EXTENDED REPORT

Choline kinase inhibition in rheumatoid arthritis

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ABSTRACT

Objectives Little is known about targeting the metabolome in non-cancer conditions. Choline kinase (ChoKα), an essential enzyme for phosphatidylcholine biosynthesis, is required for cell proliferation and has been implicated in cancer invasiveness. Aggressive behaviour of fibroblast-like synoviocytes (FLS) in rheumatoid arthritis (RA) led us to evaluate whether this metabolic pathway could play a role in RA FLS function and joint damage.

Methods Choline metabolic profile of FLS cells was determined by 1H magnetic resonance spectroscopy (1H MRS) under conditions of ChoKα inhibition. FLS function was evaluated using the ChoKα inhibitor MNS8b (IC50=4.2 μM). For arthritis experiments, mice were injected with K/BxN sera. MNS8b (3 mg/kg) was injected daily intraperitoneally beginning on day 0 or day 4 after serum administration.

Results The enzyme is expressed in synovial tissue and in cultured RA FLS. Tumour necrosis factor (TNF) and platelet-derived growth factor (PDGF) stimulation increased ChoKα expression and levels of phosphocholine in FLS measured by Western Blot (WB) and metabolomic studies of choline-containing compounds in cultured RA FLS extracts respectively, suggesting activation of this pathway in RA synovial environment. A ChoKα inhibitor also suppressed the behaviour of cultured FLS, including cell migration and resistance to apoptosis, which might contribute to cartilage destruction in RA. In a passive K/BxN arthritis model, pharmacologic ChoKα inhibition significantly decreased arthritis in pretreatment protocols as well as in established disease.

Conclusions These data suggest that ChoKα inhibition could be an effective strategy in inflammatory arthritis. It also suggests that targeting the metabolome can be a new treatment strategy in non-cancer conditions.

INTRODUCTION

Synovial inflammation, hyperplasia and joint destruction are hallmarks of rheumatoid arthritis (RA). Resident fibroblast-like synoviocytes (FLS) contribute to synovial inflammation by producing inflammatory mediators and recruiting and activating immune cells. FLS in the intimal lining and pannus are the major effectors of cartilage damage through production of extracellular matrix-degrading enzymes, such as metalloproteinases (MMP) and cathepsins. New targeted and rationally designed disease-modifying agents that modify FLS behaviour are needed to complement current therapies. However, the molecular mechanisms that regulate FLS behaviour in RA are poorly understood and represent a major obstacle for developing therapeutic interventions that modulate these functions.

Metabolomics provides a global assessment of a cellular state within the context of its immediate environment, taking into account genetic regulation, altered kinetic activity of enzymes, and changes in metabolic pathways. A metabolic strategy might provide insights that define mechanisms underlying disease, and permit development of new treatment strategies. However, few studies have addressed metabolic changes in RA, and none specifically in FLS.

In oncology, the tumour metabolome is beginning to be characterised, and such studies have already suggested several new targets and biomarkers. Among other changes, tumours display elevated phospholipid levels characterised by increases in the levels of phosphocholine (PCho) and total choline-containing metabolites (tCho) together with decreases in the glycerophosphocholine (GPC)/PCho ratio. Elevated PCho levels are partially attributed to increased activity of ChoKα, the enzyme that catalyses the first step in CDP-choline pathway and is essential for phosphatidylcholine (PtdCho) biosynthesis, the major phospholipid in eukaryotic membranes that is also required for the increased cancer cell proliferation, tumour progression and invasion. High levels of ChoKα expression and activity are often associated with malignant transformation, invasion and metastasis in some human cancers. Thus, this enzyme was recently proposed as a new target for cancer therapy and has been recognised as a prognostic marker in various human cancers. However, very little is known about ChoKα role in other non-cancer cells and diseases.

Mitogen-activated protein kinase (MAPK) and phosphoinositide 3-kinase (PI3K)/Akt are signalling pathways that regulate FLS function in RA, including effects on MMP expression and synoviocyte growth and survival. Of interest, selective inhibition of choline kinase attenuates MAPK and PI3K/Akt signalling. These considerations led us to perform an assessment of choline metabolism in RA FLS to determine if the ChoKα activity regulates their pathogenic behaviour and to determine the suitability of ChoKα as a therapeutic target.

