Caspase 9b Drives Cellular Transformation, Lung Inflammation, and Lung Tumorigenesis

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ABSTRACT

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Caspase 9 undergoes alternative splicing to produce two opposing isoforms: proapoptotic Caspase 9a and pro-survival Caspase 9b (C9b). Previously, our laboratory reported that C9b is expressed in majority of non–small cell lung cancer tumors and directly activates the NF-kB pathway. In this study, the role of C9b in activation of the NF-kB pathway in vivo, lung inflammation and immune responses, and lung tumorigenesis were examined. Specifically, a transgenic mouse model expressing human C9b in the lung pneumocytes developed inflammatory lung lesions, which correlated with enhanced activation of the NF-kB pathway and increased influx of immunosuppressive myeloid-derived suppressor cells in contrast to wild-type mice. C9b mice presented with facial dermatitis, a

Introduction

Caspase 9 is a member of the caspase family and an important mediator of intrinsic apoptosis via association with cytochrome C and Apaf-1 to form the apoptosome (1, 2). The CAS9 gene is also controversially reported as a tumor suppressor (3). Pre-mRNA produced from the CAS9 gene can be alternatively spliced by inclusion or exclusion of the four-exon cassette (exons 3,4,5,6) to produce two isoforms: proapoptotic caspase 9a (C9a) and antiapoptotic caspase 9b (C9b; refs. 4, 5). C9b mRNA lacks the exon 3,4,5,6 cassette, which encodes the large subunit and the catalytic domain, but the translated protein retains the APAF-1 and cIAP-1 association regions. C9b competes with C9a in binding to the apoptosome complex, inhibits the activation of C9a and the ability of a number of cell death agonists to induce apoptosis and the loss of cell viability (4, 5). Therefore, the

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thickened and disorganized dermis, enhanced collagen depth, and increased serum levels of IL6. C9b mice also developed spontaneous lung tumors, and C9b cooperated with oncogenic KRAS in lung tumorigenesis. C9b expression also cooperated with oncogenic KRAS and p53 downregulation to drive the full cell transformation of human bronchial epithelial cells (e.g., tumor formation).

Implications: Our findings show that C9b can directly activate NFkB pathway in vivo to modulate lung inflammation, immune cell influx, and peripheral immune responses, which demonstrates that C9b is key factor in driving cell transformation and lung tumorigenesis.

alternative splicing of caspase 9 pre-mRNA is a key step in determining the apoptotic fate of a cell (2).

In studies focused on the cellular mechanisms regulated by C9b, ablation of C9a did not produce the same biological phenotypes of enhanced cell survival, anchorage-independent growth (AIG), and tumorigenesis observed for C9b expression (6–8). These findings showed that C9b had a "gain of function" in addition to competing with C9a for the APAF-1 association. C9b was shown to activate the canonical nuclear factor-kappaB (NF-kB) pathway via direct interaction with the cellular inhibitor of apoptosis 1 (cIAP1; ref. 6).

The NF-kB pathway is the master regulator of inflammation and is tightly linked to cancer as a pro-survival and oncogenic pathway (9). Suppression of NF-kB in myeloid cells or tumor cells induces tumor regression (10). In relation to non–small cell lung cancers (NSCLC), both the Baldwin and Jacks laboratories demonstrated that the NF-kB pathway cooperates with oncogenic RAS to enhance tumorigenesis in mice (11, 12). Induction of airway epithelial NF-kB during allergic inflammation was shown to cause airway hyperresponsiveness by increasing lymphocytes and neutrophils in the bronchioalveolar lavage (BAL) fluid (13, 14). Moreover, sustained NF-kB signaling induced chronic inflammation and emphysema by four months of age. By 11 months, NF-kB activated mice develop lung adenomas, which recruit M2-polarized macrophages and regulatory T lymphocytes providing a pro-tumorigenic microenvironment (15).

Due to our finding that C9b induced activation of the NF-kB pathway and high C9b expression occurs in >78% of NSCLCs (6–8), our laboratory developed a C9b transgenic mouse model and found that C9b expression in lung pneumocytes was sufficient to activate the NF-kB pathway and induce the infiltration of lymphocytes in the lungs linked to increased IL6 levels and facial dermatitis. Furthermore, C9b expression induced lung adenocarcinoma formation in aged mice and cooperated with oncogenic KRAS to induce lung tumorigenesis demonstrating a major pro-tumor function for this caspase 9 splice variant.

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Materials and Methods

Detailed materials and methods and the list of key reagents and resources (Supplementary Table S1) can be found in Supplementary Data file.

