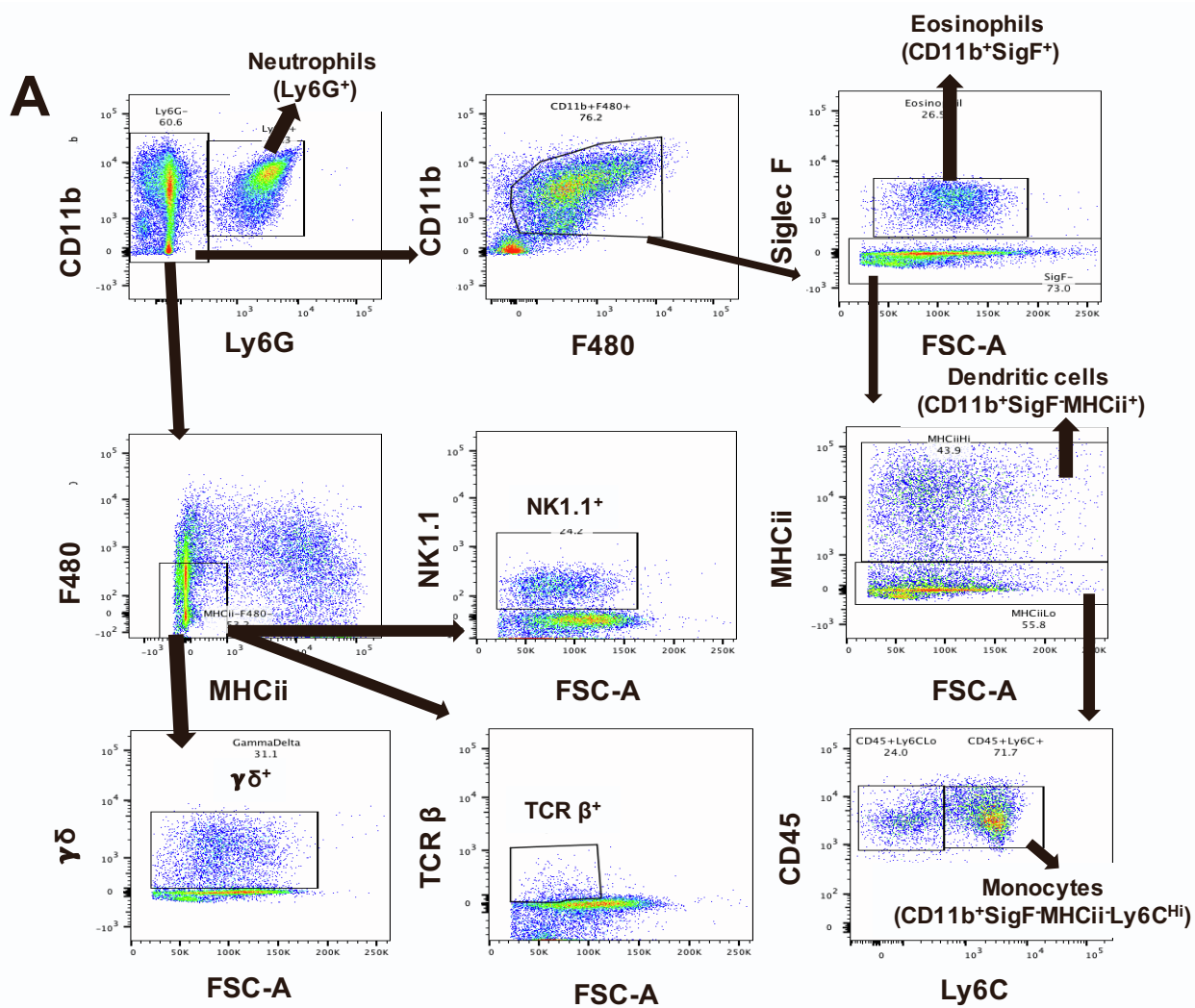
QUANTIFICATION AND STATISTICAL ANALYSIS

All statistics were performed using the Prism (GraphPad) software package. Experiments containing two groups were analyzed using an unpaired t test and data containing more than two groups were analyzed using a one-way ANOVA with a post-hoc multiple comparison test. $p < 0.05$ was considered to be statistically significant.

