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CXCL5-mediated accumulation of mature neutrophils in lung cancer tissues impairs the differentiation program of anticancer CD8 T cells and limits the efficacy of checkpoint inhibitors

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ABSTRACT

Lung tumor-infiltrating neutrophils are known to support growth and dissemination of cancer cells and to suppress T cell responses. However, the precise impact of tissue neutrophils on programming and differentiation of anticancer CD8 T cells in vivo remains poorly understood. Here, we identified cancer cell-autonomous secretion of CXCL5 as sufficient to drive infiltration of mature, protumorigenic neutrophils in a mouse model of non-small cell lung cancer (NSCLC). Consistently, CXCL5 transcripts correlate with neutrophil density and poor prognosis in a large human lung adenocarcinoma compendium. CXCL5 genetic deletion, unlike antibody-mediated depletion, completely and selectively prevented neutrophils accumulation in lung tissues. Depletion of tumor-infiltrating neutrophils promoted expansion of tumor-specific CD8 T cells, differentiation into effector cells and acquisition of cytolytic functions. Transfer of effector functions. Importantly, blocking neutrophils infiltration in the lung, overcame resistance to checkpoint blockade. Hence, this study demonstrates that neutrophils curb acquisition of cytolytic functions in lung tumor tissues and suggests targeting of CXCL5 as a strategy to restore anti-tumoral T cell functions.

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Tumor-infiltrating neutrophils; lung cancer; effector CD8 T cells; immunotherapy; CXCL5

Introduction

Tumor-infiltrating neutrophils (neu) have been associated with worst prognosis in the majority of human cancers and experimental models, with the exception of some selected tumor types.¹ Multiple direct protumorigenic functions related to cancer cell proliferation,^{2,3} metastasis⁴⁻⁷ and angiogenesis^{8,9} have been described. In addition, the capacity of neutrophils to suppress proliferation of T cells ex vivo has been largely documented in humans and experimental mouse models.¹⁰⁻¹⁴ Lung tumors are characterized by increased neutrophil counts in periphery and in tumor tissues and neu accumulation correlates to decreased CD8 T cells and reduced cytotoxicity.¹⁵⁻¹⁹ Moreover, accumulating evidences indicate that neutrophils in lung cancer determine resistance to checkpoint blockade and chemotherapy.²⁰⁻²⁴ Sparse evidences documented changes in the T cell compartment upon neutrophils depletion or drugs blocking recruitment.^{4,13,24,25} Single-cell sequencing and masscytometry recently resolved the diversity and complexity of neu subsets in lung tumor bearing hosts, unveiling multiple neu subsets with graded protumorigenic potential.^{26,27} In parallel, characterization of dysfunctional T cell states within lung tumor tissue provides novel markers to assess the interplay between the myeloid suppressive immune infiltrate and the

anticancer CD8 response.^{28,29} What causal relationship connects subsets of tumor-infiltrating neutrophils to dysfunctional endogenous tumor-specific CD8 T cell responses in vivo in lung tissues remains poorly defined.

Infiltration by neu is often caused by oncogene-driven overexpression of chemokines.³⁰ Overexpression of CXCL5 that binds CXCR2 drives neutrophils infiltration in hepatocellular carcinoma (HCC), intrahepatic cholangiocarcinoma (ICC),^{31,32} melanoma^{33,34} bladder cancer,³⁵ prostate cancer³¹ and breast cancer.^{36,37} In lung tissues, seminal studies described CXCL5 as critical to recruit neutrophils during inflammation and infection.^{38–40} In lung cancer, CXCL5 was shown to induce cancer cell proliferation, migration,^{41,42} autophagy⁴³ and epithelial mesenchymal transition⁴⁴ and it is associated with poor prognosis.^{45,46} CXCL5 is also expressed in experimental models of lung cancer,^{47,48} however the direct link to neutrophils recruitment and interference with antitumoral T cell responses has not been demonstrated.

The orthotopic Kras^{G12D/WT} Tp53 mouse adenocarcinoma model (KP) recapitulates the neutrophils structure found in human lung tumors and it is emerging as a potent preclinical tool to identify immune-cancer networks in human NSCLC.^{29,49,50} KP tumors contain a population of mature

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aged protumorigenic neutrophils expressing the lectin SiglecF,^{26,51} whose precise mechanism of accumulation and interference with anticancer CD8 responses has not been assessed.

Here, we generated an immunogenic variant of the KP model to explore the interplay between neutrophils and anticancer CD8 responses. By RNAseq, we identified CXCL5 as the dominant cancer-cell autonomous upregulated chemokine. Gene editing to delete CXCL5 expression in cancer cells resulted in tumors that selectively lack neutrophils without gross changes in the remaining immune infiltrate. CXCL5null KP clones permitted clonal expansion and activation of effector CD8 T cells functions in lung tissues, and neutrophildepleted tumors became sensitive to treatment with checkpoint inhibitors.

Results

Parental KP and immunogenic KP lung tumors are dominated by SiglecF^{high} neu

To study the interplay between neu and anticancer T cell responses in lung tumor tissue, we established an orthotopic lung adenocarcinoma model (KP cell line),^{27,52} engineered to express the OVA neoantigen (KP-OVA). As previously shown, cancer-specific CD8 T cell responses are established in early tumors but progressively decline due to defective antigen presentation by type 1 DCs⁵³ and accumulation of dysfunctional T cells.^{28,29} Flow cytometry of the immune infiltrate showed a three- to fourfold expansion of CD11b⁺/Ly6G⁺ in both parental and immunogenic KP-OVA tumors (Figure 1a). KP and KP-OVA showed a slight decrease in CD4 and B cells and KP-OVA, as expected, showed a higher frequency of CD8 T cells (Figure 1a and S1A). Staining of lung sections by anti-Ly6G antibodies confirmed that neu are deeply infiltrated in tumor nodules (Figure 1b). Neu expansion was not present in the blood, spleen or tumor draining mediastinal lymph node (mLN), suggesting that orthotopic KP-OVA lung tumors, unlike subcutaneous colon or other lung tumor models,⁵⁴ accumulate neu only locally (Figure 1c). We next analyzed expression of the lectin SiglecF on neu accumulating in orthotopic lung tumors. Up to 70% of CD11b⁺/Ly6G⁺ cells in both KP and KP-OVA tumors expressed SiglecF (SiglecF^{high} neu), whereas the majority of neu in normal lungs were SiglecF low (SiglecF^{low} neu) (Figure 1d). In agreement with previous data, we observed increased PD-L1 expression on SiglecF^{high} neu.⁵⁰ Tissue fluorescence confirmed that SiglechFhigh neu are found within nodules (Figure 1f). A small fraction of SiglecF^{high} neu was found as well in the mLN but not in the non-draining LNs, blood, bone marrow or spleen, in line with previous studies indicating that this subset differentiates specifically in tumor tissues⁵⁰ (Figure 1g).