Cell culture and in vitro assays

HBEC-3KT (obtained from ATCC) and HBEC3KT-KP cells with KRAS G12V expressing (/K) \pm p53 shRNA downregulated (/P; a gift from Drs. John Minna and Jerry Shay; ref. 16) were cultured as described (10). HBEC3KT-KP cells were authenticated via Western immunoblotting and qPCR for p53 and KRAS G12V. MLE12 cells (obtained from ATCC) were cultured as described (17). These cell lines were transduced with pLEX304-eGFP, –wild-type (WT) C9b, or -AT/ GG Mut C9b lentivirus treated with 8 µg/mL polybrene (Sigma) and selected with blasticidin. MLE12/K and HBEC-3KT/K cells are generated by transducing cells with pLenti6-KRAS^{G12V} (Addgene). Plate colony formation assays (cell survival) and soft agar colony formation assays (AIG) were undertaken as described (7). All cell lines were used within 6 passages from receipt and thawing. Cell lines were tested every 2 months for Mycoplasma (Universal Mycoplasma Detection Kit, ATCC) throughout the study starting 2 months after thawing cells received from ATCC and immediately for other cell lines herein described. Parental cell lines were authenticated by short tandem repeat profiling. Cells were treated with Bay 11-7082 (Sigma, $0-10$ µmol/L range) at the indicated doses.

Western immunoblotting and immunoprecipitation

Western immunoblotting was accomplished as described (6) using the following primary antibodies: anti–caspase 9 (Enzo Life Sciences); anti-FLAG (Sigma); RIP1, anti-IkBa, anti-Myc tag, c-Myc, anti-HSP90, and anti- β -actin (Cell Signaling Technology); anti-K-63 Ub (Abcam). Secondary antibodies were horseradish peroxidase-conjugated antirabbit IgG and anti-mouse IgG antibodies (Cell Signaling Technology). RIP1 immunoprecipitation was performed as described (6).

Mouse models

All mouse studies were conducted under the University of South Florida—approved Institutional Animal Care and Use Committee (IACUC) proposals, 8191M and 8230R (Assurance number: A-4100–01) and the James A. Haley Veterans' Hospital-approved approved IACUC protocol, #4433V.

Caspase 9b transgenic mouse model

Lung specific transgenic mouse expressing FLAG-tagged human Caspase 9b was designed and generated by the VCU Transgenic/ Knockout Mouse Shared Resource Core. Specifically, the open reading frame expressing human Caspase 9b was inserted between rabbit betaglobin introns downstream of the human surfactant protein C (SPC) promoter sequence. Genotyping PCR was performed using forward (5'-ATA TCT CCG CTG AAG CCG-3') and reverse (5'-ATC TTC GGA AGC TGG CAG T-3') primer sets.

Lung inflammation of C9b mice was semiquantitatively graded [Score 0–3 as 0 (none) and 3 (severe)] on the basis of the number and the size of peribronchial/bronchiolar/alveolar lymphoid aggregates and diffuse infiltrates observed on hematoxylin and eosin (H&E) images at 10x magnification blinded to the genotypes as described in (18). The alveolar area was calculated by morphometric analyses of trichrome stained images using ImageJ (NIH).

Cohorts of line 2 and line 4 C9b mice were compared with littermate transgene negative/control mice for the development of skin dermatitis from 3 months to 23 months of age. Mice with deep ulcerative skin lesions penetrating through both the epidermis and dermis were identified during routine daily health checks blinded to the genotype of the mice as described (19). Mice with a simple alopecia/hair loss or a focal superficial mild skin lesion or with fighting wounds were not counted.

KRAS lung tumor mouse model

Seven- to eight-week-old $KRAS^{\text{G12D L/+}}$ mice (20) were intratracheally instilled with 2.5×10^7 purified lentiviral particles. Mice were euthanized at 32 weeks post instillation.

Mouse tumor xenograft model

5-week-old female NOD/SCID (Envigo) mice were injected subcutaneously into hind flanks with 2.5×10^6 HBEC-3KT/KP cells (mixed in 50% Matrigel solution) expressing GFP ($n = 10$), WT C9b ($n = 20$), or AT/GG Mut C9b ($n = 10$). On day 40 postimplantation, mice with palpable tumors were euthanized, and tumors were excised for histologic analyses.

Histology and IHC

Mouse organs (lung, liver, spleen, and skin) and tumors were fixed in 10% formalin and embedded in paraffin for histologic and IHC analyses as described (21, 22) using the following primary antibodies: anti-FLAG, -Caspase 9, - I κ B α , p65, -pERK, -Myc Tag (Cell Signaling Technology); -SOX2 (Sigma); -Gr1 (Ly6G/6C), IL6 (Novus Biologicals); -HMGA2 (ThermoFisher, Waltham, MA); -p63, and -Synaptophysin (Santa Cruz Biotechnology). Gomori Trichrome staining (Richard-Allan Scientific) and Alcian blue-PAS staining (Statlab) were performed following the manufacturers' instructions. HMGA2 score was calculated by multiplying staining intensity score (0, no stain; 1, weak; 2, moderate; 3, strong) with % positivity score (0, <5%; 1, <10%; 2, <50%; 3, >50% tumor cells).

Flow cytometry

Flow cytometric analysis of the immune compartment in murine lung and spleen was performed as described (23). The myeloid-derived suppressor cell (MDSC) subsets within the spleen and lung were characterized by surface staining in PBS containing 5% FBS (FACS buffer) with: Gr1 (BUV395), F4/80 (BV711), CD19 (PerCP-Cy5.5), from BD Biosciences, CD11b (BV605), CD11c (Alexa488), IA/IE (Alexa647), and B220 (APC-H7) from BioLegend, incubated in 4° C for 1 hour, then washed twice with FACS buffer, and finally fixed in PBS containing 1% paraformaldehyde for flow cytometric analysis. Dead cells were excluded using the Zombie Violet Fixable Viability Kit (BioLegend) following the manufacturer's protocol. Cells were acquired on a BD FACSymphony A5 and data were analyzed with FlowJo Version 10.0 software.