RESULTS

ChoKα expression and choline metabolite profile in RA synovium

We initially determined ChoKα protein expression in synovial tissues using immunohistochemistry. ChoKα is highly expressed in osteoarthritis (OA) and RA synovium (figure 1A–C) with especially...
prominent staining in the synovial intimal lining (figure 1D) although infiltrating cells in the sublining were also positive in some samples (figure 1E, F). The initial metabolomics profile of the synovium was determined using $^{1}$H magnetic resonance spectroscopy ($^{1}$HMRS). Some RA samples clearly showed very high total choline-containing metabolites, but overall PCho levels and choline metabolite profiles of the whole RA and OA synovia were not significantly different (figure 1G–I).

**ChoKa expression and activity in RA fibroblast-like synoviocytes**

Because ChoKα was highly expressed in intimal lining cells, we evaluated its expression and regulation in cultured primary FLS. Expression of ChoKα was confirmed by Western Blot (WB) analysis (n=5 per group) (figure 2A), with similar levels in RA and OA cell lines. To determine relative levels of enzyme activity in OA and RA cell lines, we used $^{1}$HMRS to detect choline-containing compounds profile in cell extracts. Interestingly, the MRS spectra of these metabolites were similar to tumour cells, which display elevated phospholipid levels characterised by increases in PCho and total choline-containing metabolites together with a decrease in the GPC/PCho ratio <1 (a phenomenon known as the ‘GPC-to-PCho switch’). In RA and OA FLS, PCho was the predominant metabolite and the GPC/PC ratio was <1, suggesting that ChoKα is active in FLS cell lines (figure 2B). Although a few RA FLS showed higher PCho levels than OA FLS (figure 2B), choline-containing metabolites were similar in OA and RA FLS (figure 2C). We also determined if normal FLS and normal primary fibroblasts from other tissues had ChoKα activity similar to OA/RA FLS. Although we could

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**Figure 1** Choline kinase (ChoKα) expression in arthritic synovium. Osteoarthritis (OA) (A) and rheumatoid arthritis (RA) synovium (B) were immunostained with a ChoKα antibody. IgG was used as a negative control (C). (D) High magnification of ChoKα immunostaining showing a homogeneous staining in intimal lining in both OA and RA synovium sample. (E and F) ChoKα immunostaining of sublining in an OA (E) and RA (F) synovium sample. (G and H) methyl proton resonances of Cho-containing metabolites ($^{1}$HMRS spectral region 3.20–3.24 ppm) and (I) quantification of choline-containing metabolites from 5 OA and 5 RA synovium samples and glycerophosphocholine (GPC/PCho) ratio. $^{1}$HMRS, $^{1}$H magnetic resonance spectroscopy.
Figure 2  Choline kinase (ChoKα) expression in fibroblast-like synoviocytes (FLS). (A) ChoKα expression in 5 osteoarthritis (OA) and 5 rheumatoid arthritis (RA) FLS by WB and quantification. (B) 1H MRS spectra of choline-containing metabolites from 1 representative RA and 1 OA FLS samples. (C) Quantification of choline-containing metabolites from RA and OA FLS samples (n=10). (D) 1H MRS spectra of choline-containing metabolites (methyl proton resonances) from normal primary fibroblast: 14D7 and GM0321 are dermal fibroblasts; HPF and I90 are pulmonary fibroblasts. (E) Representative choline-containing metabolites profile from normal primary fibroblasts. 1H MRS, 1H magnetic resonance spectroscopy; HPF, human pulmonary fibroblasts; WB, Western Blot.
only evaluate a limited number of those cell lines, PCho was also the predominant metabolite in normal FLS and pulmonary fibroblasts but not in dermal fibroblast (figure 2D, E). Thus, our data suggests that ChoKα activity varies between normal primary fibroblasts, and might regulate fibroblast functions in other types of fibroblasts such as pulmonary primary fibroblasts.

**Effect of inflammatory mediators on ChoKα**

To determine whether inflammatory mediators implicated in RA regulate ChoKα expression, we stimulated RA FLS with tumour necrosis factor (TNF), interleukin-1β (IL-1), platelet-derived growth factor (PDGF), epithelial growth factor (EGF) or lipopolysaccharide (LPS) for 36 h and assayed ChoKα protein by WB. ChoKα expression increased 2-fold after TNF, IL-1, PDGF or EGF stimulation, whereas LPS had no effect (figure 3A). Both PDGF and TNF increased PCho levels, but only PDGF decreased the GPC/PCho ratio, suggesting a more specific increase of ChoKα activity after PDGF exposure, whereas TNF might also activate other metabolic pathways, such as PtdCho degradation via phospholipase (PL) A2 and phospholipase B (PLB) which would increase the levels of GPC and, as a consequence, the GPC/Pcho ratio remains constant (figure 3B–D).