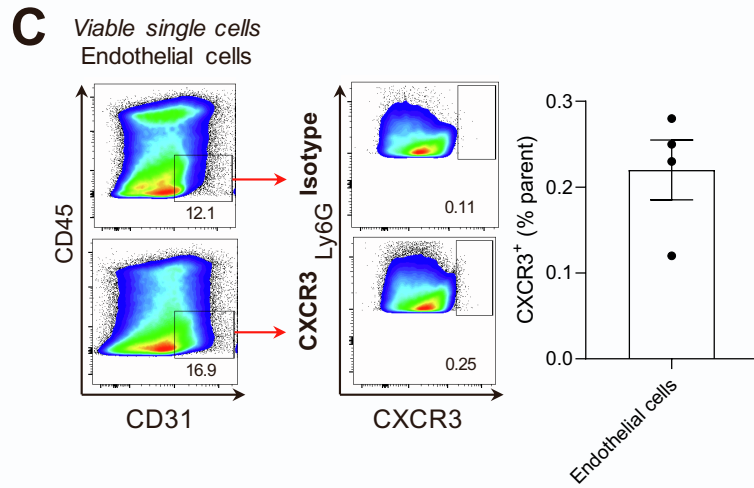
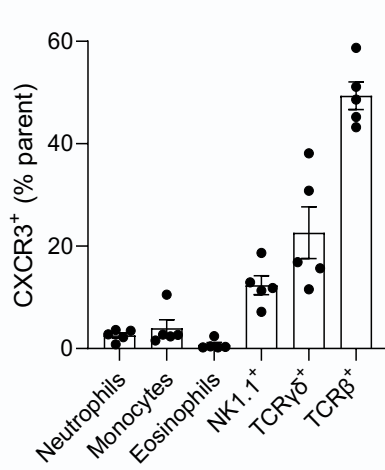
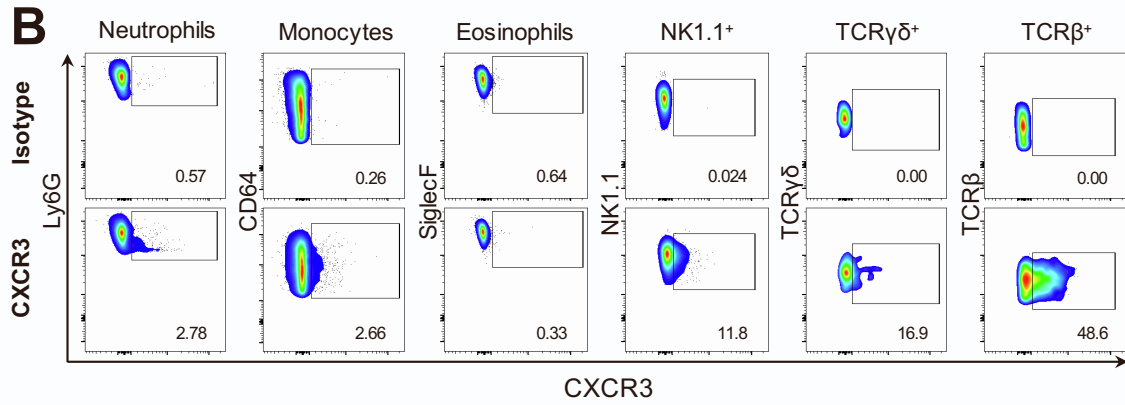
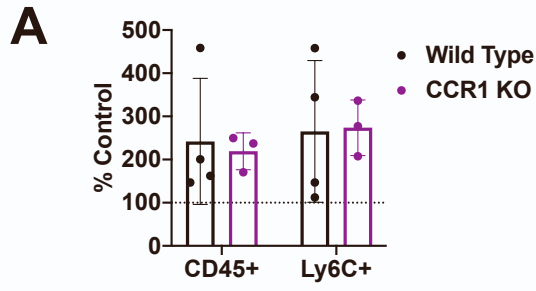
Supplemental information

Chemokine CXCL4 interactions with extracellular matrix proteoglycans mediate widespread immune cell recruitment independent of chemokine receptors

Anna L. Gray, Richard Karlsson, Abigail R.E. Roberts, Amanda J.L. Ridley, Nabina Pun, Bakhtbilland Khan, Craig Lawless, Rafael Luís, Martyna Szpakowska, Andy Chevigné, Catherine E. Hughes, Laura Medina-Ruiz, Holly L. Birchenough, Iashia Z. Mulholland, Catherina L. Salanga, Edwin A. Yates, Jeremy E. Turnbull, Tracy M. Handel, Gerard J. Graham, Thomas A. Jowitt, Ingo Schiessl, Ralf P. Richter, Rebecca L. Miller, and Douglas P. Dyer