Statistical analysis

Statistical differences between the two groups were determined by a two-tailed, unpaired Student t test. Statistical differences among \geq 3 groups (adjusted P values) were determined by ANOVA followed by Tukey multiple comparison test. All tests were performed using GraphPad Prism 9 software. P values < 0.05 were considered significant.

Results

C9b activates the NF-kB pathway and cooperates with oncogenic KRAS to induce cellular transformation

The expression of C9b in NSCLCs enhances tumor phenotypes for NSCLC cells, which required the activation of the NF-kB pathway (6).

High C9b expression is also observed in >78% of human NSCLC tumors. The question remained as to whether C9b expression was a consequence of tumorigenesis or an important oncogenic signaling factor in cooperating with oncogenes to facilitate cellular transformation. To study the functional effects of C9b expression toward full malignancy in lung epithelial cells, we used the genetically defined CDK4/hTERT-immortalized normal human bronchial epithelial cells (HBEC-3KT) \pm common lung cancer oncogenic changes (p53 downregulation via shRNA (P) and ectopic expression of oncogenic KRAS $(KRAS^{G12V}; K; Fig. 1A and B)$. Expression of WT C9b, but not GFP control or the AT/GG Mut C9b (a mutant that retains APAF-1 association and antiapoptotic capabilities, but cannot bind cIAP-1 and activate the NF-kB pathway; ref. 6), in HBEC-3KT cells increased RIP1 K^{63} -ubiquitination (Fig. 1C) and decreased IKB α (Fig. 1D) demonstrating activation of the NF-KB pathway. The effect on $I\kappa B\alpha$ was blocked by the addition of the IKK inhibitor, BAY11–7082 (Fig. 1D).

Functionally, C9b expression in HBEC-3KT cells increased cell survival/colony formation compared with GFP control and the AT/GG Mut C9b (Fig. 1E), but failed to promote AIG (Fig.1F). When KRAS^{G12V} is expressed in these cells (HBEC-3KT/K), C9b

dramatically and significantly enhanced in vitro cellular transformation (e.g., AIG) compared with GFP and AT/GG Mut C9b (Fig. 1G). WT C9b further cooperated with oncogenic KRAS to further enhance AIG when there was concomitant p53 downregulation (HBEC-3KT/KP) (Fig. 1G): cell lines which showed insignificant AIG in the GFP control similar to previous reports (Fig. 1G; refs. 16, 24). Although significantly lower, AT/GG Mut C9b also increased the AIG capacity of these cell lines compared with GFP control, suggesting a possible role for NF-kBindependent and antiapoptotic abilities of C9b in cell transformation. C9b expression in HBEC-3KT/KP cells also increased invasion through Matrigel (Fig. 1H). Similar effects were observed in MLE12 mouse lung epithelial cells (SPC⁺), a cell line derived from a transgenic mouse model with SPC promoter driven SV40 large T antigen (Supplementary Fig. S1; ref. 17). Specifically, C9b expression cooperated with oncogenic KRAS, driving AIG and invasion. The enhanced cell survival and induction of in vitro cell transformation (e.g., AIG enhancement/induction) induced by C9b were further shown to require the NF-kB pathway as BAY 11–7082 blocked these phenotypes in MLE12 cells (Supplementary Fig. S1). Enhanced AIG by C9b expression in A549 (lung cancer cell line

Figure 1.

C9b activates NF-KB in HBEC-3KT cells and cooperates with KRAS activation and p53 loss driving AIG and invasion in vitro and tumor growth in vivo. A-D, Western blot analyses with indicated antibodies of HBEC-3KT, -3KT/K (KRAS^{G12V}), -3KT/P (shTP53), and -3KT/KP cells expressing GFP, WT, or AT/GG mutant (Mut) C9b (with a Myc-tag). C, RIP1 immunoprecipitation (IP) followed by immunoblotting with K⁶³-Ub antibody shows increased RIP1 K⁶³-Ub by WT C9b in HBEC-3KT cells (in growth supplement free media). In D, HBEC-3KT cells were treated with 0, 3, 10 μ mol/L Bay 11-7082 for 15 hours. WCL, whole-cell lysates. The empty and filled triangles denote C9a and C9b proteins, respectively. E, Clonogenic assays of parental HBEC-3KT cells expressing GFP, WT, or Mut C9b. The graph shows numbers of clonal foci per well on D10. F, Bright field images (10x) of HBEC-3KT cells expressing GFP, WT, or Mut C9b grown in soft agar on D45. G, Number of colonies per low power field (LPF) view grown in soft agar on D45 (*, $P \le 0.0002$). H, Number of invaded HBEC-3KT/KP cells expressing GFP, WT, or Mut C9b through Matrigel at 22 hours per LPF view. I, SCID Tumor take rate of HBEC-3KT/KP cells expressing GFP, WT, or Mut C9b on day 30 postimplantation (top) and H&E, Myc-tag, and c-Myc IHC images (bottom). Data are means \pm SD; $n = 3$ each. Adjusted P values are determined by ANOVA Tukey multiple comparison test.