**Effect of ChoKα inhibition on FLS function**

To determine if the choline metabolite profile in FLS is secondary to ChoKα activation, we used ¹HMRSS in the presence or absence of a ChoKα inhibitor. The selective competitive ChoKα inhibitor, MN58b, was used in these and other...
MN58b inhibits proliferation of cancer cells in vitro with an IC$_{50}$ of 1–10 μM, and displays therapeutic activity against human tumour xenografts in vivo. After incubating RA FLS with MN58b using a dose chosen based on previous reports (5 μM), levels of PCho decreased and GPC/PCho ratio increased, further suggesting the role of ChoKα in choline metabolism in FLS (figure 3E, F). As ChoKα is involved in tumour cell proliferation, we then determined whether ChoKα inhibition interferes with FLS cell growth in vitro. RA FLS were pretreated with MN58b at various concentrations and then cultured in the presence of PDGF for 3 days or 6 days for Methyl Thiazolyl Tetrazolium (MTT) assay and BrdU assay, respectively. MN58b significantly decreased cell viability (figure 3G) and cell proliferation (figure 3H) in a concentration-dependent manner. This effect was most likely due to a reduced cell proliferation because there was no change in apoptosis in the MN58b-treated cells, as determined by release of histone-associated DNA fragments (see online supplementary figure S1A). We also examined the effect of the ChoKα inhibitor on G1 phase cell cycle regulators, comparing the effect of MN58b with a panPI3K inhibitor (INK631) that has been shown to inhibit cell proliferation as well. As shown in figure 3I, MN58b impaired the increase in phospho-Rb and cyclinD1 after PDGF stimulation to a greater extent than the panPI3K inhibitor, suggesting that cells were arrested in G0/G1. Partial silencing of ChoKα levels by siRNA (60%), only slightly abolished the effect on cell cycle regulation, suggesting that low levels of ChoKα activity might be sufficient to maintain a proliferative phenotype in these cells as MN58b renders a much more potent reduction of ChoK activity in several cell systems than the reduction achieved by siRNA.

ChoKα regulates p-AKT and p-MAPK activation, MMP expression, migration and cell survival

Selective inhibition of ChoKα attenuates MAPK and PI3K/AKT signalling in tumour cells. These pathways also regulate synoviocyte function in RA FLS. Figure 4A shows that phosphorylation of AKT (ser473, but not thr308), and ERK1/2, were markedly reduced by ChoKα inhibition in FLS. MN58b also decreased the expression of several MMP after TNF stimulation (figure 4B) and reduced cell migration in a migration-scratch assay (figure 4C, D). We finally evaluated whether ChoKα inhibition induces oxidative stress-induced apoptosis. FLS were challenged with 0.3 mM H$_2$O$_2$, which are conditions that induce apoptosis rather than necrosis. Pretreatment with MN58b (5 μM) induced more apoptotic cell death after H$_2$O$_2$.
Choline kinase (ChoKα) inhibition decreases severity of inflammatory arthritis in mice

We determined whether MN58b affects arthritis severity in the K/BxN passive serum transfer model, which is dependent on innate but not adaptive immunity. Based on protocols used in oncology, we injected the inhibitor daily for the first 5 days after administration of arthritogenic serum. As shown in figure 5A, MN58b-treated mice had significantly lower clinical scores from day 2. Histopathological analysis at day 7 showed markedly reduced inflammatory cell infiltration, joint destruction and cartilage damage in MN58b-treated mice compared with vehicle-treated controls (figure 5B). Confirming functional inhibition of ChoKα in the joints, 1HMRS of joint extracts showed increased PCho in arthritic joints that was abrogated in MN58b-treated joints (figure 5C, D). Importantly, MN58b treatment prevented the onset of arthritis and also successfully suppressed joint swelling in mice if treatment was initiated in established disease (figure 5E). To evaluate the influence of ChoKα inhibition on synovial inflammatory mediators, we determined relative expression of selected genes and protein levels of selected cytokines in joints from these mice on day 7 after serum transfer. mRNA for IL-1β, IL-6, MMP3 and MMP13 were significantly lower in MN58b-treated mice (figure 6A). ELISA analysis confirmed the reduction of IL-1β and IL-6 in MN58b-treated joints (figure 6B). WB analysis of joint extracts from a second group of mice on day 5 after KxB/N sera injection (day 5 scores were 11.75±1 and 5.5±0.5, p<0.01) also showed decreased phosphorylation of AKT and ERK in MN58b-treated mice (figure 6C, D). ChoKα expression was similar in control and phosphate buffered saline (PBS) or MN58b-treated arthritic joints (figure 6C).