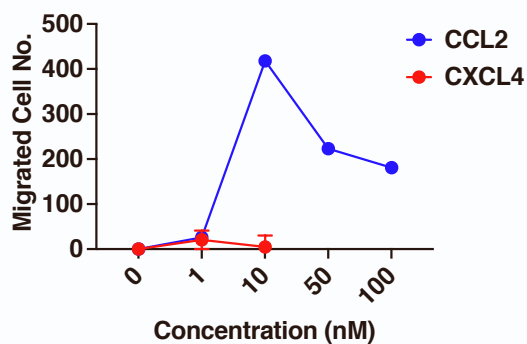
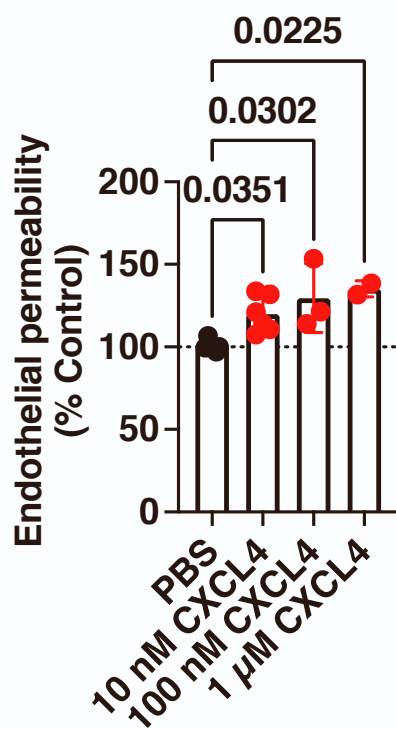
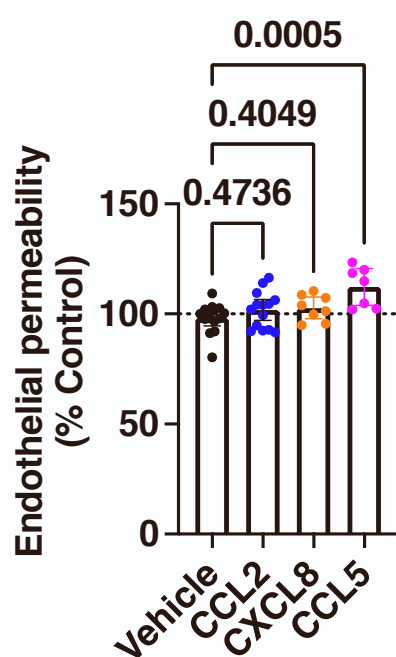
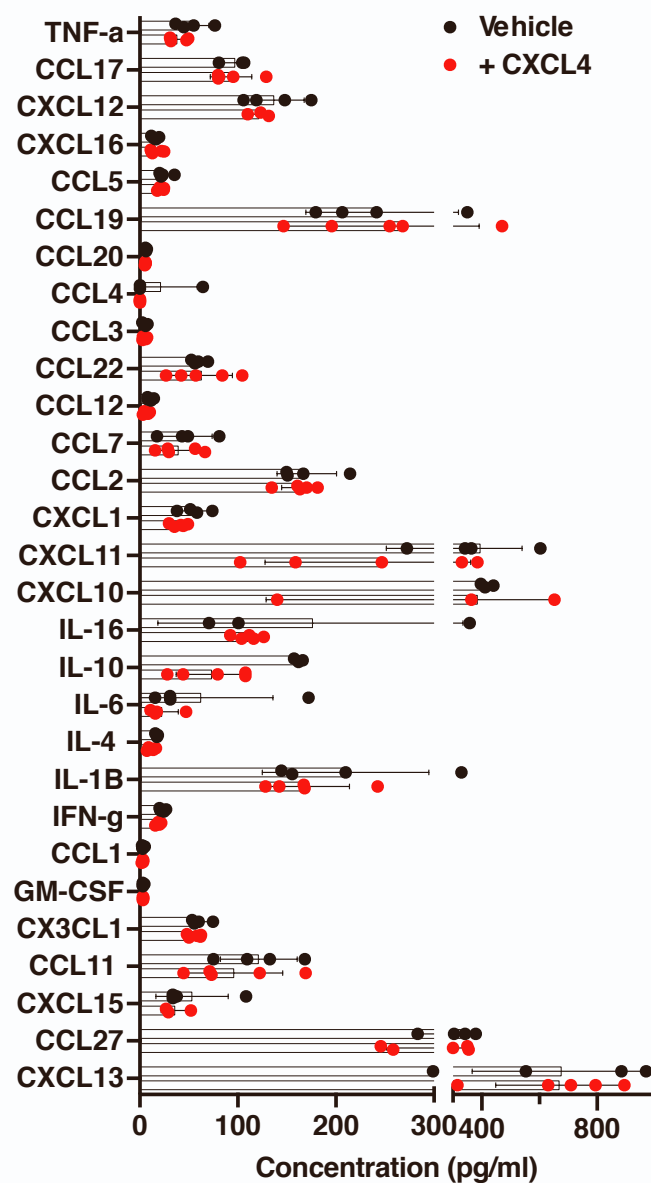
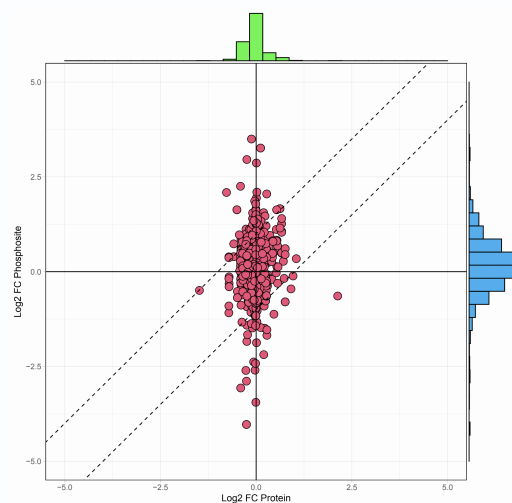


Supplementary Figure 1. Analysis of chemokine mediated recruitment of leukocytes to the murine air pouch. Related to Figure 1. (A) Gating strategy used for flow cytometry analysis of cells retrieved from the air pouch cavity. (B) Quantification of CCL2 mediated leukocyte recruitment to the air pouch. B is the mean with 95% confidence intervals and represents at least two separate experiments where data have been pooled and each dot represents an individual mouse. Data in B is normalised relative to vehicle controls. Individual p values are shown and was analysed using a one-way ANOVA with a post-hoc Sidak analysis of log-transformed data.

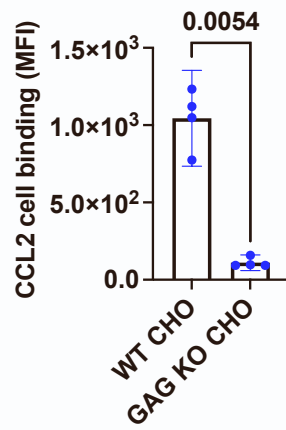
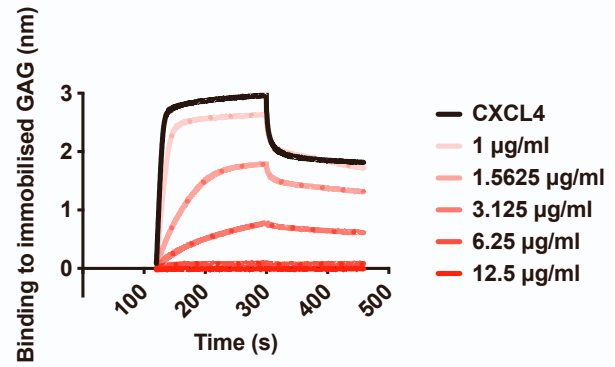
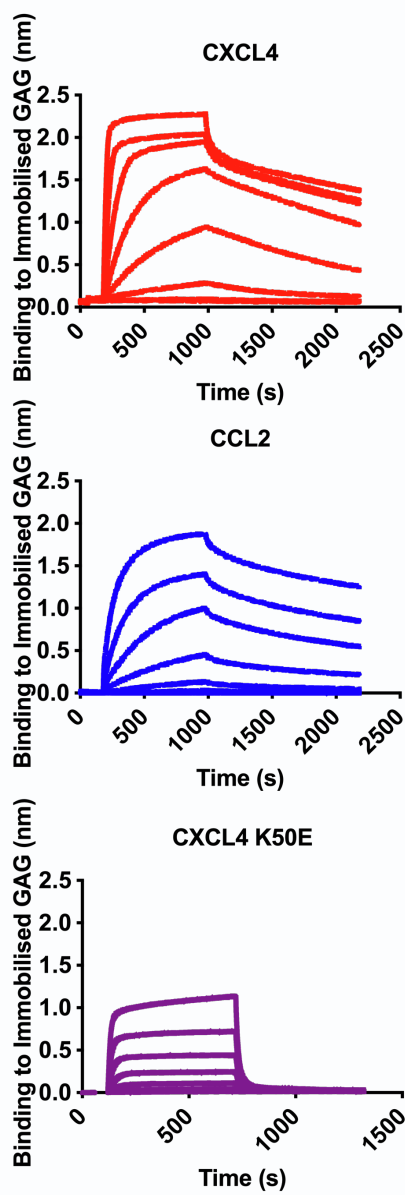
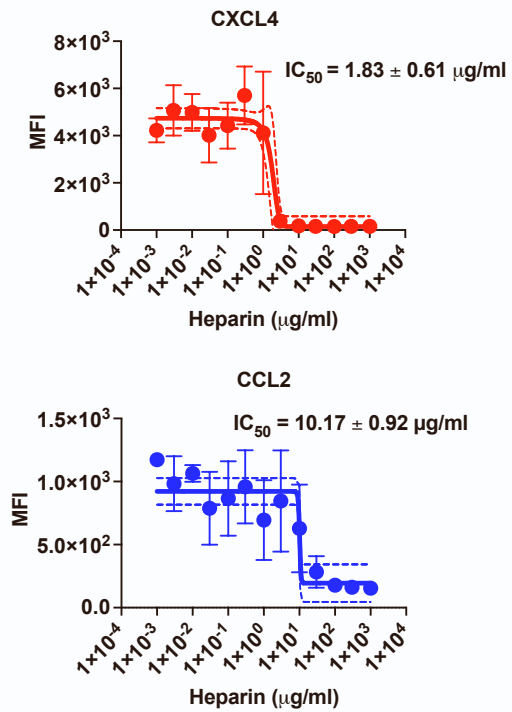
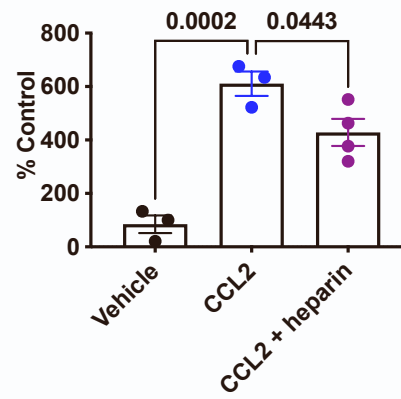


Supplementary Figure 2. Related to Figure 1. CXCL4 mediated recruitment is independent of CCR1 and CXCR3. (A) CXCL4 (5 µg) was injected into dorsal air pouches in wild type and CCR1 KO mice and 24 hours later the recruited cells were analysed using flow cytometry. (B) Indicated cell types were analysed for CXCR3 expression using flow cytometry, squares (gates) are drawn on flow cytometry plots to indicate where cells with positive staining for the leukocyte marker and CXCR3 would sit. (C) Flow cytometry analysis of endothelial (CD31⁺) expression of CXCR3.

A, B (lower panel) and C are mean with 95% confidence intervals and represent at least two separate experiments where data have been pooled, In A, B and C each dot represents an individual mouse. Data in A are normalised relative to vehicle controls.

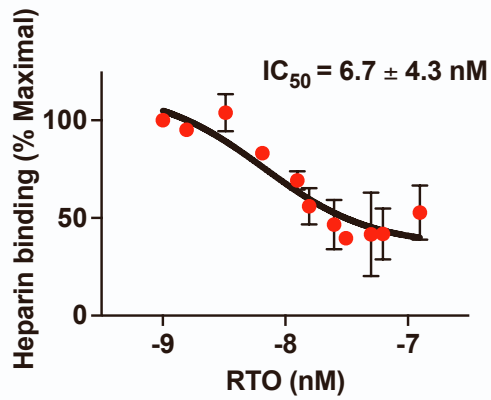
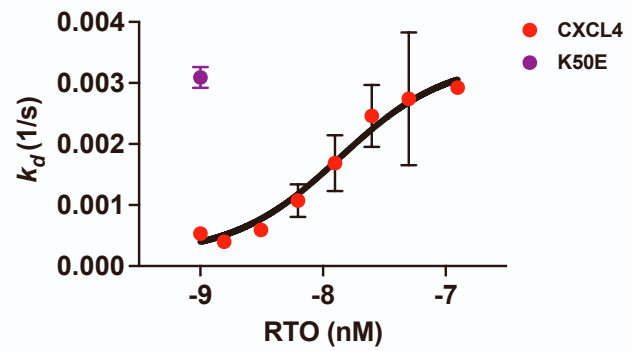
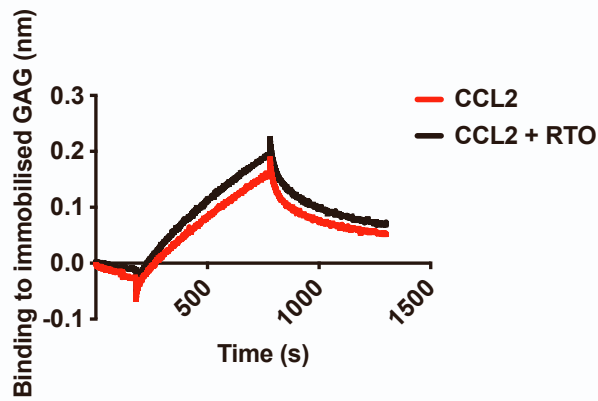
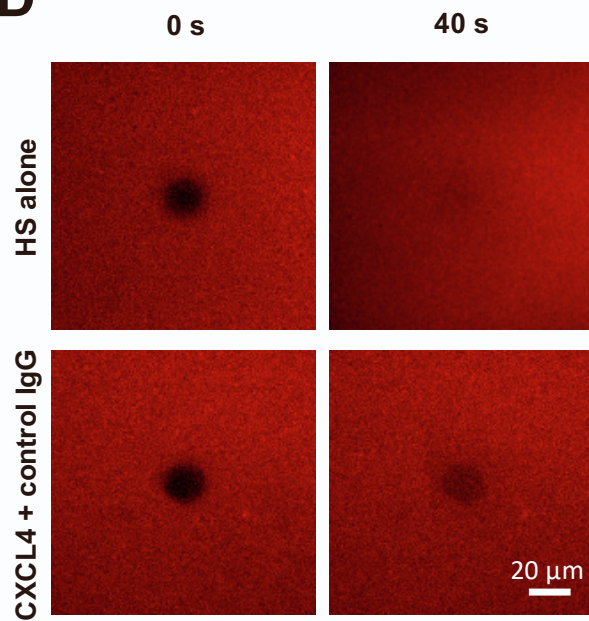
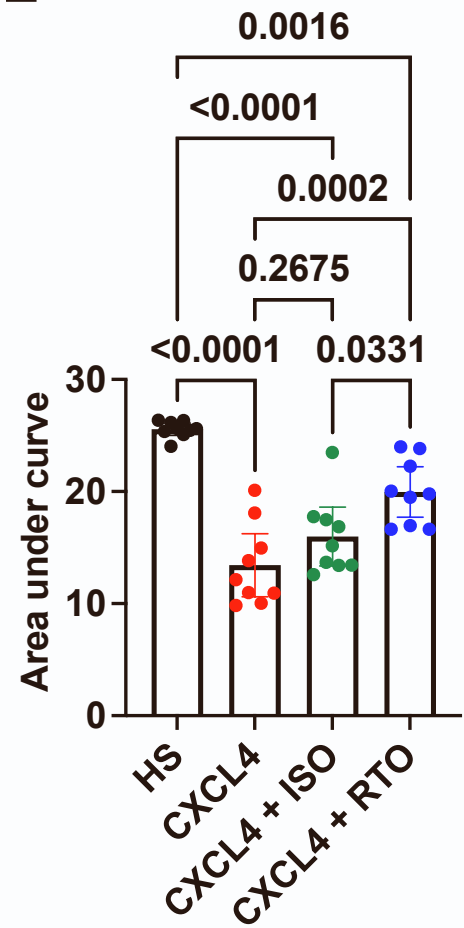
A**B****C****D****E**

Supplementary Figure 3. Related to Figure 2. CXCL4 but not CCL2 increases endothelial permeability *in vitro* and CXCL4 stimulation changes the phosphorylation state of a range of proteins. (A) Chemotaxis assay of human monocytes to CXCL4 or CCL2. (B) Endothelial permeability in response to vehicle control or different amounts of CXCL4. (C) Endothelial permeability with and without CCL2, CXCL8 or CCL5 (10 nM). (D) Air pouches injected with either vehicle or CXCL4 were flushed 24 hours later with PBS that was then analysed using a luminex kit; these data show no change in levels of the indicated proteins between vehicle or CXCL4 injected air pouches. (E) Cultured endothelial cells were stimulated with or without CXCL4 for 2 hours followed by lysis and analysed using phosphorylation enriched mass-spectrometry. The fold-change in protein and phosphorylation levels between vehicle and CXCL4 treated endothelial cells were quantified and plotted against each other. Proteins that sit outside the indicated linear relationship between changes in protein and phosphorylation levels were deemed to have changed phosphorylation status in response to CXCL4. B, C and D are mean with 95% confidence intervals and represent at least two separate experiments where data have been pooled. Each dot in B and C represents a technical replicate and each dot in D represents an individual mouse. Data in B and C is normalised relative to vehicle controls. Individual p values are shown, B and C were analysed using a one-way ANOVA with a post-hoc Sidak analysis.

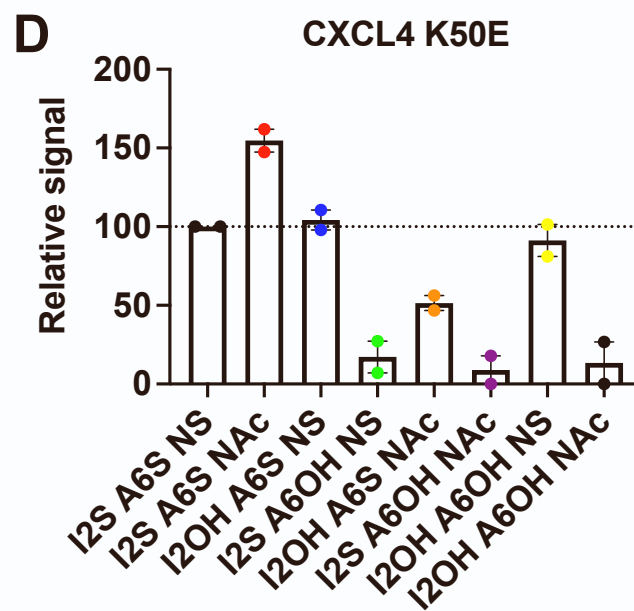
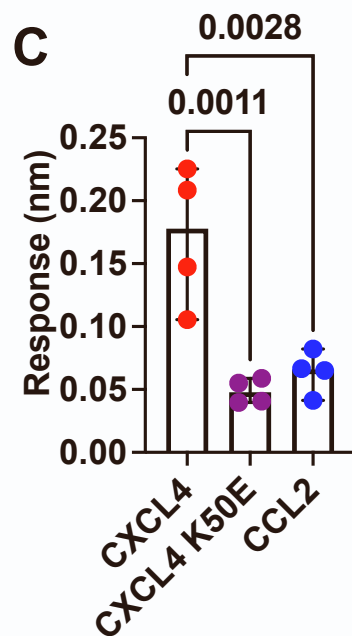
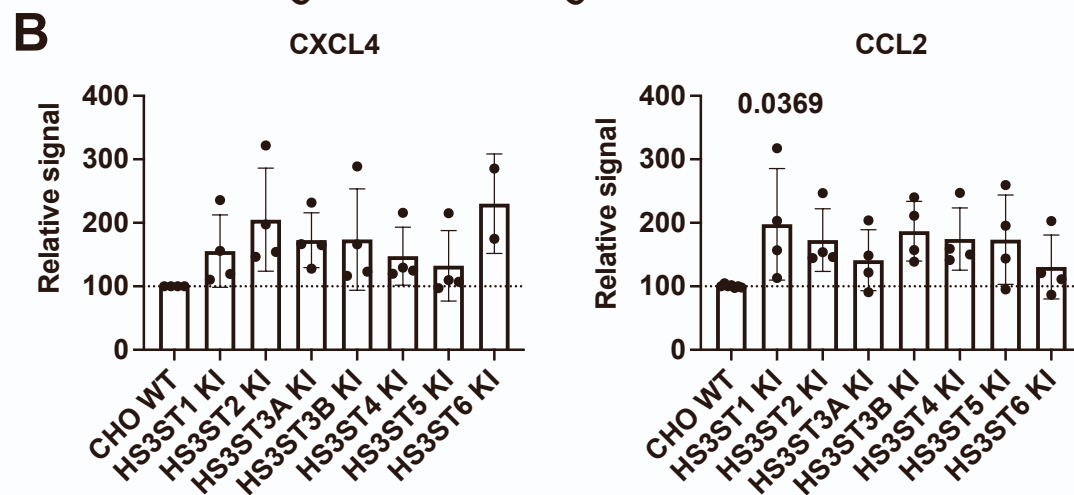
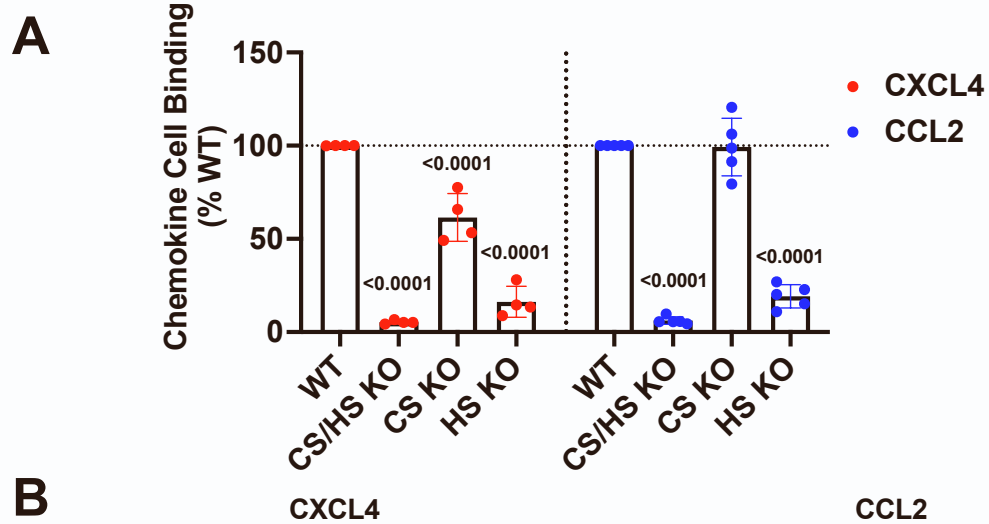
A**C****B****D****E**

Supplementary Figure 4. Related to Figure 3 and 4. Exogenous GAG inhibits CXCL4 and CCL2 binding to the cell surface. (A) CCL2 binding to CHO cells with and without surface GAGs (B4Galt7 KO). (B) CXCL4 (500, 250, 125, 62.5, 31.3, 15.6, 7.8 or 3.9 nM), CCL2 (1000, 500, 250, 125, 62.5, 31.3, 15.6 or 7.8 nM) or CXCL4 K50E (1000, 500, 250, 125, 62.5 or 31.3 nM) were passed over a heparin dp8 surface followed by buffer alone to analyse the interaction over time using BLI. (C) CXCL4 binding to immobilised heparin dp8 over time on its own and with indicated varying concentrations of competing GAG (heparin) in the solution phase. (D) The binding of labelled chemokine to CHO cells was monitored with and without pre-incubation with different concentrations of exogenous GAG (heparin) and analysed for IC₅₀ values. Solid line represents fit of the data using non-linear regression and dotted lines represent 95% confidence intervals of the fit. (E) Analysis of leukocyte (Ly6C⁺ monocyte) recruitment to the air pouch in response to CCL2 with and without exogenous heparin.

A and E are mean with 95% confidence intervals and all plots represent at least two separate experiments. Each dot in A represents a technical replicate and each dot in E represents an individual mouse. Data in E is normalised relative to vehicle controls. Individual p values are shown, A was analysed using an unpaired t-test and E was analysed using a one-way ANOVA with a post-hoc Dunnett analysis.

A**B****C****D****E**

Supplementary Figure 5. Related to Figure 4. CXCL4 oligomerisation drives its' GAG interaction and GAG cross-linking. (A) BLI assay final CXCL4 signal (after washing) is plotted with and without pre-incubation with RTO antibody. (B) Using BLI the CXCL4:dp8 off-rate (k_d) was determined and plotted with and without RTO antibody; CXCL4 mutant K50E off-rate included for comparison. (C) CCL2 (50 nM) with or without the anti-CXCL4 antibody RTO (50 nM) was passed over a heparin dp8 coated sensor and the subsequent interaction was analysed. (D) Images of the bleached region from FRAP experiments with HS alone or CXCL4 isotype (ISO) control immediately after bleaching (0 s) or 40 s after bleaching. (E) Fluorescence recovery over time analysed through the areas under the curves in Fig. 4J (between 0 and 40 s). All plots represent at least two separate experiments. A and B are mean with error where the solid line represents fit of the data points using non-linear regression analysis. Each dot in E represents a separate experiment per condition and is the mean from three recovery curves taken at different positions on the surface. E was analysed using a one-way ANOVA with a post-hoc Dunnett analysis.



Supplementary Figure 6. Related to Figure 5. GAG fine structure mediates selective binding and cell surface retention of chemokines. (A) Chemokine binding to HEK cells with no, HS alone, CS alone or HS and CS combined GAGs knocked out. (B) Chemokine binding to CHO cells with the indicated 3-O GAG sulphation enzymes knocked in. (C) Maximum BLI signal of chemokine (500 nM) binding to fully sulphated heparin (2-O, 6-O and N-sulphated). (D) BLI maximum signal of CXCL4 K50E (500 nM) binding to saturated surfaces of the indicated fully- or de-sulphated heparin samples.

Plots in A-D are mean with 95% confidence intervals and represent at least two separate experiments. Each dot in A-D represents a technical replicate. Data in A, B and D are normalised relative to wild type cells (A and B) or fully sulphated heparin fragments (D). Individual p values are shown and A, B and C were analysed using a one-way ANOVA with a post-hoc Sidak analysis. p values in A and B are comparisons of the indicated groups to WT controls.