harboring a $KRAS^{G12C}$ mutation, Supplementary Fig. S2) was suppressed by Bay 11–7082 (Supplementary Fig. S2A and S2B), which were further reduced by combined treatment with MEK162 (Ras/MEK/ERK pathway inhibitor; Supplementary Fig. S2B), supporting cooperation of oncogenic KRAS and NF-kB pathways. These cells with reduced C9b expression showed dramatic reduction in NF-kB activation and IL6 production and were less sensitive to Bay 11–7082 in clonogenic survival assays (Supplementary Fig. S2C–S2F). Finally, C9b expression in HBEC-3KT/KP cells drove subcutaneous tumor growth in SCID mice (Fig. 1I), the hallmark of full cellular transformation. Specifically, 30% of mice $(n = 6/20)$ implanted with HBEC-3KT/KP cells expressing C9b developed tumors by 30 days post injection compared with none for those with GFP or 10% for AT/GG Mut C9b $(n = 1/10)$ of a significantly smaller size than WT C9b; Fig. 1I; Supplementary Fig. S3). These tumors showed expression of Myc-tagged C9b (Fig. 1I). Thus, C9b activates the NF-kB pathway and cooperates with oncogenic KRAS and p53 downregulation to drive full cellular transformation.

Expression of caspase 9b in mouse pneumocytes activates the NF-kB pathway in vivo

As C9b enhanced tumorigenic phenotypes and cellular transformation, we next examined whether these biological phenotypes translated in vivo. To study the role of C9b in lung pathophysiology and tumorigenesis in vivo, we generated transgenic mouse lines expressing human C9b under the SPC promoter that confers lung epithelium-specific expression in alveolar type II pneumocytes and bronchioalveolar cells (refs. 25, 26; Fig. 2A). Among the independent founder lines, line 2 and line 4 mice showed germline transmission and both high C9b mRNA and protein expression restricted to the lung (Fig. 2B–D). These mice were also viable and fertile with no gross differences compared with transgene negative control mice.

Because C9b directly activated NF-kB pathway in primary lung epithelial and NSCLC cell lines (6), we tested the effects of C9b expression in mouse lung for induction of this pathway. First, C9b expression in lung epithelial cells was confirmed by IHC staining with FLAG and C9 antibodies (Fig. 2E). Importantly, lungs of these mice

Figure 2.

The SPC-driven C9b expression in mouse lung epithelium activates NF-kB pathway. A, A schematic diagram of a C9b transgene construct driven by human SPC promoter. Among the C9b transgene positive founder lines (B, PCR), line 2 and 4 C9b⁺ mice showed high levels of C9b transgene expression in lungs both for mRNA (C, real time rt-PCR, relative ratio of C9b/GAPDH) and protein (D, Western blot). E, Representative images of IHC staining with indicated antibodies (10x). Note a Flag-tagged C9b expression in lung epithelium, a decreased level of IκΒα and an increased level of p-p65 in line 2 and line 4 mice compared with control mice. Insets (40x) in p65 images show the exclusion of p65 from nuclei in control but not in C9b mice. Inset in IkBa shows its strong positivity in immune aggregates.

showed decreased levels of IKB α , increased levels of phospho-p65, and enhanced nuclei localizations of p65 compared with those of littermate transgene negative control mice (Fig. 2E). Thus, C9b mice demonstrated NF-kB pathway activation in mouse lungs showing the translation of previous in vitro findings to the in vivo setting.

Expression of human caspase 9b in mice pneumocytes induces lung inflammation and immune cell infiltration

Previously NF-kB activation induced in all lung epithelium was shown to cause an emphysema phenotype and infiltration of immune cells (15), and thus, we next histologically evaluated the effect of C9b expression on lung inflammation. Compared with control mice, C9b mice, even though C9b is only expressed in a subset of lung epithelial cells, demonstrated an emphysema phenotype in the presentation of enlarged alveolar ducts and alveolar sacs (Fig. 2E; Fig. 3A and B). The average size of the alveolar sac was significantly larger in C9b mice (6–8 months old) for both line 2 (2093 \pm 360.8, n = 5, P = 0.0283) and line 4 (2223 \pm 441.3, $n = 5$, $P = 0.0193$) compared with control mice $(832.8 \pm 82.26, n = 5; Fig. 3A and B)$. Fibrosis and fibrotic thickening of small airway walls and alveoli was evident with increased collagen deposition in C9b mice by trichrome staining (Fig. 3A). In addition, increased perivascular and peribronchiolar lymphoid aggregates and diffuse inflammation were observed in the lungs of C9b mice (Fig. 3C and D). Both Line 2 (average lung inflammation score 1.5 ± 0.189 , $n = 7, P = 0.0026$) and line 4 (1.78 \pm 0.278, $n = 9, P = 0.0010$) C9b mice showed increased numbers and sizes of immune aggregates and diffuse inflammation in lung parenchyma compared with littermate controls (0.57 \pm 0.165, n = 14, Fig. 3C and D). Analysis of the immune cell populations increased in the lungs of C9b mice included $Gr1⁺$ neutrophils, monocytes, macrophages, and MDSCs as well as $CD3\varepsilon^+$ T cells, B220⁺ B cells, CD44⁺ memory T cells, OX40⁺ effector T cells, Mac-2⁺ histiocytes (Supplementary Fig. S4A). Flow cytometry analysis of cells prepared from whole lungs and spleens showed increased effector/effector memory $CD4^+$ and $CD8^+$ T-cell populations in lung and spleen of C9b mice compared with control mice as defined by $CD44⁺$ and $CD62L⁺$ (Supplementary Fig. S4B). The most significant changes were the increased percentage of MDSCs (Gr1⁺ and CD11b⁺) in the lungs and spleens of C9b mice compared with control (Fig. 3E

Figure. 3.