DISCUSSION

In this study, we identified ChoKα as a potential therapeutic target in RA FLS that could contribute to the aggressive phenotype of these cells in vivo. Importantly, ChoKα inhibition by a selective small molecule decreased FLS migration and proliferation as well as abrogated joint inflammation and damage in a murine model of RA. Small molecule inhibitors that inhibit key signalling molecules in RA have the potential to improve efficacy and to overcome some limitations of antibody-based therapeutic approaches, such as parental treatment and cost. Based on these data, we propose that inhibiting ChoKα could complement current RA therapies by suppressing pathogenic FLS behaviour. Although ChoKα inhibitors could potentially target other cell types, current studies only assess their effect on FLS.

![Figure 5](https://example.com/figure5.png)

Figure 5  Choline kinase (ChoKα) inhibition treatment successfully abrogated joint inflammation and destruction. WT mice were injected with 150 μL serum from adult K/BxN mice on day 0. MN58b was administrated every day from day 0 to 5 at 3 mg/kg. (A) Ankle thickness and clinical score in PBS-treated animals (black circles, n=5) and MN58b-treated animals (black squares, n=5) injected with 150 μL of K/BxN serum on day 0. Values are means±SD. ***p<0.001. (B) Histological scores for joint inflammation, erosion and cartilage damage in PBS-treated and MN58b-treated mice (5 mice/group) on day 7 after serum transfer. (C) Representative 1HMRS spectra (methyl proton resonances of choline-containing metabolites) and (D) quantification of choline-containing metabolites in joints from naive and arthritic joints at day 7 with or without daily MN58b treatment. (E) WT mice were injected with 150 μL serum from adult K/BxN mice on day 0. MN58b was administrated every day from day 4 to 7 at 3 mg/kg. Shown clinical score in PBS-treated animals (black circles, n=5) and MN58b-treated animals (black squares, n=5). Values are means±SD. *p<0.05, **p<0.01. 1HMRS, 1H magnetic resonance spectroscopy; PBS, phosphate buffered saline; WT, wild type.
Many signalling pathways activated under inflammatory and hypoxic conditions have profound effects on cellular metabolism to support cell growth and survival. The study of cancer cell metabolism has successfully identified cancer-specific metabolic changes that provide new therapeutic targets. Other cell types, for instance, lymphocytes are subject to major metabolic challenges upon activation and, therefore, adopt specific metabolic programmes to adapt to changing environmental conditions. However, there are little data about targeting the metabolome in non-cancer conditions or regarding metabolic changes in FLS. We postulate, based on our results, that ChoKα and choline metabolism are activated in inflamed joints, and that targeting the metabolome can be a new treatment strategy in inflammatory diseases. Although much is known about the role of ChoKα in tumorigenesis and cancer cell proliferation, very little information is available about its activity in normal cells or in inflammatory diseases. However, there is little data about targeting the metabolome in non-cancer conditions or regarding metabolic changes in FLS. We postulate, based on our results, that ChoKα and choline metabolism are activated in inflamed joints, and that targeting the metabolome can be a new treatment strategy in inflammatory diseases. Although much is known about the role of ChoKα in tumorigenesis and cancer cell proliferation, very little information is available about its activity in normal cells or inflammatory diseases.

Figure 6 Choline kinase (ChoKα) inhibition treatment decreases inflammatory mediators in the KxB/N model. (A) PBS and MN58b-treated WT mice were injected with 150 μL of K/BxN serum on day 0 and they were sacrificed at day 7. Clinical scores at day 7 were 11±1.2 and 3±1.2, respectively (p<0.001). RNA isolation of naive and arthritic joints at day 7 with or without daily MN58b treatment, were prepared and analysed for the expression of the indicated genes. Results are expressed as means of 4 mice/group±SD. (B) PBS and MN58b-treated WT mice were injected with 150 μL of K/BxN serum on day 0 and they were sacrificed at day 5. Clinical scores at day 5 were 11.75±1 and 5.5±0.5, respectively (p<0.01). Joint protein from naive and arthritic mice of PBS and MN58b-treated mice was extracted and analysed by ELISA and (C) by WB for the presence of the indicated proteins. (D) Quantitative analysis of western blots (arbitrary densitometry units) after normalising results to total protein. PBS, phosphate buffered saline; WB, Western Blot; WT, wild type.

