Combined toll-like receptor 3/7/9 deficiency on host cells results in T-cell-dependent control of tumour growth

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Toll-like receptors (TLRs) are located either on the cell surface or intracellularly in endosomes and their activation normally contributes to the induction of protective immune responses. However, in cancer their activation by endogenous ligands can modulate tumour progression. It is currently unknown how endosomal TLRs regulate endogenous anti-tumour immunity. Here we show that TLR3, 7 and 9 deficiencies on host cells, after initial tumour growth, result in complete tumour regression and induction of anti-tumour immunity. Tumour regression requires the combined absence of all three receptors, is dependent on both CD4 and CD8 T cells and protects the mice from subsequent tumour challenge. While tumours in control mice are infiltrated by higher numbers of regulatory T cells, tumour regression in TLR-deficient mice is paralleled by altered vascular structure and strongly induced influx of cytotoxic and cytokine-producing effector T cells. Thus, endosomal TLRs may represent a molecular link between the inflamed tumour cell phenotype, anti-tumour immunity and the regulation of T-cell activation.

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Toll-like receptors (TLRs) are a conserved family of receptors, well recognized for their ability to respond to pathogenic structures, also known as pathogen-associated molecular patterns. TLRs are located either on the cell surface or intracellularly in endosomes. Although surface TLRs such as TLR2 and TLR4 primarily recognize bacterial proteins, endosomal TLRs primarily detect viral and bacterial nucleic acids. Triggering of TLRs initiates a complex intracellular signalling cascade in activated cells. Among others, MyD88 and TRIF are major adaptor molecules in this cascade. In the immune system, TLR ligation results in the activation of myeloid cells and subsequent induction of anti-pathogenic immunity. In this context, the activation of myeloid immune cells via TLRs represents a link between innate and adaptive immunity.

This immunostimulatory potential of TLR ligation has been used to develop cancer immunotherapies based on synthetic or natural TLR ligands. Examples for both therapeutic ligands of membrane bound as well as endosomal TLRs exist. Compounds targeting endosomal TLRs and mimicking viral and bacterial RNA viral RNA and DNA have attracted a somewhat broader interest. Examples include the use of imiquimod and CpG, ligands for the endosomal TLRs 7 and 9. Imiquimod is a short synthetic RNA and is clinically used to treat actinic keratosis, external genital warts and superficial basal cell carcinoma. CpG, through binding to TLR9, has strong adjuvant activity and has been applied in numerous clinical trials in the treatment of allergy, cancer and infectious diseases.

In the context of these therapeutic applications the expression of TLRs on tumour cells and the consequences of TLR activation on cancer cells have received increasing attention. Studies in this area quickly revealed the dichotomous nature of tumour cell stimulation with TLR ligands. On the one hand, triggering of TLRs has been reported to induce cell death in tumour cells. This cell death may be anti-tumoural in two ways: First, as a direct consequence, the number of tumour cells is reduced. Second, via a process termed ‘immunogenic cell death’, additional activation of anti-tumour immunity may occur. However, some evidence suggests that this type of immunogenic cell death may rather be associated with the effects of TLR ligands on RIG-I-like helicases, another class of pattern recognition receptors responsive to synthetic and pathogenic nucleic acids. Conversely, under certain conditions, TLR ligands may also elicit anti-apoptotic effects, which can even be associated with additional escape from cytolysis by immune cells.

Although the effects on cancer (and epithelial) cells are fairly well understood, the consequences of TLR signalling in fibroblasts and mesenchymal cells remain poorly described. Some studies suggest tumour-promoting effects of TLR expression on cancer-associated fibroblasts. For example, TLR4 expression by stromal fibroblasts in colorectal cancer was associated with poor prognosis and overexpression of TLR3 in fibroblasts results in upregulation of the oncoprotein c-Myc. Other studies showed that high TLR9 messenger RNA expression on fibroblast-like cells in breast or oesophageal squamous cell carcinoma was associated with reduced metastasis and invasion.

In addition to foreign pathogenic structures, the TLRs also recognize non-foreign structures commonly referred to as damage-associated molecular pattern molecules (DAMPs). DAMPs are self-proteins of the host, which are released during pathogenic conditions such as chronic inflammation, autoimmune diseases, sepsis and cancer.

In cancer, DAMPs are released as a consequence of continued cancer-related inflammation, tissue destruction and cell death. The release of the nuclear DNA-binding molecule high-mobility group box 1 (HMGB1) by dying tumour cells activates TLR2 and TLR4 on DCs, resulting in an efficient processing and cross-presentation of tumour antigens and tumour regression. Similarly, members of the S100 family are released during cellular stress and cancer. In immune cells, this leads to TLR4-dependent release of pro-inflammatory cytokines and receptor for advanced glycation endproducts (RAGE)-dependent migration. In addition, histones, if translocated from the nucleus to the extracellular space, function as DAMPs and increased levels of circulating histones were found, for example, in serum of pancreatic cancer patients. Likewise, nucleic acid fragments released from necrotic cancer cells or adjacent injured epithelial cells can act as DAMPs.