Lungs of C9b mice show emphysema, fibrosis, and immune infiltration. A, H&E and trichrome staining show enlarged alveoli and increased peribronchovascular collagen deposits in line 2 and line 4 mice compared with control. Average area of alveoli (B , $n = 5$ per group) is measured using Image J from trichrome stained slides avoiding large vessels and bronchi (A, bottom). C and D, Scoring systems for immune aggregates and infiltrates (C, 10X) and average lung inflammation score for each group ($n = 14$ for control, 7 for line 2, and 9 for line 4 mice, D). E and F, Increased % of MDSCs (Gr1+/CD11b+) in lung and spleen of C9b mice ($n = 3$) compared with Control ($n = 3$) by FACS analysis. Data are means \pm SEM. Adjusted P values are determined by ANOVA Tukey multiple comparison test for **B** and **D** and by unpaired t-test for F.

and F). Thus, C9b mice demonstrated significant lung inflammation indicative of constitutive activation of the NF-kB pathway through a presentation of remodeling of lung epithelium with emphysema, fibrosis, increased lymphoid aggregates, and increased infiltration of MDSCs.

Expression of human caspase 9b in mice pneumocytes is linked to skin dermatitis

One unexpected phenotype observed among C9b mice was the development of severe skin dermatitis (Fig. 4A) as the C9b transgene was not expressed in the keratinocytes, fibroblasts, and melanocytes in the skin of C9b mice (Supplementary Fig. S4C). Specifically, a cohort of Line 2 ($n = 74$) and line 4 ($n = 36$) C9b mice were compared with littermate control mice ($n = 68$) from 3 months to 23 months of age. Although skin dermatitis is common around the neck and body of female C57BL/6 strain mice (19), C9b mice developed severe ulcerative lesions often on the face with deep scabs warranting euthanasia, affecting both male and female mice (Fig. 4; Supplementary Fig. S5). Kaplan–Meier dermatitis-free survival curves show significantly increased severe dermatitis development both in line 2 ($P < 0.001$, Log-rank test) and line 4 $(P = 0.0029)$ compared with control mice (Fig. 4B). Histologic analysis of the skin lesions showed the epidermal hyperplasia, serocellular crusting, and dermal inflammatory infiltrates (Fig. 4C) as well as thickened and disorganized dermal collagen layers (Fig. 4C). Thus, C9b mice with C9b expression in lung pneumocytes showed systemic inflammatory phenotypes.

To begin to determine the mechanistic underpinnings of how C9b mice develop skin inflammatory phenotypes distal from lungs

where it is expressed, the levels of cytokines/chemokines in blood circulation were assessed. Various cytokines/chemokines such as CXCL13, IL16, IL1Ra, CCL5, CCL2, G-CSF, CXCL1, and TIMP1 were all upregulated in the plasma of C9b mice with dermatitis compared with control (Supplementary Fig. S6A). Previously, we identified CXCL1 and CXCL2, downstream targets of NF-kB pathway that were associated with atopic dermatitis (27), as genes regulated by C9b (6). Here, we observed increased expression of Cxcl1 and/or Cxcl2 in lungs of C9b mice that were suppressed by IkBa super repressor (Supplementary Fig. S6B–S6D). The NF-kB pathway is also known to activate the proinflammatory cytokine IL6, which is a known regulator of the survival and expansion of MDSCs as well as skin dermatitis and wound healing (28, 29). We found that reduced C9b expression and Bay 11–7082 treatment in A549 cells suppressed IL6 secretion and IL6 mRNA expression as well as CXCL1 expression (Supplementary Fig. S2). IL6 levels in the blood of control versus C9b mice were significantly increased [both line 2 (204.8 \pm 51.40, $n = 23$, $P = 0.0018$) and line 4 (82.65 \pm 17.11, $n = 11$, $P = 0.0057$] as well as significantly increased plasma IL6 levels (Supplementary Fig. S6E) compared with control groups $(20.72 \pm 8.78 \text{ pg/ml})$. When mice were grouped on the basis of the presence and absence of skin dermatitis, IL6 levels were increased compared with control in line 2 (298.2 \pm 97.12, $n = 10$, $P = 0.0192$) and line 4 (72.66 \pm 9.816, $n = 8$, $P = 0.0012$) C9b mice without dermatitis (Supplementary Fig. S6F). These results show that the increase of plasma IL6 is not caused by dermatitis, but by C9b expression in the lung driving not only a localized, but also a more systemic inflammatory response. Because

Figure 4.

C9b mice develop severe skin dermatitis on the face and the back skin requiring euthanasia (A). B, Kaplan– Meier dermatitis-free survival curves show early onset of dermatitis in line 2 ($n = 74$) and line 4 mice ($n = 36$) compared with control ($n = 68$, P value: Log-rank test). C, H&E and trichrome staining of skins show the significantly thickened dermal area with increased collagen (blue in trichrome) in dermatitis lesions of C9b mice.

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these are only correlative observations, identification of causal cytokines/chemokines driving skin phenotypes of the C9b mice require further studies.

C9b mice develop lung cancer with aging

To investigate whether C9b expression in lung epithelium promotes lung tumorigenesis, aged C9b mice (>21 months old) were compared with age-matched control mice. Consistent to the low spontaneous lung cancer incidents (1% in males, 3% in breeding females, and 0% in virgin females) in C57BL/6J mice (strain background of C9b mice), none of the control ($n = 0/20$) mice developed primary lung cancers (30). In stark contrast, various histologic subtypes of lung cancers were observed in C9b mice $(n = 4/20$ for line 2 and $n = 2/13$ for line 4; Fig. 5A and B) in addition to emphysema, fibrosis, and severe inflammation associated with the increased infiltration of macrophages and lymphocytes (Supplementary Fig. S7A). Histologic subtypes include squamous cell carcinoma ($n = 1$, p63-positive) and carcinoid tumor $(n = 1,$ synaptophysin-positive), and adenocarcinomas. Adenocarcinoma was the most common subtype observed ($n = 3/20$ for line 2 and $n = 2/13$ for line 4), which were negative for adenocarcinoma differentiation marker Nkx2.1, but positive for acidic/neutral mucin by Alcian blue-PAS staining (Fig. 5C). Unexpectedly, all these tumors were negative for SPC and FLAG-tagged C9b expression while surrounding normal epithelium were positive for SPC and FLAG (Fig. 5C). Most of these tumors were positive for SOX2 that were normally expressed in conducting, but not in respiratory airways. Thus, our data suggest that these tumors are derived from conducting airway cells or bronchioalveolar stem cells (BASCs) at the bronchioalveolar duct junction (BADJ) that can generate both conducting and respiratory airway cells including type II pneumocytes $(SPC⁺)$

To examine the modulating effect of C9b on the tumor microenvironment, the number of MDSCs in the C9b mice lung tumors were examined as MDSCs (Gr1⁺/CD11b⁺) have been shown to promote tumor growth (31). C9b mice both with $(Gr1⁺$ cells per 20x field, 47.5 \pm 11.7) or without lung tumors (57.1 \pm 20.5) show significantly increased $Gr1^+$ cells infiltrating into lung parenchyma compared with control (7.26 \pm 4.14; Fig. 5D and E), suggesting that elevation of MDSCs is not consequence of the tumor formation, but is more likely

Figure 5.

C9b mice develop lung tumors with aging. Macroscopic (A) and microscopic lung tumors observed in C9b mice over 21 m old of age are summarized in B. *, 1 mouse developed a carcinoid tumor and adenocarcinoma. C, Histologic analyses including H&E (a-c), Alcian Blue-PAS (d), and IHC staining with indicated antibodies (e-i) of lung tumors identified adenocarcinoma (a, d, g-i), squamous cell carcinoma (SCC; b and e), and carcinoid tumors (c and f). Images are taken at 10x magnification. Inset in b is at 40x magnification. D and E, Gr1⁺ cells (D, IHC) in the lung per low power field (20x, 3 views per mouse) were counted in control (n = 9) and C9b mice with (red, $n = 5$) or without (n = 11) lung tumors (E). Data are means \pm SEM. Adjusted P values are determined by ANOVA with Tukey multiple comparison test.

due to C9b expression. In addition, type II pneumocytes, not tumor cells, express IL6 (Supplementary Fig. S7B). These data show that C9b enhances the recruitment of MDSCs into the lung allowing for a protumor microenvironment.

C9b cooperates with KRAS in lung tumorigenesis

On the basis of our finding that C9b will cooperate with oncogenic KRAS to induce cellular transformation and provides a protumorigenic microenvironment, we hypothesized that C9b expression will cooperate with oncogenic KRAS to drive lung tumorigenesis. To test this hypothesis, KRAS G12D L/ $+$ mice were intratracheally injected with modular lentiviral particles (32) to simultaneously express Cre recombinase and a coding sequence of either GFP ($n = 12$), WT C9b $(n = 10)$, or AT/GG Mut C9b $(n = 11;$ Fig. 6A and B). At 32 weeks post injection, lungs from GFP control mice showed tumor foci with ERK activation, a sign of KRAS activation (Fig. 6C). WT C9b expression led to the development of larger sized and increased tumor multiplicity compared with GFP controls as assayed by percent replacement of normal lung parenchyma by tumor cells based on H&E and pERK