Basic and translational research

We also show that inhibiting this pathway suppresses inflammatory arthritis in the passive KxB/N model. This model is dependent on FLS44 as well as other innate immune cell types, such as macrophages, neutrophils and mast cells. Unfortunately, CHKA-deficiency is embryonic-lethal and CHKAF/FP mice are not available to study in the specific cell types that confer protection from arthritis. Although most of these studies are focused in the ChoK1 isoform, the recent discovery of the human ChoKβ isoform raises the possibility that this newly identified kinase might also be relevant. However, ChoK activation was not able to induce tumour growth under conditions, while ChoK1 does.45 46 Furthermore MN58b showed a much higher specificity against ChoK1 (IC50=5 μM) than against ChoKβ (IC50=107.5 μM), this is 21.5 times more potent against ChoK1 than ChoKβ isoform. These observations suggest that the effect we detect is mostly through ChoKβ.

Unique metabolomic profiles have been identified in the serum of patients with several diseases. For instance, the serum metabolic fingerprint in established RA was clearly different from that of healthy controls.47 Lactate and lipids were important discriminators of inflammatory burden. Of interest, a number of metabolites, such as choline, were also found to contribute strongly to the correlation. Other studies have also determined changes in phospholipid metabolites in serum of RA patients.9 The phospholipid composition of the synovial fluid (SF) has also been recently described.47 SF from patients with RA had higher content of total phospholipids, major phospholipid classes and phospholipid species than control SF. These data suggest altered phospholipid metabolism in RA and that cells such as FLS, which mediate the synthesis and release of phospholipids,19 increase their production and modify their classes under inflammation.

Taken together, our in vitro and in vivo studies suggest that choline metabolism is activated in RA FLS under pro-inflammatory conditions, and that selectively blocking ChoKβ might be beneficial in inflammatory arthritis by suppressing FLS functions, including migration and resistance to apoptosis. Therefore, the data provide a rationale for strategies that inhibit ChoKβ as a therapeutic approach in RA.

MATERIAL AND METHODS

Mice

KRN T-cell receptor transgenic mice were a gift from Dr D Mathis and Dr C Benoist (Harvard Medical School, Boston, Massachusetts, USA) and Institut de Génétique et de Biologie Moléculaire et Cellulaire (Strasbourg, France). Mice were on C57Bl/6 background. Mice used in these experiments were 8–12 weeks old. All animal protocols received advance approval for the use of laboratory animals. The protocol for these studies was approved by the Committee on the Use of Laboratory Animals of the University of Texas Health Science Center at San Antonio (UTHSCSA) and National Institutes of Health Grant P30 CA54174 (CTRC at UTHSCSA).

Reagents

MN58b (1,4-(4′-Bis-((4-(dimethylamino)pyridinium-1-yl) methyl)-diphenyl) butane dibromide) was obtained from JCL. MN58b inhibits proliferation of cancer cells in vitro with an IC50 of 1–10 μM, and its specificity has been addressed in previous reports.20 43 46 INK631, a pan PI3K inhibitor was obtained from Intellikine (La Jolla, California, USA). Concentrations for MN58b and INK631 were chosen based on previous reports. Cytokines and PDGF-BB were obtained from R&D Laboratories (Minneapolis, Minnesota, USA).

Preparation of synovium and synoviocytes, and cell lines

Synovium and FLS were obtained from patients undergoing total joint replacement or synovectomy who meet the 1987 revised American College of Rheumatology criteria for seropositive RA or patients with OA as previously described.50–52 Human pulmonary fibroblasts (HPF) cell line was obtained from ScienceCell Research Laboratories. 190, 14D7 and GM0321B cell lines were donated by Ben Yu’s laboratory.