CXCL5 is highly expressed by mouse lung tumors and correlates to neutrophil infiltrate in human cancers

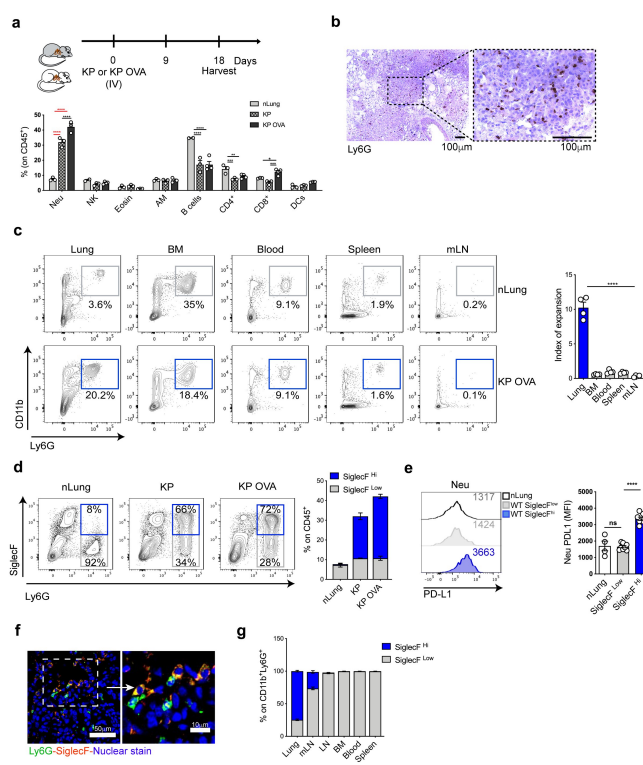
We tested a classical protocol for neutrophil depletion by administration of anti-Ly6G antibodies into KP-OVA bearing mice.^{4,55} Despite efficient depletion in the blood, a large

fraction of neu persisted in lung tumor tissues as previously reported,⁵⁶ precluding further analysis (Fig S2A-D). We thus turned into gene expression data to identify mechanisms of neu accumulation in KP tumors. We identified three chemokines (Cxcl5, Cxcl16 and Ccl21a) in common between the top 10% highly expressed chemokine genes in isolated KP cells and KP-OVA tumor tissues (Figure 2a). Of these, Cxcl5 that is directly implicated in neu recruitment in various cancer tissues and in inflamed lungs was chosen for subsequent analysis. A 20-fold induction of Cxcl5 transcripts in tumor lung tissues was also confirmed using a gene expression array (Fig S3A). To isolate the source of Cxcl5 we analyzed tumor tissues upon fractionation into CD45 negative and positive cells and in isolated KP cells ex vivo. We found high expression of Cxcl5 primarily in CD45 negative cells in cancer tissues and in KP cells (Figure 2(b,c)). We conclude that murine KP lung tumors express high levels of Cxcl5 by a cancer-cell autonomous mechanism.

To explore the correlation between CXCL5 expression and neutrophils recruitment in human lung tumors, we next compiled a large compendium of lung adenocarcinoma (LUAD) (Table 1), that integrates multiple gene expression datasets. Patients were stratified using CIBERSORT ⁵⁷ to create three categories corresponding to low, medium and high neu density. Interestingly, we observed a high significant correlation between *CXCL5* expression and neu density (Figure 2d). Moreover, expression of a gene signature corresponding to tumor-infiltrating neu²⁷ highly correlates to *CXCL5* expression (Figure 2e). In line with previous reports,⁴⁶ we confirmed that expression of CXCL5 is associated with worst prognosis also in this large compendium (Figure 2f). Together, these data indicate that in humans, as in mice, enrichment of neu in lung cancer tissues is associated with overexpression of *CXCL5*.

CXCL5 deletion inhibits accumulation of tumor associated SiglecF^{high} neu

We next tested the possibility to interfere with neu recruitment by targeting CXCL5 expression in KP cells by CRISPR/CAS9 genome editing (Figure3b). WT and KO^{CXCL5} KP-OVA clones were selected by ELISA and validated for OVA expression and growth rate (Figure 3a and S3B-D). Lung tissues of animals challenged with parental KP-OVA cells showed an early peak of Cxcl5 expression that persisted in late tumors. In contrast, lungs challenged with the CRISPR-targeted clone had only a modest increase in Cxcl5, both at early and late time points (Figure 3b). These data demonstrate efficient gene deletion and confirm chemokine production primarily by cancer cells and not by stromal cells conditioned by tumor environmental factors. To test whether deletion of CXCL5 was sufficient to blunt neu arrival we analyzed the immune infiltrate at two consecutive time points. Tumors formed by the parental KP-OVA line showed the expected accumulation of SiglecF^{high} neu already at initial stages, which slightly intensified at later stages (Figure 3c). In sharp contrast, tumors induced by CXCL5 deficient cells were devoid of neu at early stages and showed only a modest increase at later stages. Of the few infiltrating neu in KO^{CXCL5} tumors, a minor fraction expressed SiglecF (Figure 3c). Immunohistochemistry confirmed lack of Ly6G⁺ cells in



Ly6G-SiglecF-Nuclear stain

Figure 1. Neutrophils density in KP and KP OVA orthotopic lung tumors. (a-f) Mice were challenged intravenously with KP, KP-OVA cells or PBS (nLung) as depicted in the scheme. The immune composition was evaluated by flow cytometry in normal lungs (nLung) and established lung tumors, 18 d upon inoculation. (a) Relative abundance of each subset expressed as a fraction of live CD45⁺ cells (neu = neutrophils, Eosin = eosinophils, AM = alveolar macrophages). Data represent mean ± SEM of two to four mice each group. Significance was determined by two-way ANOVA with $*p \le 0.05$; $**p \le 0.01$; $***p \le 0.001$; $***p \le 0.001$. (b) Representative 10× image of paraffine-embedded KP-OVA tumor tissue labelled with Ly6G antibody (brown) and the corresponding 40× magnification. Scale bars (100 µm) are showed. (c) Representative dot plot of neu in lungs (gated on live CD45⁺,Lin⁻), BM (bone marrow), blood, spleen and mLN (mediastinal lymph node) of normal and KP OVA tumor bearing mice. The expansion index represents the ratio between neu in KP-OVA and nLung. Data represent mean ± SEM of four mice each group. (d) Representative dot plots showing SiglecF expression on lung neu (gated on CD45⁺, Lin⁻, Ly6G/CD11b⁺ cells). Numbers in quadrant show percentages of SiglecF^{high/low} among CD11b⁺Ly6G⁺ cells. Bars show quantification of SiglecF^{high/low} neu frequencies on CD45⁺ total lung cells. Data are mean ± SEM of three to five mice/group. (e) PD-L1 expression on lung-infiltrating neu. Representative histogram of PD-L1 expression on neu from nLung or KP-OVA WT tumors. PD-L1 MFI (median fluorescence intensity) are indicated in the histogram and plotted on the right as mean ± SEM of two experiments with two to four mice each group. Significance was determined by one-way ANOVA with ns p > 0.05, **** $p \le 0.0001$. (f) Representative cryo-sections showing Ly6G (green) and SiglecF (red) in tumor nodules and the corresponding magnification. Scale bars (50 or 10 µm) are indicated. (G) Percentage of SiglecF^{high/low} neu within CD11b⁺Ly6G⁺ cells in the indicated organs isolated from KP OVA tumor bearing mice (mLN = mediastinal lymph node, LN = inguinal lymph node, BM = bone marrow). Data represent the mean ± SEM of three to six mice each group.