signal (Fig. 6C and D; GFP 4%, WT C9b 37%). Mut C9b also significantly enhanced these parameters, but to a lesser extent than WT C9b (Mut C9b 21%). Interestingly, tumors driven by WT C9b showed dramatically stronger HMGA2 expression in a high percentage of tumor cells compared with tumors from control GFP or Mut C9b expression (Fig. 6E), which is a hallmark of a shift to poorly differentiated and more invasive tumors. We also observed a case of distal metastatic lung tumor with desmoplastic reaction (Fig. 6F) with only WT C9b. In addition, numerous clusters of immune cell foci were observed in or near tumors in WT C9b group. C9b expression in the context of KRAS activation in vivo also increased the number of $Gr1⁺$ cells infiltrated into lungs compared with GFP (36.8 \pm 21.2 vs. 4.9 \pm 4.6, $P = 0.0022$) and to Mut C9b (8.3±4.0, $P < 0.0001$; Fig. 6G; Supplementary Fig. S7). Mut C9b did not increase Gr1^+ cells in lung compared with GFP ($P = 0.5607$; Fig. 6G). To assess whether this increase in Gr1⁺ cells correlates with increased MDSCs, we analyzed % of Gr1⁺ CD11b⁺ cells in the lungs by FACS analyses ($n = 2$ each; Supplementary Fig. S8). Lungs with WT C9b showed a higher percentage of MDSCs compared with GFP or Mut C9b. Thus, WT C9b

Figure 6.

C9b cooperates with KRAS activation driving lung tumorigenesis. $KRAS^{GI2D} L/+$ mice with intratracheal injection (B) of lentiviral particles containing GFP-, WT-, or Mut C9b-Cre(T2a)Luc (A) developed lung tumors (C) at 32 weeks post injection. IHC staining (C) with indicated antibodies shows the expression of Myc-tagged WT or mutant C9b, ERK activation (pERK), and HMGA2 in tumors and the presence of Gr1⁺ cells in the lung (black arrowheads). Orange arrows point to the clusters of immune cells observed. The percent replacement of normal lung parenchyma by tumor cells based on H&E and pERK signal $(n = 12$ for GFP, 10 for WT C9b, and 11 for Mut C9b, D), the HMGA2 positivity score (E), and the number of Gr1+ cells per view (20X magnification, G; $n = 4$ for GFP, 6 for WT C9b and Mut C9b, average of 3 views per mouse for E and G) in each group are shown. F, Metastasis observed in the liver (H&E, 10x magnification) in a mouse with WT C9b expression. Data in D, E and G are means ± SEM combined from 2 independent intratracheal injection experiments. Adjusted P values are determined by ANOVA Tukey multiple comparison test.

cooperated with KRAS activation, driving more poorly differentiated lung tumors with an increase in infiltrating MDSCs (or $Gr1^+$ high immune infiltrates), while Mut C9b also moderately enhanced lung tumorigenesis, but without increasing MDSCs. These data suggest a role for NF-kB activation by C9b via cIAP1 in recruiting or the expansion of MDSCs, which would provide a more pro-tumor microenvironment.

Discussion

In this study, C9b expressed in the lung pneumocytes of mice activated the NF-KB pathway in vivo, induced lung inflammation, the infiltration of immune cells (e.g., immunosuppressive MDSCs conducive to tumor growth), affected the peripheral immune system, and induced systemic inflammation. Importantly, C9b induced lung tumor formation in aging mice and cooperated strongly with oncogenic KRAS to enhance both cellular transformation and lung tumorigenesis. HMGA2 expression was dramatically increased in tumors from the oncogenic KRAS mice by WT C9b. HMGA2 is upregulated in lung cancer, a hallmark of higher grade tumors, highly expressed in poorly differentiated and invasive cancers, and inversely correlated with survival (33, 34). Thus, C9b is driving the formation of high grade lung tumors, which required the ability of C9b to associate with cIAP1 as expression of a well-characterized, structurally intact mutant of C9b lacking the ability to bind cIAP1 showed significantly less ability to enhance lung tumorigenesis in cooperation with KRAS. Therefore, the presented study corroborates the previous findings of both the Jacks and Baldwin Laboratories in the cooperation of the NF-kB pathways in oncogenic KRAS-induced lung tumorigenesis and induction of higher grade tumors (11, 12), but importantly, provides a physiologic and mechanistic link to the alternative splicing of CAS9 to produce C9b and activate this system. This finding translated to the ability of C9b to induce cellular transformation in cooperation with oncogenic KRAS and oncogenic KRAS/TP53 downregulation. Thus, the C9b/cIAP1 interaction site may be a plausible personalized therapeutic target for NSCLC expressing higher levels of C9b without the global effects on suppressing the immune system such as the concerns for general NF-_{KB} inhibitors.

As we observed the spontaneous induction of lung tumors in the C9b mice in stark contrast to WT mice, C9b may be, by definition, a protooncogene, explaining the enhanced C9b expression observed in a large percentage of NSCLC tumors. On the other hand, many of the spontaneous tumors from the aging C9b transgenic mice showed expression of SOX2, and all lacked expression of SPC and our C9b transgene (expressed via an SPC promotor). SOX2 is a transcription factor playing essential roles in lung development and maturation. SOX2 is expressed in the trachea, airway/bronchiolar epithelium and the conducting airways of the adult lung, but is completely absent from the respiratory airways (35). Previously, SOX2 was proposed to differentiate KRASG12C driven tumors as derived from Clara cell antigen 10 $(CC10)^+$ cells, not from SPC⁺ cells (36, 37). Our data suggest that these tumors are either derived from transformed CC10 expressing cells or BASCs at the BADJ coexpressing CC10 and SPC. SPC (and thus, C9b under an SPC promotor) is downregulated as tumors progress (38). This possibly explains the loss of C9b expression in the spontaneous lung tumors observed in the C9b mice if C9b is a protooncogene. Conversely, C9b expression alone does not confer in vitro cellular transformation to the HBEC-3KT cells, and thus, the role of C9b may be as an extrinsic paracrine role in modulating the microenvironment allowing tumor growth to occur in a more spontaneous manner. Additional studies are needed to determine whether C9b is truly a protooncogene or acts to promote a strong protumorigenic microenvironment.

Mechanistically, C9b plays NF-kB–dependent and –independent functions as we have previously reported (6). NF-kB activates myriads of downstream target genes such as c -Myc and $C/EBP\alpha$ and various cytokine/chemokines (39, 40). Thus, C9b is expected to alter many of these genes leading to the inflammatory phenotypes observed. In fact, we observed that reduced C9b expression (via siRNA targeting C9b) downregulated C/EBPa (Supplementary Fig. S9A). Bay 11–7082, NFkB pathway inhibitor, suppressed C/EBPa level in A549 (Supplementary Fig. S9B). These data suggest that $C/EBP\alpha$ may be downstream of C9b activation of the NF-kB pathway (as one of the many transcription factors regulated by NF-kB such as c-Myc). In addition, the NF-kB– independent, antiapoptotic function of C9b did significantly enhance tumor area/multiplicity induced by oncogenic KRAS as shown by expression of the Mut C9b, which retains the ability to bind APAF1, block activation of C9a, and inhibit intrinsic apoptosis. Our findings suggest that loss of intrinsic apoptosis in cells such as would be observed by loss of the CAS9 gene would support cooperation with oncogenes to drive cell transformation and lung tumor formation. Currently, the role of the CAS9 gene as a tumor suppressor is controversial with Lowe and colleagues reporting the loss of the CAS9 enhances the cellular transformation of murine embryonic fibroblasts induced by oncogenic manipulations (3); a finding that was not confirmed by Strasser and colleagues (41). Our findings suggest that in certain contexts, C9a will act as a tumor suppressor, at least in constraining the ability of tumors to transition to higher grades. This study also found that expression of the Mut C9b significantly enhanced in vitro cell transformation in cooperation with oncogenic KRAS, albeit to a much lesser extent than WT C9b that retains the ability to also drive the NF-kB pathway. On the other hand, Mut C9b only induced full cellular transformation in cooperation with oncogenic manipulations in one instance, and thus, a full tumor suppressive role for the antiapoptotic functions of the CAS9 gene requires more exploration in different cellular and oncogenic contexts.

Another novel finding was the systemic nature of inflammation that was observed by early activation of the NF-kB pathway in only lung pneumocytes induced by C9b expression. Indeed, systemic inflammation was observed as shown by increased serum levels of various cytokines (Ex. CXCL1 and IL6) and facial dermatitis. NF-kB family of transcription factors have been implicated in the pathogenesis of several autoimmune diseases including rheumatoid arthritis, inflammatory bowel disease, type I diabetes, systemic lupus erythematosus and multiple sclerosis (42). Interestingly, mouse models with altered NF-kB pathway have shown skin dermatitis phenotypes. For example, IkBa -/- mice was reported to develop widespread severe skin dermatitis with increased $Cd11b⁺$ macrophages/monocytes in spleen (43) and $cRel-/-p50-/-p65+/-$ mice developed spontaneous dermal and intestinal inflammation in association with chronic neutrophilia (40). Clinical links are also reported between facial rashes and lung inflammation as to asthma and viral pneumonia induced by COVID-19 (44–46). Whereas the clinical manifestation of the facial rash was suggested as a plausible link to the later development of asthma, our findings suggest the opposite in that lung inflammation, not yet manifested or detectable by current clinical tests, is prognostic for later development of asthma. In these cases, clinical testing for systemic IL6 may be suggested to treat the lung inflammation to block the development of chronic lung disease. Of note, this study is mainly showing correlative observations between C9b, NF-kB activation in the Type II pneumocytes, and systemic inflammation linked to facial dermatitis. The

identification of causal cytokines/chemokines driving skin phenotypes of the C9b mice require further studies, but our findings suggest a very strong interplay between organ systems and systemic inflammation.

In conclusion, this study found that C9b induces the activation of the NF-kB pathway in the lungs, the infiltration of lymphocytes into the lungs, and lung inflammation linked to increased IL6 levels, systemic inflammation, and facial dermatitis. C9b induced lung tumor formation in aged mice demonstrating that C9b is either a protooncogene, promotes a highly pro-tumor microenvironment, or both, leading to spontaneous tumor formation. C9b also cooperated with oncogenic KRAS to induce lung tumorigenesis, which identified the C9b/cIAP1 interaction as a plausible new target for personalized treatment of NSCLC presenting with high C9b expression.

Authors' Disclosures

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Authors' Contributions

M. Kim: Formal analysis, supervision, validation, methodology, writing–original draft. N.T. Vu: Data curation, formal analysis, validation, investigation, methodology.

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