Choline-containing metabolites profiling using 1H NMR

RA and OA FLS and synovium were isolated and prepared for the 1H NMR analyses as described.53–54 MRS spectra were acquired at 16.4 T (700 MHz) on a Bruker Avance spectrometer (Bruker BioSpin, Billerica, Massachusetts, USA) equipped with a TCI cryoprobe and high-throughput robotics. One-dimensional 1H NMR spectra were acquired using at least 512 scans and 32 dummy scans, 32 K data points, and a spectral width of 9.8 kHz. Excitation sculpting pulse sequence was implemented to suppress the water signal. All the MRS datasets were processed using MetaboLab55 in the MATLAB programming environment (MathWorks, Natick, Massachusetts, USA). MRS spectra were normalised based on the probabilistic quotient normalisation method. Assignment of MRS resonances and metabolite quantification were performed using Chenomx nuclear magnetic resonance (NMR) Suite and other available databases.27–29 For clarity, only the choline-containing metabolite resonances generated by the methyl protons of PCho, GPC and Cho (MRS spectral region between 3.20 and 3.24 ppm) are shown in the figures. We quantified the metabolite concentrations of intracellular fibroblast cell lines and synovium tissue extracts, and the results are presented as relative concentrations of choline-containing metabolites. Ratio of choline-containing metabolites (PCho/total choline and GPC/PCho) are also included.

Statistical analysis

Data are expressed as means±SD. The analysis used unpaired Student’s t test for comparing two groups, and analysis of variance (ANOVA) for multiple group comparisons. Results were considered significant if p<0.05.

Acknowledgements

This work was supported by grants from the National Institutes of Arthritis and Musculoskeletal and Skin (MG: 1K08AR064834), the National Institute of Allergy and Infectious Diseases (GSF; R01AI070555) and Ministerio de Economía y Competitividad (JCL: SAF2011–29699, RD12/0036/0019). Support for the NMR facility was provided by the University of Texas Health Science Center at San Antonio (UTHSCSA) and National Institutes of Health Grant NCI P30 CA54174 (CTRC at UTHSCSA).

Contributors

MG and ES-L designed the project, performed most of the experiments and analysed the data. MG also wrote the manuscript; AL and ST performed and interpreted MRS analysis. RG-C performed IHC staining. MK helped to interpret the data and in manuscript preparation. JCL provided MN58b, participated in the design of some experiments, helped to interpret the data and in manuscript preparation. GSF designed the study, analysed the data, supervised the overall project and wrote the manuscript.

Competing interests

MG, ES-L, ST, JCL, MK and GSF have filed a patent application.

Provenance and peer review

Not commissioned; externally peer reviewed.

REFERENCES

Supplementary Figure 1. Choka inhibition regulates cell death. (A) On day 3, apoptosis was determined by histone associated DNA fragmentation. There was no effect on apoptosis under these conditions. (B) FLS were exposed to 0.3mM H$_2$O$_2$ for 6 hours with or without pretreatment with MN58b (5µM). Apoptosis was determined by histone associated DNA fragmentation. (C) FLS were exposed to 0.3mM H$_2$O$_2$ for 6 hours with or without pretreatment with MN58b (5µM). Representative images after 6 hours of treatment. Quantification (as the average of number of dead cells in three different fields from 3 different RA FLS cell lines) of apoptotic cells after 0.3 mM (D) or 0.15 mM (E) H$_2$O$_2$ with or without pretreatment with MN58b (5µM). Representative of 3 different experiments.
Supplementary Figure 2 Scheme of choline metabolism, its regulatory pathways and downstream effectors. PCho: Phosphatidylcholine; DAG: Diacylglycerol; Pchol: phosphocholine; PA: Phosphatidic Acid; LPA: Lysophosphatidic acid; CDP-Cho: cytidine 5′-diphosphocholine; GPC: Glycerophosphocholine. Cho: Choline.
**Supplementary methods**

**Immunohistochemistry.** Paraffin sections prepared from RA and OA synovial tissue were incubated overnight at 4°C with mouse anti-human ChoKα antibody [58] at a 1:50 dilution. Citrate buffer (S1700; Dako) was used at 96°C for 40 minutes for antigen retrieval. The signal was developed using diaminobenzidine and sections were counterstained with hematoxylin.

**ELISA.** IL-1β, IL-6 and MMP3 has evaluated by DuoSet enzyme-linked immunosorbent assay (ELISA; R&D Systems, Minneapolis, MN), following the manufacturer's protocol.