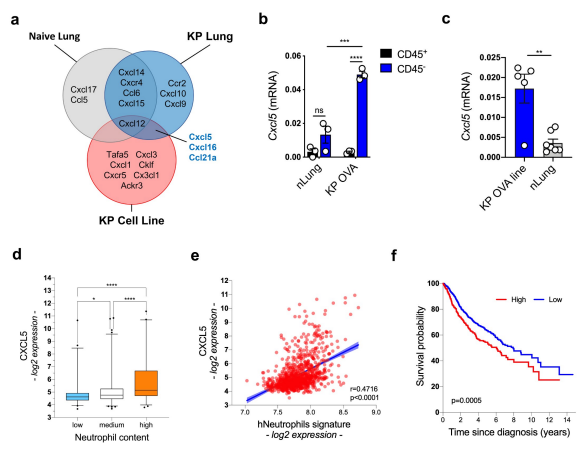


Figure 2. CXCL5 is highly expressed in mice tumors and human LUAD. (a) Venn diagram showing chemokines and chemokine receptors genes that were common between nLung tissues, KP lung tissues and the KP cell line. Top 10% quantile was used as a cut-off for highly expressed genes in each sample. nLung data are from normal C57Bl6/J lung of the Expression Atlas; KP-OVA lung and KP cell line data are from in-house RNA-seq. (b) *Cxcl5* transcripts in CD45⁺ or CD45⁻ cells isolated from nLung or KP OVA tumor bearing lungs were quantified by RT-PCR. Graph shows mean \pm SEM of three replicates, significance was determined by one-way ANOVA with *** $p \le 0.001$; *** $p \le 0.0001$. (c) *Cxcl5* transcripts in the KP-OVA cell line and in nLung. Mean \pm SEM are from five to eight independent RNA extractions, significance was determined by unpaired *t*-test with ** $p \le 0.01$. (d) Patients from the compendium in Table 1, were stratified based on low, medium or high neu content based on CXCL5 or CXCL5 for overall survival of human cancer patients in the compendium (n = 989), comparing top and bottom quartiles.

Table 1. Datasets composing the lung cancer compendium.

| Dataset | Data source | Affymetrix array | Samples | | |
|---------|---|------------------|---------|-----|--|
| | | | AC | SCC | |
| Duke | GSE3141 | HG-U133 Plus 2.0 | 58 | 53 | |
| DKFZ | GSE10245 | HG-U133 Plus 2.0 | 40 | 18 | |
| OCI | GSE14814 | HG-U133A | 28 | 52 | |
| ЕМС | GSE19188 | HG-U133 Plus 2.0 | 40 | 24 | |
| NCCRI | GSE31210 | HG-U133 Plus 2.0 | 226 | - | |
| HLM | GSE68465 | HG-U133A | 92 | - | |
| MI | | | 187 | - | |
| DFCI | | | 82 | - | |
| MSKCC1 | http://cbio.mskcc.org/public/lung_array_data/ | HG-U133A | 107 | _ | |
| MSKCC2 | | | 129 | - | |

AC: adenocarcinoma; SCC: squamous cell carcinoma.

nodules of KO^{CXCL5} challenged mice, both at initial stages and in more advanced tumors (Figure 3d). Lack of CXCL5 secretion by KP-OVA cells had no impact on the frequency of circulating and splenic neu whereas it resulted in higher neu content within the bone marrow, likely reflecting reduced mobilization (S3E-G). We conclude that CXCL5 is a non-redundant mechanism driving neu accumulation in KP lungs. To ascertain that lack of neu recruitment in KO^{CXCL5} tumors is causally linked to

chemokine expression, we reintroduced CXCL5 expression by lentiviral transduction, creating KO^{CXCL5(lenti-CXCL5)} or control KO^{CXCL5(lenti-vec)} (Fig S4A). Reconstituted KP-OVA cells were validated for CXCL5 chemokine production *in vitro* and expression in lung tissues (Fig S4B,C). Importantly, recruitment of neu was restored in the TME of KO^{CXCL5(lenti-CXCL5)} tumors, but not in control KO^{CXCL5(lenti-vec)}, indicating that chemokine expression alone is sufficient to induce neu influx (Fig S4D).

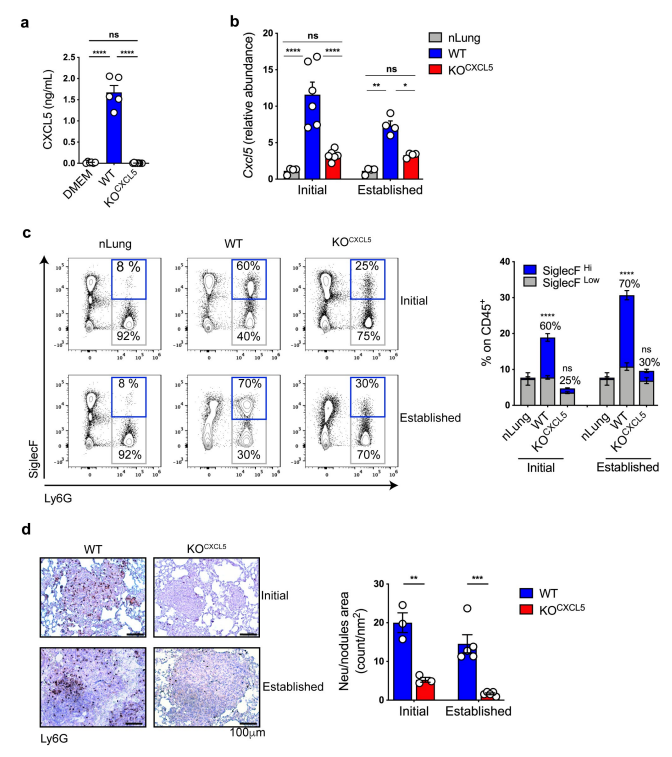


Figure 3. CXCL5 deletion blunts neu recruitment in immunogenic KP lung. (a) Secretion of CXCL5 by parental and gene deleted KP-OVA clones was assessed by ELISA. Data represent the mean \pm SEM of five to nine independent measurements. Significance was determined by a one-way ANOVA with **** $p \le 0.0001$. (b–d) Mice were inoculated with WT or KO^{CXCL5} KP-OVA clones. Lung tissues were harvested after 9 (initial) or 18 (established) d for analysis. (b) Relative abundance of *Cxcl5* transcripts in total lung were evaluated by qRT-PCR. Data are expressed as fold induction over nLung. Data represent the mean \pm SEM of four to six independent RNA extraction. Significance was determined by two-way ANOVA with * $p \le 0.05$, ** $p \le 0.01$, **** $p \le 0.0001$. (c) Representative dot plots showing infiltrating neu and SiglecF expression (gated on CD45⁺CD11b⁺ cells). Numbers in quadrants refer to percentages of SiglecF⁺ on CD11b⁺/LyGG⁺ cells. Bars show the frequencies (%) of neu on live CD45⁺ lung cells. The fractions of SiglecF^{Low} within neu is indicated. Data represent the mean \pm SEM of two independent experiment, three to four mice each group. Significance was determined by a two-way ANOVA with *** $p \le 0.0001$. (d) Representative IHC tissue sections of initial or established nodules labelled with Ly6G antibody (brown dots). Scale bars (100 µm) are indicated. Graph on the right represents quantification of Ly6G⁺ on nodule area (count/nm²). Data are mean \pm SEM of three to five mice each group. Significance was determined by a two-way ANOVA with *** $p \le 0.001$: *** $p \le 0.001$: *** $p \le 0.001$; *** $p \le 0.001$.

Neutrophil density inversely correlates with CD8 T cells in tumor nodules

Given the efficient and selective depletion of infiltrating neu observed in KO^{CXCL5} tumors, we moved to examine the consequences on tumor-specific CD8 T cell responses. Previous studies showed exclusion of CD8 T cells by infiltrating neu in a genetic model of colorectal cancer.²⁵ To explore the relative distribution of the two cell types in lung tumors, we classified nodules into three categories

corresponding to small (area <0.009 mm²), medium (area between 0.0091 mm² and 0.02 mm²) and large (area >0.021 mm²). We found that CD8 T cells outnumbered neu in small nodules and were equal to neu in medium nodules, whereas the proportion was inverted in large nodules that contained more neu than CD8 T cells (Figure 4a), suggesting T cell exclusion. Higher magnification images revealed several neu-CD8 synapses and some areas where CD8 T cells were surrounded by multiple neu, indicative of tight intercellular interactions (Figure 4b).

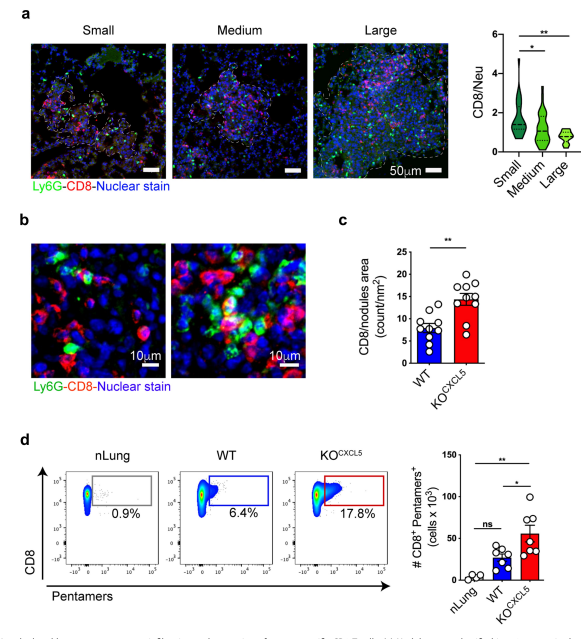


Figure 4. Neu depleted lung tumors promote infiltration and expansion of cancer-specific CD8 T cells. (a) Nodules were classified into 3 categories based on the area (small<0.009 mm², 0.0091 mm² < medium<0.02 mm², large>0.021 mm²). Scale bars (50 µm) are indicated. On the right, quantification of the relative abundance of CD8 and neu within nodules of different sizes. Data are expressed as ratio of CD8/neu (mean \pm SEM of 90 nodules from two independent experiments with 2 mice each). Significance was determined by one-way ANOVA with * $p \le 0.05$, ** $p \le 0.01$. (b) High magnification examples of representative lung sections from WT KP OVA tumors co-labelled with antibodies to CD8 (red) and Ly6G (green) and counterstained with Hoechst to visualize nuclei. (c) Quantification of infiltrating CD8 (CD8 count/nm²) in WT or KO^{CXCL5} KP OVA nodules. Data are from lung tumor sections from two independent experiments with five mice each group (mean \pm SEM). Significance was determined by an unpaired *t*-test with ** $p \le 0.01$. Scale bars (100 µm) are indicated. (d) Representative dot-plot showing OVA class-1 pentamer⁺ cells are plotted. Data are mean \pm SEM of two independent experiments with 3/4 mice/group. Significance was determined by one-way ANOVA with * $p \le 0.05$, ** $p \le 0.01$.

Moreover, the density of CD8 T cell was significantly higher in KO^{CXCL5} tumor nodules that lack neu (Figure 4c). In line with this observation, flow cytometry to measure the fraction of tumor specific, endogenous, CD8 by pentamer staining showed a threefold expansion in neu depleted tumors (Figure 4d).

Neutrophils inhibit markers of effector T cell differentiation and cytolytic functions

To further explore the CD8 compartment in growing lung tumors, animals were challenged with WT or KO^{CXCL5} KP-OVA cells and tissues were harvested at the onset of the T cell responses. The fraction of CD8 T cells carrying activation (CD69) and effector memory differentiation (CD44^{high}/ CD62L^{low}) markers were significantly higher in the lung of KO^{CXCL5} tumors as compared to WT (Figure 5(a,b). Deeper analysis of intracellular transcription factors showed a significantly higher fraction of cells committed toward the effector fate (T-bet⁺/Eomes⁻), in neu depleted tumors (Figure 5c). In both WT and KO^{CXCL5} tumors, CD8 T cells upregulated the exhaustion markers PD-1, indicating exposure to cognate antigen, which was more pronounced in KO^{CXCL5} tumors (Figure 5d). We next investigated expression of TCF-1 transcription, which defines a state of not terminally exhausted cells still retaining proliferative and effector functions. The majority of CD8 T cells in the lung of tumor-bearing mice expressed TCF-1, in agreement with recent data.²⁹ Among PD1⁺/TCF-1⁺ CD8, the fraction of cells producing granzyme was significantly higher in CXCL5^{KO} tumors, indicating maintenance of effector functions and reduced exhaustion (Figure 5e). To further understand whether endogenous CD8 T cells displaying effector functions were tumor specific, we restimulated lung cell suspensions ex vivo with OVA class-I peptide. Notably, tumor tissue resident CD8 T cells secreted twice as much IFN-y in neu depleted tumors than in control tumors (Figure 5f). Finally, to establish if neu actively suppresses effector functions in CD8 T cells, we generated tumor-specific effector CD8 T cells in KP-OVA neu-depleted tumors. Effector cells were then transferred into mice carrying WT or KO^{CXCL5} tumors and the production of IFN-Y by transferred CD8 T cells in the lung was evaluated 2 d later. IFN-Y production was strongly decreased upon transfer in hosts carrying WT tumors, indicating that effector functions are inhibited in the presence of tissueinfiltrating neu (Figure 5(g,h)).

Lung tissue-infiltrating neu limit the capacity of anti PD-L1 antibodies to reduce tumor growth

We next examined the impact of selective deletion of infiltrating neu on growth of KP-OVA lung tumors. The size of tumor nodules and the number of proliferating tumor cells was evaluated in initial and established tumors. At early stages, nodules were significantly reduced in KO^{CXCL5} tumors and the fraction of proliferating cancer cells was diminished (Figure 6a). However, tumor growth was similar between the two genotypes at later time points, indicating that tumor containment by cytotoxic CD8 T cells was not sufficient to control growth at later stages. By decreasing the cancer cells input, we found delayed tumor growth also at later

time point in KO^{CXCL5} tumors (Fig S5). The efficacy of checkpoint blockade was recently shown to inversely correlate to mutations in Stk11, CXCL5 levels and neu recruitment in NSCLC patients.^{20,58} Experimental data further suggested that neu depletion can increase the benefit of chemotherapy by reverting T cell suppression²⁴ and radiation therapy was shown to synergizes with rewiring of aged neu in lung tumors.⁵¹ Moreover, a recent study demonstrated that NETs counteract checkpoint blockade by inducing CD8 T cells exhaustion.⁵⁹ To evaluate whether blocking neu recruitment could synergize with anti-PD-L1 treatment in our model, we inoculated mice with WT or KO^{CXCL5} KP-OVA followed by administration of antibodies to PD-L1 or isotype every 3 d (Figure 6b). In line with previous studies,⁶⁰ the treatment had no impact on the growth of WT KP tumors. In contrast, PD-L1 blockade effectively decreased the growth of KO^{CXCL5} tumors (Figure 6(c,d)). We concluded that CXCL5-driven accumulation of neu in KP lung tumors prevent acquisition of anticancer CD8 effector functions and limits the activity of checkpoint blockade.

Discussion

This study shows that blunting neu recruitment in lung cancer tissues allows expansion, effector state commitment and cytoxicity of tumor-specific CD8 T cells. These findings expand and complement previous studies by providing a more detailed analysis of neu-driven changes in anti-tumoral CD8 responses in vivo, in the original environment of the tumor. Recent advances unveiled the complexity of neu and CD8 T cells states in cancer tissues, revealing the importance of studying tumors in the native tissue of origin and the need to increase the markers used to define effector functions and exhaustion.^{27,29} The negative association of neu density with patient outcome in lung adenocarcinoma remains mostly correlative, thus we focused on a valid lung cancer experimental model to address the crosstalk between neu and tumor-specific CD8 T cell responses in tissues. In line with past data, transplantable KP lung tumors and, as demonstrated here, immunogenic KP-OVA, accumulated a large fraction of SiglecFhigh neu, a mature subset with protumorigenic functions.^{26,51} Since neu depletion by conventional antibodies to Ly6G failed in lung tissues, we sought to identify targetable pathways guiding their recruitment to the lung. The chemokine CXCL5 turned out to be the dominant CXCR2 ligand expressed in KP cancer cells, in line with previous data showing upregulation of chemokines downstream oncogenic Kras.⁴⁸ Despite other pathways such as G-CSF⁵¹ and IL-17⁶¹ may operate in neu recruitment to the lung, genetic deletion of CXCL5 in murine cancer cells was sufficient to completely inhibit accumulation of mature neu. SiglecF^{high} neu accumulate as well in slowly progressing autochthonous KP tumors,⁵¹ yet it remains to be explored whether CXCL5 acts as a dominant factor controlling recruitment also in this context. Of relevance, CXCL5 expression in human lung cancer patients is emerging as a biomarker for prognosis and response to therapy^{45,58,62} and we found that its expression highly correlates to neu density and predicts survival across a large human compendium suggesting a similar nonredundant role in lung cancer patients. It still remains to be established, however, what is the major cellular source of the chemokine in the human context.

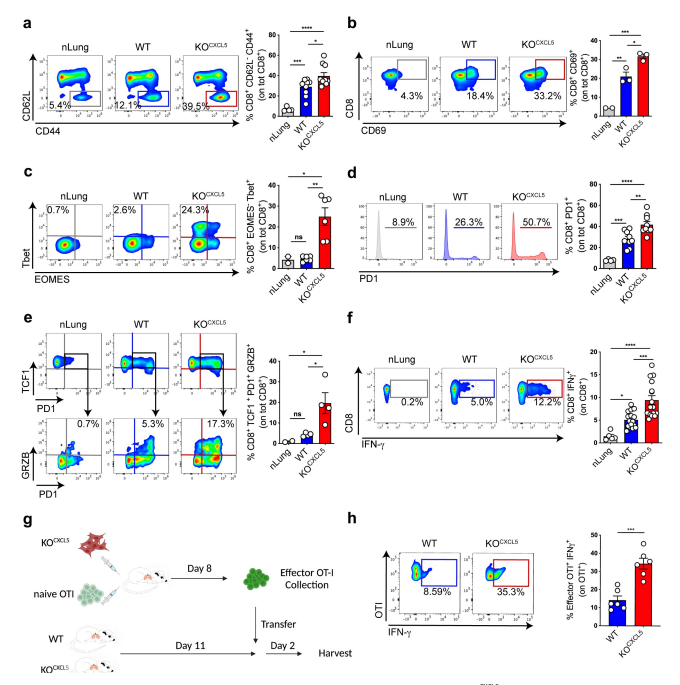


Figure 5. Lung-infiltrating neu restrict differentiation into effector CD8 T cells. (a–e) Mice were challenged with WT or KO^{CXCLS} KP-OVA cells and endogenous T cell responses in the lungs were analysed by flow cytometry. (a) Representative plots showing CD62L and CD44 expression on lung CD8 T cells and corresponding quantification. (b) Representative plots and quantification of CD69 in lung CD8 T cells. (cC) Expression of T-bet and EOMES in lung CD8 T cells.) (d) Representative histograms and quantification of PD-1 expression on total lung CD8 T cells. (e) Example and quantification of the percent of granzyme and PD-1 expression levels on TCF1-expressing lung CD8 T cells. (f) Single-cell lung suspensions were restimulated ex-vivo with OVA class-1 peptide and intracellular staining was performed to determine IFN- γ expression levels. Representative dot-plots and quantifications are shown. Data (a–f) represent mean ± SEM of two to eight mice/group. Significance was determined by one-way ANOVA with ns p > 0.05, * $p \le 0.05$, * $p \le 0.01$, *** $p \le 0.001$. (g) Scheme of OVA-specific CD8 effector adoptive transferred into CD45.1 recipient carrying WT or KO^{CXCL5} cancer cells and adoptively transferred with OT-1. Effector OT-1 cells were sorted from the mLN after 8 d and re-transferred into CD45.1 recipient carrying WT or KO^{CXCL5} tumors. Two days later lung tissues were harvested to perform intracellular staining for IFN- γ as described in (f). (h) Dot plot and quantification of effector OT-1 on total lung OT-1 T cells. Data represent mean ± SEM of four to six mice each group. Significance was determined by unpaired t-test with * $p \le 0.05$, *** $p \le 0.05$,

Tumors lacking a neu infiltrate, offered a well-controlled approach to compare the phenotype of CD8 T cells in tissues. Tumor devoid of neu showed a larger expansion of tumorspecific CD8 T cells and a significant upregulation of surface markers and transcription factors defining effector T cells, as compared to neu-infiltrated tumors. Based on recent evidences describing diverse levels of T cell dysfunction in lung tumors,^{28,63} we propose that neu contribute to induce a hyporesponsive state with reduced effector functions in CD8 T cells. Consistently, neu confer resistance to PD-L1 blockade, which is in line with the correlation between neu density and response to immunotherapy in human cohorts of NSCLC treated with ICB.⁶⁴ One possibility is that neu depletion lower the load of PD-L1 molecules reducing terminal exhaustion and allowing reactivation upon therapy. Alternatively, other inhibitory signals arising from mature tissue neu render T cells

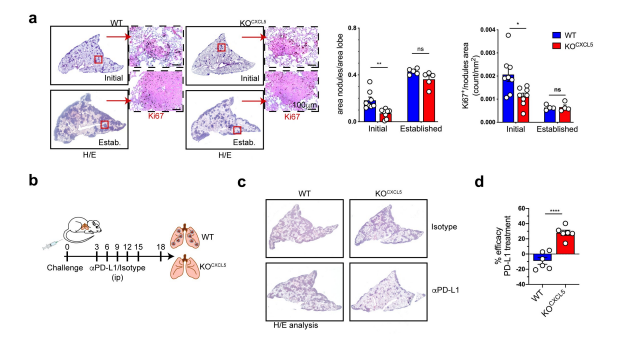


Figure 6. Targeting CXCL5-mediate neutrophils recruitment increase the effectiveness of checkpoint blockade. (a) Mice were challenged with WT or KO^{CXCL5} KP-OVA cells and lungs were harvested after 9 or 18 d to score tumor burden (area nodules/area lobe) and cancer cell proliferation. Representative images of tumor nodules labelled by HE and Ki67 staining. Bars show quantification of tumor burden (area nodules/total lobe area) and quantification of the number of Ki67⁺ cells (expressed as number of cells/nodule area). Data are mean \pm SEM of one to two experiments with four to five mice each group. Significance was determined by two-way ANOVA with ns p > 0.05, * $p \le 0.05$, *p

refractory to reactivation by checkpoint inhibitors. Previous studies proposed several mechanisms of T cell suppression by neutrophils including NOS and ROS, Arginase1, Fas-FasL induced expression apoptosis, PD-L1 and PGE2 production.4,11,12,65,66 The suppressive mechanisms operating preferentially in vivo in different cancer tissues need to be further elucidated and are likely to be context dependent. The present study suggests that future analysis should be performed in vivo using multiple parameters to define the extent of T cell dysfunction, beyond assays of ex-vivo T cell suppression. This study also provides a proof of concept that neutralizing pathological overexpression of CXCL5 by antibodies may be viable and more specific than blocking chemokine receptors ²⁴ to reinforce the efficacy of checkpoint inhibitors.

In conclusion our findings uncover that SiglechF^{high} neu accumulating in lung tissues curb development of effector CD8 T cell functions and impede reactivation by checkpoint block-ade, suggesting CXCL5 as an interesting axis to target in order to restore an immune responsive tumor environment.

Material and methods

Mice

C57BL/6 and OT-I (C57BL/6-Tg(TcraTcrb)1100Mjb/J) mice were purchased from Envigo or Jackson Laboratories, respectively. Animals were maintained in sterile isolators at the ICGEB animal Bio-experimentation facility (12 h/12 h light and dark cycle, $21^{\circ}C \pm 2^{\circ}C$).

Sample size was determined based upon prior knowledge of the intragroup variation of tumor challenges by our laboratory and published studies^{49,50} and was sufficient to detect meaningful biological differences with good reproducibility. The study was approved by International Centre for Genetic Engineering and Biotechnology (ICGEB) board for animal welfare and authorized by the Italian Ministry of Health (approval number 1133/2020-PR, issued on 12/11/2020). Animal care and treatment were conducted with national and international laws and policies (European Economic Community Council Directive 86/609; OJL 358; December 12, 1987). All experiments were performed in accordance with the Federation of European Laboratory Animal Science Association (FELASA) guidelines

Cell lines

The KP cells (LG1233) were generated from lung tumors of C57BL/6 KP mice (K-ras^{LSLG12D/+};p53^{fl/fl} mice) and kindly provided by Dr Tyler Jacks (Massachusetts Institute of Technology, Cambridge, USA).⁶⁷ KP OVA cells were generated by transducing with the lentiviral vector Pdual-liOVAha-PuroR as described in.⁵³ To generate KO^{CXCL5} cells, the KP OVA cells were transiently co-transfected with pSpCas9(BB)-(PX458) and pZac2.1-U6sgRNA-CMV-ZsGreen plasmids, carrying 5'-caccgCTGCCGCAGCATCTAGCTGA-3' guide. The ZsGreen⁺ cells were sorted and single clones were tested for CXCL5 expression by ELISA (Abcam ab100719).

To rescue CXCL5 expression in KP OVA KO^{CXCL5} cells were transduced with a lentivirus carrying the expression vector pLVX-IRES-G418-CXCL5 or empty vector pLVX-IRES-G418 as control. After antibiotic selection, CXCL5 expression was tested by ELISA.

All cell lines were maintained in DMEM media (containing 1 g/L of glucose) supplemented with 10% fetal bovine serum (FBS, Euroclone) and Gentamicin (50 μ g/mL, Gibco) and routinely tested for mycoplasma contamination. The growth rate *in vitro* was assessed by crystal violet assay.

Tissue preparation for flow cytometry

Lung tissues from control or tumor bearing mice were harvested after PBS lung circulatory perfusion, mechanically cut into small pieces and digested with 0.1% Collagenase type 2 (265 U/mL; Worthington) and DNase I (250 U/mL; Thermo Scientific) at 37°C for 30'. Cells were filtered using a 70- μ m cell strainer (Corning), to obtain single-cell suspensions. Blood was collected through subclavian vein puncture, followed by red blood cell lysate. Spleen, lymph nodes (mediastinal and inguinal) were smashed and filtered with a 70 - and 40 - μ m cell strainer, respectively. Bone marrow was extracted from femur and tibia, flush to obtain a single-cell suspension.

Flow cytometry

For cell staining, FcR binding sites were blocked by using aCD16/CD32 and viability of cells was assessed by staining with LIVE/DEAD dye). The antibodies used for the experiments are listed in Supplementary Table 1. To analyze *Cxcl5* expression, CD45⁺ or ⁻ populations from healthy or tumorbearing lungs were stained with CD45-A647 antibody and sorted. MHC-I-OVA pentamers (SIINFEKL/H-2Kb Pro5, Proimmune) were used to identify OVA specific CD8 T cells following manufacturer's instruction. For intracellular detection of IFN γ , single-cell suspensions were stimulated with OVA peptide (SIINFEKL) 2 μ M 37°C for 4 h in the presence of Golgi Stop (monensin, BD Biosciences). Upon extracellular staining, cells were fixed and permeabilized using Cytofix/ Cytoperm solution (BD Biosciences) following manufacturer's instructions.

To identify effector EOMES⁻Tbet⁺ CD8⁺ T cells, we performed a nuclear staining analyzed using Foxp3/transcription factor staining buffer set (Thermo Fisher) following manufacturer's instructions.

To measure OVA expression, cells were fixed and permeabilized using Cytofix/Cytoperm solution (BD Biosciences) following manufacturer's instructions, stained with ratmonoclonal α HA and then with α RAT-AF488. Where indicated, absolute cell count was analyzed by adding TrueCount Beads (BioLegend) to the samples following manufacturer's instructions. Samples were acquired with FACS Celesta (BD Biosciences) and analyzed with FlowJo software (Tree Star, Inc.).

Flow cytometry experiments are based on objective measurements and blinding was not required.

Real-time PCR

RNA from total lungs or sorted cells was extracted using Trizol reagent (Thermo Fisher Scientific), according to manufacturer's instruction. cDNA was synthesized using SuperscriptII (ThermoFisher) and quantitative real-time PCR (qRT-PCR) was performed using SsoFast EvaGreen Supermix (Biorad) with specific primers: Gapdh For (AGAAGGTGGTGAAGCAGGCAT) Rev (CGAAGGTGGAAGAGTGGGAGT), Cxcl5 For (GCT GCC CCT TCC TCA GTC AT) Rev (CAC CGT AGG GCA CTG TGG AC). Gene expression profiling of inflammatory cytokines and receptors of normal and KP-OVA tumor bearing lungs was performed by custom RT² Profiler PCR Array (Qiagen, cat. no. 330221) following manufacturer's instructions.

Immunohistochemistry

To assess tumor burden, lung tissues were harvested and fixed in formaldehyde 10% and paraffine embedded following standard procedure. Consecutive sections of 8 µm were dewaxed and rehydrated and stained with the H&E using (Bio-Optica, Milano Spa). The area of tumor nodules was quantified manually over consecutive sections and averaged (three sections/sample).

To identify neutrophils or proliferating cells within nodules, sections were treated with antigen-retrieval solution (Vector laboratories) for 20 min at 120°C. Slides were treated for 10 min in H₂O₂ and after blocking in 10% goat serum in 0.1% Tween 20 for 30 min and incubated overnight at 4°C with specific antibody diluted in PBS 0.1%Tween 20: antimouse Ly6G (1A8, BD Pharmingen, cat. no. 551459), or antimouse Ki67 (D3B5, Cell Signaling, cat. no. 12202s). Detection was performed using the ImmPRESS polymer detection system (Vector Laboratories), according to manufacturer's Instructions. Automatic thresholding and measurements were performed using Ilastik or ImageJ software, respectively. Images were acquired by Leica microscope. For tumor burden, neutrophil and proliferating cells measurements slides were scored blindly by two independent operators.

Immunofluorescence staining

To identify SiglecF^{high} neu, tumor tissues were intratracheally perfused with 1% paraformaldehyde (PFA), fixed in 4% PFA and embedded in a frozen tissue matrix following standard procedure. Sections of 5 μ m were dried 10' at RT, fixed with 4% PFA for 15' and permeabilized for 15' with PBS 0,5% Triton. After blocking in 5% mouse serum in PBS 1%BSA/0,1%NP-40 for 30', slides were incubated overnight at 4°C with specific antibody diluted in PBS 1%BSA/0,1%NP-40: anti-mouse Ly6G-PE and anti-mouse SiglecF-BB515 (listed in Supplementary Table 1). Nucleus were labeled by Hoechst 15' at RT. Images were acquired with LSM880 META reverse microscope.

To assess spatial distribution of neutrophils and CD8 T cells, tumor tissues were paraffine embedded. Sections of 8 μ m were dewaxed, rehydrated, treated with antigenretrieval solution, and incubated overnight at 4°C with rat antimouse CD8 (4SM15, Invitrogen) and rabbit anti-mouse Ly6G

(E6Z1T, Cell Signaling) antibodies diluted in PBS 0.1% Tween 20 followed by α -rat 647 (Invitrogen) and α -rabbit AF488 (Invitrogen) nucleus were labeled by Hoechst.

Images were acquired with C1 Nikon reverse microscope. Automatic thresholding was performed by Ilastik. ImageJ software was used to quantify CD8 T cells and neu and to measure nodule's area. Nodules having an area $<0.09 \text{ mm}^2$ were classified as small, the ones having an area ranging from 0.091 to 0.2 mm² as medium and those with an area $>0.2 \text{ mm}^2$ as large nodules. Spatial distribution of neu and T cells was performed blindly on code-labeled slides.

In vivo tumor challenge and blocking studies

To establish the adenocarcinoma tumor models C57BL/6 WT female mice at 8–10 weeks of age, were intravenously injected with 7×10^4 KP cells or with 2×10^5 KP OVA variants (WT or KO^{CXCL5}, KO^{CXCL5} (lenti-CXCL5)</sup> or KO^{CXCL5} (lenti-vec)</sup>). In one experiment (supplementary Figure 5) mice were challenged with 5×10^4 cells. Otherwise indicated, mice were sacrificed at initial (9 days) or at established stage of tumorigenesis (18 d). To deplete neutrophils *in vivo*, KP OVA bearing mice were intraperitoneally treated every 3 d, starting from the 6th to 12th d, with 200 µg of anti-Ly6G antibody (InVivo Plus, clone 1A8, Bio X Cell) or isotype control (Rat IgG2a isotype control, clone 2A3, Bio X Cell) and sacrificed at d 13.

In PD-L1 blockade experiments, mice were challenged with KP OVA WT or KO^{CXCL5} cells, intraperitoneally treated every 3 d with 200 μ g of α PD-L1 (InVivoMab, clone 10 F.9G2, BioXcell) or isotype control (InVivoMab, rat IgG2b isotype control, clone LTF-2, BioXcell). Mice were sacrificed at d 18.

Adoptive transfer of effector OVA-specific CD8

2x10⁶ naïve OTI T cells were intravenously injected into C57BL6/CD45.2 congenic mice followed by inoculation of 4×10^5 KP-OVA KO^{CXCL5} tumor cells. After 8 d, the mLNs were collected and effector CD44⁺ CD62l⁻ CD8⁺ T cells were isolated by cell sorting and intravenously transferred into CD45.1 recipient (0.1 × 10⁵/mouse) mice that had been challenged with WT or KO^{CXCL}KP-OVA cells 10 d before. Lung tissues were harvested 2 d after adoptive transfer and processed for intracellular IFN- Υ detection as described above.

RNA seq and data analysis

RNA from total lungs was extracted using RNeasy Micro Kit (Qiagen), according to manufacturer's instruction. Libraries were prepared using TruSeq Stranded mRNA LT Sample Prep Kit and sequenced on an Illumina platfor. Raw sequences files' quality was checked via FastQC (v 0.11.9, http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) and was found to satisfy quality check. We have performed reads alignment to the mouse reference genome using Salmon alignment tool, used for transcript quantification, with the following parameters: read length of 100, Mus musculus GRCm39.103 (GTF file), Mus musculus GRCm39 DNA primary assembly fasta (Sequence file). Data was further elaborated performing a trimming with Salmon.

We have therefore obtained the generated gene counts, which was analyzed using R's package DESeq2. Using vsd method we generated the normalized expression matrix which was investigated with Principal Component Analysis (PCA) together with a dimensionality reduction algorithm present in the DESeq2 package. Chemokines and chemokine receptors were ranked based on their normalized expression in KP lung and KP cell line and top 10% expressed genes were selected. The same ranking and cut-off were applied to TPM values of naïve lung (nLung), as retrieved from a C57Bl6/J mouse background from Expression Atlas.

Collection and processing of human lung cancer gene expression data

A lung cancer compendium has been created from seven major datasets comprising microarray data of lung cancer samples annotated with clinical outcome. All data were measured on Affymetrix arrays and have been downloaded from NCBI Gene Expression Omnibus (GEO, http://www.ncbi.nlm.nih.gov/geo/) GSE3141, GSE10245, GSE14814, GSE19188, GSE31210, GSE68465, and form the Ladanyi and Gerald Laboratories Lung Adenocarcinoma microarray repository (http://cbio. mskcc.org/public/lung_array_data/). Prior to analysis, we eliminated duplicate samples and renamed all original sets after the medical center where patients were recruited. This reorganization returned 1,136 unique samples from ten independent cohorts comprising 989 adenocarcinomas, 778 of which with complete clinical outcome information (Table 1). The type and content of clinical and pathological annotations of the compendium samples have been derived from the original cohorts. Since raw data (.CEL files) were available for all samples, the integration, normalization and summarization of gene expression signals has been obtained applying the procedure described in Rustighi et al.⁶⁸ Briefly, expression values were generated in R from intensity signals using a custom CDF obtained merging HG-U133A, HG-U133A2 and HG-U133 Plus2 original CDFs and transforming the original CEL files accordingly. Intensity values have been background-adjusted, normalized using quantile normalization, and gene expression levels calculated using median polish summarization (multiarray average procedure, RMA). Gene expression data for a total of 21,995 probe sets have been collapsed to 12,391 unique gene symbols using the hgu133a.db annotation package (version 3.2.3) and the aggregate function of R stats package. To remove possible batch effects due to the different cohorts, we applied the ComBat function of the sva Bioconductor package to the merged matrix. ComBat was used with default parameters and the batch variable imputed as a vector of dataset labels (see Table 1). Clinical information among the various datasets has been standardized as described in Cordenonsi et al.⁶⁹ Average signature expression has been calculated as the average expression of all signature genes in sample subgroups.

Human neutrophil fraction analysis

Neutrophil cell fractions have been quantified using CIBERSORT⁵⁷ on the lung adenocarcinoma samples of the *LUAD compendium*. Briefly, the non-log linear expression

matrix with the 989 cases was uploaded to the *CIBERSORT* R script (version 1.04) as a mixture file and *CIBERSORT* was run in absolute mode with the LM22 signature gene file, 100 permutations, and quantile normalization. In absolute mode, CIBERSORT scales relative cellular fractions into a score that reflects the absolute proportion of each cell type in a mixture. Although not expressed as a fraction, the absolute score can be directly compared both between- and within-samples.⁷⁰ Samples have been divided into three groups based on the upper (high neutrophil content) and lower (low neutrophil content) quartiles of the absolute scores for the neutrophil cell type. Samples that fell in between the quartile cutoffs have been termed the medium neutrophil content group.⁷¹

Survival analysis

To evaluate the prognostic level of CXCL5 expression, we separated the samples into two groups based on CXCL5 standardized expression.⁶⁹ Tumor samples have been classified as 'Low' if CXCL5 standardized expression was negative and as 'High' if the CXCL5 standardized expression was positive. The overall survival probability in the two groups has been estimated using the Kaplan–Meier method and the Kaplan–Meier curves compared using the log-rank (Mantel–Cox) test. P values were calculated according to the standard normal asymptotic distribution. Survival analysis was performed in GraphPad Prism.

Statistic

Primary data were collected in Microsoft Excel, and statistical analysis was performed using GraphPad Prism 8 software. Values reported in figures are expressed as the standard error of the mean, unless otherwise indicated. For comparison between two or more groups with normally distributed datasets two-tailed Student's *T*-test, multiple *T*-test, one-way ANOVA or two-way ANOVA were used as appropriate. For the comparison of matched groups, we used Wilcoxon test. The nonparametric Kruskal–Wallis test with Dunn's multiple comparison was performed to compare three or more unmatched groups. *p* Values >0.05 were considered not significant, *p* values ≤0.05 were considered significant: **p* ≤ 0.001, *****p* ≤ 0.0001.

No exclusion criteria or data have been performed.

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Disclosure statement

No potential conflict of interest was reported by the author(s).

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Contributors

FS and GMP performed experiments. FS and GMP analyzed data, prepared figures and contributed to manuscript writing. AC provided CXCL5 expression construct. SB and RA performed bioinformatic analysis. NC set up the tumor model and generated initial observations. FB ideated and supervised the study and wrote the manuscript.

Ethical approval

The study was approved by International Centre for Genetic Engineering and Biotechnology (ICGEB) board for animal welfare and authorized by the Italian Ministry of Health (approval number 1133/2020-PR, issued on 12/11/2020). Animal care and treatment were conducted with national and international laws and policies (European Economic Community Council Directive 86/609; OJL 358; December 12, 1987). All experiments were performed in accordance with the Federation of European Laboratory Animal Science Association (FELASA) guidelines

Data availability statement

All data relevant to the study are included in the article or uploaded as supplementary information. The datasets used and analyzed during the current study are included in the article or uploaded as Supplementary Information. They are available from the corresponding author on reasonable request.

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