Thus, despite some uncertainties, the continued triggering of TLRs by endogenous ligands is mainly associated with tumour-promoting effects. These effects are mostly linked to TLR2 and TLR4. The role for the endosomal TLRs such as TLR3, 7 and 9 is less clear and is investigated in this study. We use syngeneic wild-type (WT) transplantable tumour cells and TLR-deficient recipient mice, to specifically address the role of TLR expression on host cells. During the first 7–10 days, tumour growth is virtually identical in WT and Tlr3/7/9-deficient mice. Unexpectedly, at day 10 a clear tumour rejection is observed in Tlr triple knockout (KO) mice. This phase is associated with a dramatic local inflammatory reaction and substantial changes in the tumour microenvironment, ultimately enabling an unexpected T-cell-dependent complete tumour rejection. In sum, our data suggest that TLR signalling may provide a molecular link between tumour-associated inflammation and anti-tumour immunity.

**Results**

**Tumour rejection in the absence of TLR signalling.** To investigate the role of the endosomal TLRs 3, 7 and 9 on tumour growth, a murine head and neck cancer cell line (murine oropharyngeal carcinoma cell, MOPC) was injected into single KO, double KO and Tlr3/7/9 triple KO mice (Fig. 1a and Supplementary Fig. 1). In the majority of TLR-deficient and WT control mice, a palpable tumour developed within 1 week. Data from single KO mice were comparable to WT mice and thus were not presented in the figures. Unexpectedly, at later points in time, tumours declined in size in Tlr3/7/9 triple KO, but not single or double KO mice, and fully disappeared between day 14 and day 30 (Fig. 1a and Supplementary Fig. 1). In contrast, only one out of six Tlr3/9/7/11 mice could reject the tumour and only two single KO mice did not develop a tumour after 3–4 weeks. Histologic analysis of tumour sections, taken from WT and Tlr3/7/9/11 mice at different time points during the initial growth and the later rejection phase, confirmed tumour regression (Fig. 1b). To exclude cell line-specific effects, two additional syngeneic tumour cell lines (TC1, MB49) were injected into Tlr triple KO mice. Cell numbers for the inoculum were determined according to titration experiments performed in WT mice. Again, all TLR3/7/9-deficient mice and WT mice developed tumours during the initial 1–2 weeks post injection (depending on the cell line used), but only Tlr triple KO mice rejected tumours within 40 days (Supplementary Fig. 2).

To test whether tumour rejection is caused by tumour cell death, we performed TdT-mediated dUTP nick end labelling (TUNEL) staining on tissue sections. Although only little cell death could be detected in both WT and Tlr triple KO mice during the first week of tumour growth, a dramatic increase in TUNEL-positive cells was observed in TLR-deficient mice at day 10 and day 14 (Fig. 1c,d), suggesting substantial tumour cell death during regression.

Tumour growth may be influenced by the local tissue microenvironment. To explore whether this phenomenon is limited to...
the subcutaneous (s.c.) injection site, we next extended our analysis to an orthotopic model recently developed by our group. In this model, the tumour cells are injected into the floor of the mouth and the weight of the mice is recorded as a measure of tumour growth. WT and Tlr3/7/9−/− mice started to lose weight during the initial growth phase as a consequence of impaired food uptake. Around day 10, analogous to the s.c. model, Tlr3/7/9−/− mice started to recover to normal body weight. In contrast, WT mice showed progressive tumour growth and were killed according to animal welfare regulations (Fig. 1e). These data indicate that the

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**Figure 1 | Tlr3/7/9−/− mice reject MOPC tumours within 3 weeks through induction of cell death.** (a) MOPC tumour cells (0.5 × 10^6) were s.c. injected into the right flank of 8- to 16-week-old C57BL/6 WT (n = 19) and Tlr3/7/9−/− mice (n = 24). Mean of tumour growth ± s.e.m. and tumour take (percentage of mice with a palpable tumour) are shown. Three stars (*** ) indicate statistical difference P ≤ 0.0005 (t-test). (b) Frozen sections of s.c. tumours, dissected at different time points after tumour cell injection (day 1 to day 28), were stained by haematoxylin–eosin staining. One representative example for each time point is shown. Scale bar, 1,000 µm. (c) Quantification of dead cells in MOPC tumours was determined by calculating the area of TUNEL-positive cells in relation to the entire tumour area at day 3, day 6, day 10 and day 14 after tumour cell injection (n = 3–6). Single values and mean (horizontal line) are shown. Results were considered significant at *P ≤ 0.05 and *** P ≤ 0.0005 (t-test). (d) Representative examples of the TUNEL staining at day 10 and day 14 of tumours of WT and Tlr3/7/9−/− mice. Green: TUNEL, red: 7-AAD nuclear staining. Scale bar, 100 µm. (e) Tumour growth of 0.5 × 10^6 MOPC cells after orthotopic (ortho) injection into the floor of the mouth in 8- to 12-week-old C57BL/6 WT (n = 4) and in Tlr3/7/9−/− mice (n = 5). Relative weight of the mice is shown. Mice were killed if they lost 20% of their initial weight, indicated by the dotted red line.
mechanism of tumour rejection is not limited to the s.c. site, but rather maintained at the orthotopic site.

**Immune and non-immune cells are involved in tumour rejection.**

The stromal tumour microenvironment consists of immune cells and non-immune stromal cells. Although immune cells are of haematopoietic origin, the remaining stromal cells are largely non-haematopoietic and contain vessels and mesenchymal fibroblastoid cells. We performed classical bone marrow chimera experiments, to test whether tumour rejection is mediated via TLR deficiency on haematopoietic or non-haematopoietic cells. As depicted in Fig. 2a, tumour rejection only occurred in irradiated Tlr\(^{-/-}\) mice, which received Tlr\(^{-/-}\) bone marrow (BM; designated KO control). Both chimera types (WT BM into KO host and KO BM into WT host) were permissive for tumour outgrowth, suggesting that TLR deficiency on both haematopoietic and non-haematopoietic stromal cells is required for complete tumour rejection. When we quantified tumour growth by caliper measurements on day 30 of the experiment (Fig. 2b), we found an expected great variance in tumour size between individual animals. No or very small tumours could be measured in KO control mice, whereas tumours were clearly measurable in WT control and in both types of chimera. These data suggest that cooperative activity of both haematopoietic and non-haematopoietic, radioresistant TLR-deficient stromal cells is required for tumour rejection.

Blood vessels are a major component of the non-haematopoietic stromal tumour microenvironment. When we analysed the vessel composition by CD31 immunohistochemistry between day 6 and 14, we found substantial differences between WT and Tlr\(^{-/-}\) mice. At day 10 and 14, gross microscopic inspection (Fig. 3a) already suggested a considerable increase in vessel diameter in Tlr\(^{-/-}\) mice. This increase was then confirmed and quantified using digital image analysis showing a twofold increase in vessel diameter in the Tlr\(^{-/-}\) group compared with WT mice (Fig. 3b). During the regression phase (days 10 and 14), the enhanced vessel diameter was accompanied by a decrease in the number of vessels per microscopic field (Fig. 3c). However, on day 6, before the regression phase, no differences in vessel diameter and density were observed. To substantiate these findings from two-dimensional immunohistochemical analysis, we used light sheet microscopy and three-dimensional (3D) analysis (Fig. 3d and Supplementary Movies 1 and 2). Tumours in WT mice showed a typical pro-angiogenic dense microvascular network, which was distributed through the entire tumour. Interestingly, during the rejection phase (day 10), tumours in Tlr\(^{-/-}\) mice were more heterogeneously vascularized and large tumour areas appeared to be devoid of a typical tumour-associated vascular network (Fig. 3d, lower panels). To examine whether endothelial cell death could be involved in this phenomenon, we combined CD31 staining with caspase staining. Most endothelial cells were negative for caspase in tumours of WT and Tlr3/7/9\(^{-/-}\) mice, making substantial endothelial caspase-mediated cell death during tumour rejection unlikely (Fig. 3e). We next analysed the vascular maturation level in both tumour types by using the established pericyte markers desmin and z-smooth muscle actin. In accordance with the enhanced vascular diameter found in Tlr\(^{-/-}\) mice, at day 14 intratumoural microvessels of Tlr\(^{-/-}\) mice displayed a mature vascular network with strong recruitment of desmin-positive mural cells and indicative of vessel normalization. In contrast, total loss or partial lack of pericyte coverage was observed in tumours of WT mice (Fig. 3f). Quantification of vessel maturation indicated no difference at day 10 (WT 52 % Desmin\(^{+}\) vessels; Tlr\(^{-/-}\) 59 % Desmin\(^{+}\) vessels), but a clear difference became apparent at day 14 (WT 36 % versus Tlr\(^{-/-}\) 91 %) (Fig. 3g). These data establish that the tumour regression in Tlr\(^{-/-}\) mice is accompanied by vessel normalization, whereas tumour growth in WT mice is associated with a typical tumour-associated pro-angiogenic vessel phenotype.

**Inflamed tumour cell phenotype in TLR-deficient mice.** Based on these findings, we speculated that differences in the microvascular endothelium could be further associated with a differential infiltration of the tumours by immune cells. We found that tumours in WT mice were moderately infiltrated by immune cells (Fig. 4a,b). Tlr\(^{-/-}\) mice showed a similar moderate infiltration until day 6 post injection. Post day 6, however, there was a dramatic increase in the number of tumour-associated leukocytes in Tlr\(^{-/-}\) mice. This strongly induced immune infiltrate was composed of both myeloid (CD11b, Gr-1) and lymphoid (CD4, CD8) cells. Interestingly, this change in tumour infiltration was not accompanied by changes in the composition of peripheral blood immune cell subsets (Supplementary Fig. 3a). To test whether the inflammatory growth of tumours in Tlr\(^{-/-}\) mice was indeed caused by enhanced and active migration of immune cells into the tumour tissue, we performed adoptive transfer experiments. To this end, tumour-bearing WT and Tlr\(^{-/-}\) recipient mice were injected with fluorescently labelled splenocytes derived from tumour-bearing donor mice (both WT
One day later, tumours were explanted, cleared and subjected to 3D light sheet microscopy. As shown in Supplementary Fig. 3b, a substantial number of tumour-infiltrating splenocytes was only detectable in $Tlr^{-/-}$ mice, which received adoptively transferred cells from TLR-deficient mice. These data, in agreement with the BM chimera experiment (Fig. 2), suggest that cooperative mechanisms between non-haematopoietic stroma and immune cells are responsible for the inflammatory immune cell recruitment in TLR-deficient mice.

In an attempt to better understand this inflammatory activity in tumours of $Tlr^{-/-}$ mice, we performed a pilot experiment (four mice, day 6) and analysed gene expression in tumour tissue.
using RNA microarrays. Evaluation of cytokine gene expression suggested an induction of interleukin (IL)-6, IL-24 and interferon (IFN)-γ in KO over WT animals (Fig. 4c). These candidate genes were then further investigated in a time course and on the protein level (Fig. 4d,e). In agreement with the previous findings, induction of IL-6 and IL-24 started between day 4 and day 6,
and was followed by induction of IFN-γ. Release of IFN-γ and IL-6 was also confirmed on the protein level using a model of ex vivo cultured tumour explants (Fig. 4e). Induction of IL-24 seemed to be restricted to the RNA level and was not confirmed by enzyme-linked immunosorbent assay (ELISA) analysis of tumour explant supernatants (SNs).

T-cell-dependent tumour rejection in TLR-deficient mice. In the next series of experiments, we explored the activation state, phenotype and functionality of T cells in WT versus Tlr" mice. Proliferative activity of both CD4 and CD8 T cells from tumour-draining lymph nodes was higher in TLR-deficient mice compared with WT controls (Fig. 5a). In tumours, >40% of CD4 cells in TLR-deficient mice displayed an activated phenotype, defined as CD62L⁻/⁻CD154⁺ (Fig. 5b). Similarly, the percentage of cytotoxic CD8 T cells (positive for granzyme B, CD107a or perforin) was strongly induced in TLR-deficient mice over WT mice (Fig. 5b and Supplementary Fig. 3d). In line with these findings, a substantial number of CD4 and CD8 cells from tumours of Tlr" mice, but not from WT mice, also expressed PD-1, LAG3 and Tim3 were not differentially expressed exhaustion markers and the frequency of intratumoural cytotoxic molecules, we also tested the expression of CD11b and CD154 in TLR-deficient mice. Proliferative activity of both CD4 and CD8 T cells from tumour explants (Fig. 4e). Induction of IL-24 seemed to be restricted to the RNA level and was not confirmed by enzyme-linked immunosorbent assay (ELISA) analysis of tumour explant supernatants (SNs).

Discussion

TLRs are important sensors of pathogen-associated molecular patterns and help to protect the host from foreign intruders. Engagement of TLRs links innate with adaptive immunity and results in the induction of protective immunity. This principle has subsequently been used and exploited therapeutically, making TLR agonists important components in today’s cancer immunotherapy. Conversely, and somewhat counterintuitive, the triggering of TLRs by endogenous ligands in some cases does not result in anti-tumour immunity but rather contributes to tumour progression. This can be explained by the fact that in these cases TLR activation is a continuous and chronic process, which is induced by either tumour-inducing pathogens or by endogenous TLR ligands released during cancer-related inflammation. More recently, the interconnection of the so-called microbiome with endogenous and therapy-induced anti-tumour immune responses has increasingly gained attention. In a pioneering paper using murine melanoma models, Sivan et al. identified distinct commensal microbiota, which were associated with improved endogenous anti-tumour immunity and synergized with checkpoint-blockade immunotherapy. In a model of therapy-induced anti-tumour immunity, the interaction of commensal microbiota and myeloid cells influenced response to immunotherapy. All in all, these data highlight the complex and highly context-dependent interconnection of TLR signalling and anti-tumour immunity.

Both tumour cells and non-malignant cells of the tumour host can be targets for TLR ligands. It has been described that triggering of TLR responses in tumour cells can lead to the induction of tumour cell death, which, under certain conditions, is associated with induction of anti-tumour immunity by mechanisms referred to as immunogenic or immunostimulatory cell death. Alternatively, activation of TLR on tumour cells can also promote tumour growth by inducing anti-apoptotic mechanisms, immune escape or chemoresistance.

In this study, we specifically focused on the role of the endosomal TLRs 3, 7 and 9 expressed on non-malignant host cells. To this end, we used transplantable WT tumour cell lines and TLR-deficient host mice. We show that the absence of those TLRs on host cells results in the induction of protective CD8 T-cell-mediated immunity and complete regression of WT tumour cells in multiple tumour models. Tumour regression required the cooperative activity of immune and radioreistant host cells. The anti-tumour immune response was associated with significant alterations in the tumour microenvironment such as vessel normalization, enhanced numbers and strong activation of tumour-infiltrating lymphocytes.

Until today, endogenous triggering of TLRs on host cells has been studied mainly for TLR2 and 4, whereas less is known on the endosomal TLRs 3, 7 and 9. This is partly due to the fact that the endogenous ligands, commonly referred to as DAMPs, for TLR2/4 are by far better characterized than their endosomal nucleic-acid-sensing counterparts. Examples for TLR2/4 DAMPs include HMGB1, histones, syndecans, heat-shock proteins and S100 proteins. Interestingly, effects of TLR2 and TLR4 activation by those endogenous DAMPs seem to be context-dependent...
as well. In a transplantable model of orthotopic head and neck cancer in C3H mice, the absence of TLR4 on host cells resulted in accelerated tumour growth. In the context of chemo- or radiotherapy, the expression of TLR4 and activation of MyD88 mainly on dendritic cells were crucial for efficient induction of anti-tumour immunity. Conversely, in models of melanoma and bladder cancer, TLR4 signalling, supposedly via tumour-derived heat-shock proteins and tumour-associated macrophages, accelerated the growth of lung metastases. Similarly, tumour-derived versican has been proposed as a driver of metastasis by promoting the pro-metastatic activity of myeloid cells via TLR2 (ref. 39). In contrast, the activation of TLR2 by HMGB1 during

Figure 5 | T-cell activation in lymph nodes and tumour. (a) Lymph nodes: number of proliferating cells within the tumour-draining lymph nodes (LNs) of 6- to 12-week-old C57BL/6 WT and Tlr3/7/9−/− mice was assessed at day 6 and day 10 by immunofluorescence double staining of frozen sections using αCD4 and αKi67 antibodies. Quantification of positive cells was performed by observer-assisted counting on four randomly selected ×200 magnified images per sample (n = 3). (b,c) Tumour: flow cytometric analysis of T-cell activation of tumour-infiltrating lymphocytes of 6- to 12-week-old C57BL/6 WT and Tlr3/7/9−/− mice at day 6 and day 10. (b) Activation status of CD4+ cells was analysed by staining of CD62L−/CD154+ cells. Activated CD8+ cells are determined by showing Granzyme B+ (GzmB) and CD107a+ expression. (c) Cytokine production of tumour-infiltrating CD4+ and CD8+ cells was assessed after ex vivo stimulation with αCD3/CD28 antibodies and subsequent flow cytometry staining using αIFN-γ and αTNF-α antibodies. Single values and mean (horizontal line) are shown. Results were considered significant at *P ≤ 0.05 and **P ≤ 0.005 (t-test).
brain tumour therapy was reported to contribute to anti-tumour immunity via activation of dendritic cells.

Unfortunately, and in contrast to TLR2/4, the endogenous ligands for TLR3/7/9 remain largely enigmatic. Some studies suggest that (mitochondrial) DNA, mostly in the context of severe injury and sepsis, could be implicated in pathological activation primarily of TLR9 (ref. 37). In addition, HMGB1 has been suggested as a DAMP for TLR9 (ref. 38), although it is difficult to determine whether HMGB1 or associated DNA is the recognized molecular structure.

In mice mutated for TLR3, 7 and 9, the activation primarily of TLR9 has been demonstrated. In addition, HMGB1 has been identified as a DAMP for TLR9 in a model of wound-induced skin cancer, the absence of TLR5 on leukocytes has been suggested as a driver of metastasis. In a model of wound-induced skin cancer, the absence of TLR5 on leukocytes reduced tumour prevalence in BM chimeras. Similarly, the absence of TLR7 on BM-derived cells reduced carcinogenesis in a pancreatic cancer model.

We tried to determine whether TLR deficiency on haematopoietic or non-haematopoietic radioresistant host cells was required for induction of tumour regression. Unexpectedly, only the combined deficiency of both compartments resulted in induction of anti-tumour immunity. This was somewhat surprising, as previous papers suggested a predominance of the haematopoietic cells of the host, Kim et al. identified the inflammatory versican-TLR2-macrophage axis as a driver of metastasis. In a model of wound-induced skin cancer, the absence of TLR5 on leukocytes reduced tumour prevalence in BM chimeras. Similarly, the absence of TLR7 on BM-derived cells reduced carcinogenesis in a pancreatic cancer model. Conversely, implanted prostate cancer cells grew faster in Tlr3−/− mice, although the relative contribution of BM-derived versus radioresistant cells or immune versus non-immune cells was not assessed in that study.

In our study, tumours in WT mice were mostly non-inflamed, with low numbers of tumour-infiltrating T cells, which showed little activation. In TLR-deficient tumour regressor mice, we observed clear signs of T-cell activation in both lymph nodes and tumours. This activation was accompanied by strongly enhanced numbers of T cells infiltrating tumours from TLR-deficient mice. In addition to T-cell activation itself, T-cell recruitment to the tumour tissue and anti-tumour immunity may also be influenced by vessels and stromal fibroblasts. Although the role of the so-called cancer-associated fibroblasts in tumour progression and modulation of tumour immunity is fairly well established, the influence of endothelial cells and especially pericytes on effector
cell recruitment to the tumour is only beginning to emerge. In an interesting study by Johansson et al. 48 TNF-α and IFN-γ were targeted to the tumour vasculature. Although IFN-γ primarily had anti-vascular pro-apoptotic activity, low-dose TNF-α stabilized the tumour-associated vascular network and enhanced CD8 effector T-cell activity. During the phase of tumour regression, we observed altered vessel structure and increased pericyte coverage, which could indicate a potential role of the altered vasculature in enhanced recruitment of effector T cells into the tumour microenvironment. However, initial experiments performed in this model did not suggest induced vessel activation or altered vascular permeability and leukin in TLR-deficient mice. Thus, at this stage it remains unclear whether vascular alterations and enhanced T-cell influx are mechanistically linked or represent rather independent parallel events. It is also important to note that changes in vessel structure, T-cell influx, T-cell activation and tumour regression seem to be rather parallel events in this model. All of these were observed around day 10, but were not yet present at day 6 after tumour injection.

In clinical studies and human cancer patients, mostly expression of TLR on tumour cells has been analysed and evaluated for prognostic impact and clinical relevance. Few studies investigated TLR expression on stromal cells. For example, in an observational clinical study, TLR9 on stromal cells was associated with good prognosis in oral squamous cell carcinoma (OSCC). Conversely, expression of TLR4 on stromal fibroblasts associated with good prognosis in oral squamous cell carcinoma (OSCC).59 Conversely, expression of TLR4 on stromal fibroblasts was associated with poor prognosis in colorectal cancer.64 Similar data were reported for hepatocellular carcinoma, where TLR9 expression on tumour-associated fibroblasts was associated with reduced overall survival.60 Thus, our study is one of the first to experimentally demonstrate a role for radiosensitive (stromal) cells in regulation of tumour progression via endosomal TLR signalling.

Collectively, our data uncover a novel function of endosomal nucleic-acid-sensing TLRs for the regulation of anti-tumour immune responses. The combined absence of three TLRs on both BM-derived and radiosensitive cells of the tumour host was required for tumour regression. Our data also suggest a previously unknown role for endosomal TLRs in limiting endogenous adaptive and protective T-cell responses in the tumour host.

Methods

Animals. All animal experiments were approved by the animal ethics committee of the state of North Rhine-Westphalia and carried out according to German guidelines for experimental animal welfare. Six-to-17-week-old male and female C57BL/6 WT and Tlr3/7/9−/− mice were obtained from the animal facility of the University Hospital Essen or from Harlan Winkelmann GmbH (Borchen, Germany) and housed under standard conditions in individually ventilated cage boxes. Six-to-12-week-old male and female Tlr single and Tlr double KO mice were provided by Stefan Bauer, Philipps-University Marburg.

Tumour cell culture. The murine oropharyngeal cell lines (MOPCs) were kindly provided by J. Lee (Sanford Research/University of South Dakota) and cultured in medium containing DMEM high glucose and Ham’s F12 (3:1) (Thermo Fisher Scientific, Karlsruhe, Germany) supplemented with 10% (v/v) heat-inactivated FCS (Biochrom, Berlin, Germany), 100 IU ml−1 penicillin, 100 μg ml−1 streptomycin (Thermo Fisher Scientific), epidermal growth factor (5 μg ml−1) (Biochrom), insulin (5 μg ml−1), transferrin (5 μg ml−1), cholera toxin (0.0084 μg ml−1), hydrocortisone (0.5 μg ml−1) and tri-sodium-thyronine (0.00136 μg ml−1) (Sigma-Aldrich, Taufkirchen, Germany). MOPC-EGFP cells were generated by lentiviral gene transfer using a pCL6EGWo empty vector52 and enriched to > 90% enhanced green fluorescent protein (eGFP) positivity by FACS analysis. The murine lung carcinoma cell line T47C was kindly provided by Z. Friedenlander (Hadassah Medical Center, Israel) and the murine bladder cancer cell line MB49 by K. Eswaranathan (University of Singapore, Singapore). Both cell lines were cultured in DMEM high glucose supplemented with 10% (v/v) FCS, 100 IU ml−1 penicillin, 100 μg ml−1 streptomycin and 1 mM sodium pyruvate (Thermo Fisher Scientific). All cells were routinely tested for mycoplasma contamination using the Veron Mycoplasma Detection Kit (Minerva Biolabs, Berlin, Germany).

Tumour models. Synergetic murine cell lines were either injected s.c. into the flank or into the Musculus mylohyoideus (orthoplastic injection). Tumour cells (0.5 x 104–2 x 105) were applied in PBS using a 27-gauge syringe under anaesthesia. After s.c. injection, tumour volume was measured every other day in two dimensions and tumour volume was estimated according to the following formula

\[ V = \frac{4}{3} \pi \times (\text{width})^2 \times (\text{length}) \]

After orthotopic injection, weight of the mice was measured daily. Mice were killed after a weight loss of > 20% in accordance with animal welfare guidelines.

Isolation of primary spleen cells and BM cells. For spleenocyte and lymph node cell isolation, tissue was dissociated with a plunger. BM was collected from femurs by flushing with the bone with a 23-gauge needle. Erythrocyte lysis in spleenocyte and BM was performed by applying distilled water to the cell pellet for 20 s before readjusting osmolality with 2 x PBS. Cell suspensions were passed through a 50 μm filter and cultured in RPMI-1640 (Thermo Fisher Scientific) supplemented with 10% (v/v) heat-inactivated FCS, 100IU ml−1 penicillin and 100 μg ml−1 streptomycin or in PBS. BM cells were isolated with the PanT cell isolation Kit (Miltenyi, Bergisch Gladbach, Germany) according to the manufacturer’s instructions. Purity was > 90% in all experiments.

Isolation of cells from tumour tissue. Tumour tissue was cut into 2–3 mm pieces in PBS, centrifuged at 300 g for 7 min and resuspended with 1 ml enzyme mix containing 0.2 mg ml−1 Collagenase D, 0.2 mg ml−1 Dispase 1 and 0.1 mg ml−1 DNase (all from Sigma-Aldrich). After 45 min incubation at 37°C, the cells were passed through a 100 μm filter and kept in RPMI-1640, supplemented as described above, until analysis.

Generation of BM chimeric mice. C57BL/6 WT and Tlr3/7/9−/− mice were exposed to dose (9.5–10.5 g) total body irradiation, from a 800- beam 60Co source (Philips, Germany). Donor BM cells from C57BL/6 WT and Tlr3/7/9−/− mice were isolated as described above. Recipient mice were injected intravenously with 5 x 106 BM cells in 0.2 ml PBS 1 day after irradiation. BM from WT mice was injected into TLR-deficient mice and vice versa. Receiving BM of the same genotype served as control. To achieve haematopoietic reconstitution, mice were allowed to recover for 8 weeks until 0.5 x 107 MOPC tumour cells were injected s.c. into the flank of the mice. Tumour growth was measured every other day for 1 month.

Antibody-mediated depletion. zGr-1 antibody (RB6-8C5, BioXCell, West Lebanon, USA) was used to deplete Gr-1-positive cells by intraperitoneal injection of 200 μg antibody diluted in 0.9% NaCl. Treatment started 3 days before tumour cell injection and was repeated every third day throughout the experiment. For depletion of CD8+ or CD4+ T cells, 100 μg antibody (CD8, 2.43, BioXCell and CD4, GK1.5, BioXCell) was used the same way with application intervals of 7 days. Control groups were treated with isotype antibody (rat IgG2a LTF-2, BioXCell) using the same regimen as applied for the specific antibody. Depletion was verified using peripheral blood from tail veins and flow cytometry.

Cytokine analysis of cell culture SNs. Explanted tumour pieces with a mean weight of 12 mg were cultured at 20 mg ml−1 medium for 48 h to generate tumour explant SNs. SNs were collected, centrifuged to remove cell debris and stored at − 20°C until analysis.

The murine cytokines IL-6 and IL-24 were detected using DuoSet ELISA Kits (R&D Systems, Wiesbaden, Germany) and mouse IFN-γ was detected using OptEIA Set ELISA Kits (BD Biosciences, Heidelberg, Germany) according to the manufacturer’s protocol. Absorbance at 450 nm was measured with a Synergy II microplate reader (BioTek, Bad Friedrichshall, Germany).

Flow cytometry. Cells were stained with fluorochrome-coded antibodies in PBS supplemented with 5% murine serum and 10% FCS for 20 min at 4°C. The following antibody conjugates and respective clones were used: CD11b-PE-Cy7 (clone 2A3, 2 μg ml−1, Abcam, Cambridge, UK), Ly6G-PerCP-Cy5.5 (clone 1A8, 2 μg ml−1, BioLegend, Fall, Germany), CD4-V450 (clone 2B11, 4 μg ml−1, BD Biosciences), CD4 (BV605, RM4-5, 0.25 μg ml−1, BioLegend), CD8 (AF700, 0.25 μg ml−1, BioLegend), CD26 (BV510, Mel14, 0.25 μg ml−1, BioLegend), CD154 (PE, MR1, 0.25 μg ml−1, e Bioscience), CD244.2 (APC, eBio244F4, 1 μg ml−1, e Bioscience), LAG3 (PE, eBioC987BW, 0.25 μg ml−1, eBioscience),
second antibodies were used: rat anti-CD31 (clone MEC13.3, 1:1,000), rat anti-CD3 (clone M1/70.15, 1:1,000), rat anti-CD11b (clone M1/70.15, 1:1,000; Thermo Scientific), and rat anti-CD123 (clone 2G12, 1:200, Thermo Scientific). Nuclei were visualized by 4,6-diamidino-2-phenylindole (DAKO, Hamburg, Germany). Donkey anti-rabbit Alexa594 (1:800) and goat anti-rabbit Alexa648 (1:200) (both from Thermo Fisher Scientific).

Double florescence staining of tumour tissue was performed after fixation with BD CytoPerm Cytopermit Kit (BD Biosciences) and incubation with hydrogen peroxide (DAKO, Hamburg, Germany). Slices were incubated for 60 min at room temperature with the primary monoclonal antibody in a humid chamber. Subsequently, samples were incubated with secondary antibodies for 30 min and for 10 min with AEC Single ABC and C176 Solution (Thermo Fisher Scientific). Nuclei were visualized by Shandon Instant Hematoxylin (Thermo Fisher Scientific)28. The following primary (rat- or mouse-) and soluble secondary antibodies were used: rat-γCD31 (clone MEC13.3, 1,1000), rat-γCD11b (clone M1/70.15, 1,1000; Thermo Scientific), rat-γCD3 (clone KT15, 1,000, AB德Serbio/Rio-Bad, Düsseldorf, Germany), rabbit-γDesmin (ab15200, 1,600, Abcam), rabbit-γ-rat HPRO, goat-γ-rabbit HPRO (both 1,1000, Dianaova, Hamburg, Germany), donkey-γ-rabbit-Alexa594 (1,800) and goat-γ-rabbit-Alexa648 (1,200) (both from Thermo Fisher Scientific).

For quantification of tumour-infiltrating lymphocytes and vessels, images were taken using Axioskop 2 (Carl Zeiss, Jena, Germany) and Axiovision (Carl Zeiss) software as indicated in the experiment. Cells were quantified by manual counting of individual cells of four microscopic fields taken at ×200 magnification. Quantification was done by independent investigators in a blinded manner. Quantification of perfusion was performed by manual counting of up to five fields per tumour, including 100 vessels in total, using the mean of three tumour sections per tumour (top, middle and base) of concordant distance. Vessel analysis was performed by using the BX51 (Olympus, Hamburg, Germany) and corresponding Cell P Software. For representative pictures, images were processed in the same file simultaneously in Photoshop CS5 (Adobe Systems, San Jose, USA).

**Determination of cell death.** Apoptosis of cultured cells was detected using Annexin-V:PE Apoptosis Detection Kit (BD Biosciences) via flow cytometry, according to the manufacturer’s protocol. In frozen sections of tumour tissue, total amounts of apoptotic cells were quantified by APO-DIRECT TUNEL staining (Becton Dickinson). 7-Aminoactinomycin D (AlA-D) (BD Biosciences) was used as nuclear staining. To determine the number of TUNEL-positive cells in relation to the whole tumour area, digital images taken at ×25 magnification were analysed using Image J software (NIH, Bethesda, USA).

**Tumour clearing and ultramicroscopy.** MOPC-ε-GFP cells (1 × 10⁶) were injected s.c. into WT and Tlr3⁻/⁻ mice. If vessel staining was required, 10µg of the rat-γ-mouse CD31 (clone MEC13.3, BioLegend) antibody coupled with Alexa Fluor 647 (BioLegend) was injected retrobulbarly in each mouse (20 µg ml⁻¹). Injection was carried out under anaesthesia by intraperitoneal application of ketamine and xylazine dose at 100 and 10 mg per kg body weight, respectively, and diluted in 0.9% NaCl for injection of 10 µl per g body weight. Twenty minutes later, mice were killed and perfused by injection of 20 ml 5 mM EDTA/PBS (pH 7.2) into the left ventricle of the heart followed by 20 ml 4% paraformaldehyde (PFA). Tumours were dissected and the 3DISCO protocol was used for clearing44. Briefly, tumours were incubated in tetrahydrofuran with increasing concentration (30, 50, 80, 100% (v/v)/), for 30 min each. Subsequent incubation in dibenzyl ether was performed overnight. Vascularization of the tumours was analysed using the lightsheet illumination technique of the LaVision BioTec UltraMicroscope (Olympus). Data were analysed using the Image) and Imaris 8.0.2 Software (Imaris, Cologne, Germany).

**Adoptive transfer.** Recipient WT and Tlr3⁻/⁻ mice were s.c. injected with 1 × 10⁶ MOPC-ε-GFP tumour cells 6 days before adoptive transfer. Donor mice were s.c. injected with 1 × 10⁶ MOPC-ε-GFP tumour cells 10 days before harvesting of donor splenocytes. Splenocytes of WT and Tlr3⁻/⁻ donor mice were isolated and labelled with 1 µM CellTracker Deep Red Dye (Thermo Scientific). Labelled splenocytes (1 × 10⁷) were injected intravenously into recipient mice. One day later, tumours were dissected and cleared. Infiltration of transferred splenocytes was examined by generating 3D images using ultramicroscopy.

**Statistical analysis.** Statistical significance was assessed with the Student’s t-test for comparison of two groups or analysis of variance if more than two groups were compared. Number of mice per group is indicated for each experiment and individual mice are shown for most experiments. Sample size was determined based on previous preliminary data. No animals were excluded from analysis and no explicit method of randomization was used to assign animals to experimental groups. Results were considered significant at *p < 0.05, **p < 0.005 and ***p < 0.0005. Hierarchical clustering of gene expression data was performed with R software (The R Foundation for Statistical Computing, Vienna, Austria).

**Data availability.** Microarray data supporting the findings in this article were deposited in NCBI’s Gene Expression Omnibus under the GEO Series accession code GSE92358. All other data generated or analysed during this study are included in this published article and its Supplementary Information files or available from the corresponding author upon reasonable request.

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Author contributions
J.C.K. planned and performed experiments, analysed and interpreted data, and wrote the manuscript. K.M. and G.Z. provided financial support and conceived of the study, interpreted the data and wrote the manuscript. S.B. provided financial support and most KO mice and contributed to experimental planning and data interpretation. I.H. performed experiments, analysed and interpreted data. J.C.K. planned and performed experiments, analysed and interpreted data, and wrote the manuscript. K.M. and G.Z. provided financial support and conceived of the study, interpreted the data and wrote the manuscript.

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