**MTT assay.** For the MTT assay, 3 × 10^5 FLS/well were plated into 96-well plates in 10% FBS/DMEM. After 24 hours, the medium was replaced with low-serum medium (0.1%
FBS/DMEM) for 24 hours for synchronization. On day 0, medium was replaced with 1% FBS and cells were treated with ChoKα inhibitor at the indicated concentration or with PBS for 1 hour. PDGF or medium alone was added to the appropriate wells. The experiment was performed in triplicate wells. Cell viability was estimated on day 3 after incubation with MTT for 4 hours and was read at 550 nm with a spectrophotometer.

**BrdU assay.** For the BrdU assay, 3 × 10³ FLS/well were plated into 96-well plates in 10% FBS/DMEM. After 24 hours, the medium was replaced with low-serum medium (0.1% FBS/DMEM) for 24 hours for synchronization. On day 0, medium was replaced with 1% FBS and cells were treated with ChoKα inhibitor at the indicated concentration or with PBS for 1 hour. PDGF or medium alone was added to the appropriate wells. The experiment was performed in triplicate wells. Cell growth was estimated on day 6 after incubation with BrdU for 24 hours and was read at 550 nm with a spectrophotometer (Cell Proliferation ELISA kit by Roche Applied Science, Mannheim, Germany).

**Cell Survival and Apoptosis Assays.** For cell survival and apoptosis assays, 1 × 10⁶ FLS/well were plated onto 6-well plate in 10% FBS/DMEM, then after 24 hours was replaced with starving medium (0.1% FBS/DMEM) for 24 hours for synchronization. Cells were incubated with MN58b inhibitors for 1 hour, then cells were treated with 300µM H₂O₂ for 6 hours and viability was determined in H₂O₂ treated cells by phase contrast light microscopy. Apoptosis was determined using a Cell Death Detection ELISAPLUS kit (Roche Applied Science, Mannheim, Germany).

**Migration scratch assay.** FLS were seeded onto 24-well plates and allowed to come to confluence. Cells were incubated with MN58b for 1 hour. A single scratch wound was induced through the middle of each well with a sterile pipette tip. Cells were
subsequently exposed to PDGF (10ng/ml). SFC migration across the wound margins from 36h was assessed, photographed and measured by ImageJ.

**Real-time quantitative PCR (qPCR).** FLS, synovium and joints were collected. For FLS, cells were collected after FLS stimulation. Total RNA was extracted with Trizol (Invitrogen) and reverse-transcribed with random hexamers and Superscript II Kit (Invitrogen). qPCR was performed with SYBR Green PCR Master Mix Kit (Applied Biosystems, Foster City, CA). The relative amounts of transcripts were compared to those of HRPT and normalized to untreated samples by the $\Delta\Delta^{\text{Ct}}$ method.

**Western blot analysis.** FLS, synovium or joints were disrupted in lysis buffer (PhosphoSafe™, Novagen, Gibbstown, NJ) containing a protease inhibitor cocktail. Proteins were separated by SDS-PAGE and transferred to a nitrocellulose membrane. Blots were probed with antibodies against ChoKα (Proteintech), phospho-ERK1/2, phospho-AKT, ERK1/2, phosphoRb and Rb (Cell Signaling Technology, Danvers, MA), and cyclin D1 and total AKT (Santa Cruz) Biotechnology Inc, Santa Cruz, CA). Horseradish peroxidase-conjugated anti-IgG (Cell Signaling Technology, Danvers, MA) was used as secondary antibody. Membranes were developed using a chemiluminescence system (ECL detection reagent: Amersham Life Science, Aylesbury, UK). Densitometry analysis was done by using Quantity One 1-D analysis software (Bio-Rad).

**Serum transfer and arthritis scoring.** Sera from arthritic adult K/BxN mice were pooled and recipient mice were injected intraperitoneally (i.p.) with 150 µl of K/BxN serum on day 0. Clinical arthritis scores were evaluated as described [59]. MN58b (3mg/kg) was injected daily i.p. beginning on day 0 (for 5 days) or day 4 after serum administration.
**Histology analysis.** Joints were fixed in 10% formalin, decalcified in 10% EDTA for 2-3 weeks and paraffin embedded. Sections were prepared from the tissue blocks and stained with hematoxylin and eosin (H&E) and O-safranin. A blinded semiquantitative scoring system was used to assess synovial inflammation, extra-articular inflammation, bone erosion and cartilage damage (0-5 scale), as previously described [59].

Reference: