Musculoskeletal Pathology

Characterization of the KRN Cell Transfer Model of Rheumatoid Arthritis (KRN-CTM), a Chronic Yet Synchronized Version of the K/BxN Mouse

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In this study, a chronic yet synchronized version of the K/BxN mouse, the KRN-cell transfer model (KRN-CTM), was developed and extensively characterized. The transfer of purified splenic KRN T cells into T cell-deficient B6.TCR.C\textsuperscript{a−/−}H-2\textsuperscript{b−/−} mice induced anti-glucose 6-phosphate isomerase antibody-dependent chronic arthritis in 100% of the mice with uniform onset of disease 7 days after T cell transfer. Cellular infiltrations were assessed by whole-ankle transcript microarray, cytokine and chemokine levels, and microscopic and immunohistochemical analyses 7 through 42 days after T cell transfer. Transcripts identified an influx of monocytes/macrophages and neutrophils into the ankles and identified temporal progression of cartilage damage and bone resorption. In both serum and ankle tissue there was a significant elevation in interleukin-6, whereas macrophage inflammatory protein-1 \(\alpha\) and monocyte chemotactic protein-1 were only elevated in tissue. Microscopic and immunohistochemical analyses revealed a time course for edema, synovial hypertrophy and hyperplasia, infiltration of F4/80-positive monocytes/macrophages and myeloperoxidase-positive neutrophils, destruction of articular cartilage, pannus invasion, bone resorption, extra-articular fibroplasia, and joint ankylosis. The KRN cell transfer model replicates many features of chronic rheumatoid arthritis in humans in a synchronized manner and lends itself to manipulation of adoptively transferred T cells and characterizing specific genes and T cell subsets responsible for rheumatoid arthritis pathogenesis and progression. (Am J Pathol 2010, 177:1388–1396; DOI: 10.2353/ajpath.2010.100195)

Rheumatoid arthritis (RA) is characterized by chronic inflammation of the distal joints with hypertrophy and hyperplasia of the synovial epithelium, increased synovial fluid volume, and radiographic evidence of bone erosion. Infiltration of inflammatory cell types such as monocytes/macrophages, neutrophils, fibroblasts, T cells, and dendritic cells into the synovial, subsynovial, and peri-articular tissues leads to complement activation, production of pro-inflammatory chemokines and cytokines, and eventual destruction/remodeling of cartilage and bone. Indeed, it is the influx of monocytes/macrophages that mediates and amplifies this tissue damage, thereby maintaining the clinical signs of inflammation.\(^1,^2\) This process begins when chemokines drive monocyte emigration out of the blood,\(^3–8\) and is amplified when activated macrophages produce additional cytokines once they enter the joint.\(^9–13\) The extent of macrophage infiltration and their activation state is known to correlate with joint pain and the general inflammatory status of the patient, and most therapies currently available for RA decrease the number of macrophages in synovial tissue.\(^14–16\) Despite this, a complete picture of how inflammatory cells and their mediators contribute to the stepwise pathogenesis of RA has not been established, and the reliance on rodent models for obtaining this information is substantial.

Murine models of arthritis are essential components of basic research and preclinical testing of disease-modifying antirheumatic drugs. One of these models, the K/BxN murine model, exists in two different forms. In the first form, KRN TCR transgenic mice are bred to nonobese diabetic mice and the resultant F\(_1\) (K/BxN) mice develop a severe, destructive arthritis by 4 to 5 weeks of age.\(^17,18\) However, this model is not synchronized and the disease onset is not uniform across the mice.

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In the second form, serum transferred from K/BxN mice into normal recipient mice induces arthritis due to the presence of arthritogenic antibodies specific for the ubiquitously expressed glycolytic enzyme glucose 6-phosphate isomerase (GPI). Arthritis in this second, K/BxN setously expressed glycolytic enzymeglucose 6-phosphate isomerase (GPI). Arthritis in this second, K/BxN setously expressed glycolytic enzyme is induced due to the presence of arthritogenic antibodies specific for the ubiquitously expressed glycolytic enzyme glucose 6-phosphate isomerase (GPI). Arthritis in this second, K/BxN setously expressed glycolytic enzyme glucose 6-phosphate isomerase (GPI). Arthritis in this second, K/BxN setously expressed glycolytic enzyme glucose 6-phosphate isomerase (GPI).

Materials and Methods

Animals

KRN TCR Tg mice on a C57BL/6 background were originally obtained from Mathis and Benoist as described. K/BxN mice were generated by crossing KRN mice with nonobese diabetic females. The B6.G7 mouse strain was initially obtained from Mathis and Benoist as described. KRN TCR Tg mice on a C57BL/6 background were originally obtained from Mathis and Benoist as described. KRN TCR Tg mice on a C57BL/6 background were initially obtained from Mathis and Benoist as described. KRN TCR Tg mice on a C57BL/6 background were generated by crossing KRN males with nonobese diabetic females. The B6.G7 mouse strain was obtained from Mathis and Benoist as described.  

Reagents

Anti-F4/80 antibody was obtained from AbD Serotec, Oxford UK (cat # MCA 497G, 1:400 dilution), anti-myeloperoxidase (MPO) antibody was obtained from Ventana Medical, Tucson AZ (cat # 760-2659 RTU). 

Adoptive T Cell Transfer and Measurement of Joint Swelling

Naive KRN T cells (H-2b) were isolated from the spleens of male mice age 8 to 10 weeks old using CD4 (L3T4) MicroBeads (Miltenyi Biotec, Auburn, CA) according to the manufacturer’s instructions. By passing the T cells through a second column, they were purified to >98% CD4+ T cells. The percentage of these T cells that expressed the KRN TCR was determined by fluorescence-activated cell sorting (FACS) analysis using anti-Vβ6 and ranged from 85% to 95%. The T cells were washed extensively with 1× HBSS then resuspended in 1× HBSS at a concentration of 3.5 × 10⁶ KRN T cells/ml. Male B6.TCR.Cα–/- H-2b/g7 (recipient) mice age 10 to 14 weeks old were anesthetized with a mixture of ketamine/xylazine, and 100 μl of the cell suspension was injected into the retro-orbital sinus. The two rear ankles of the recipient B6.TCR.Cα–/- H-2b/g7 mice were measured starting at day 0 post T cell transfer. Measurement of ankle thickness was made axially across the tibiotarsal (ankle) joint using a Kafer Dial Thickness Gauge J15 (Long Island Indicator Service Inc., Hauppauge, NY). Ankle thickness was rounded off to the nearest 0.1 mm.

Anti-GPI Antibody Enzyme-Linked Immunosorbent Assay

Mice were bled once a week starting on day 0 post T cell transfer. Sera were stored at −20°C before analysis. Sera were plated at an initial dilution of 1:100 followed by serial half log dilutions in Immulon II plates (Fisher Scientific, Pittsburgh, PA) coated with 5 μg/ml recombinant murine GPI. Donkey anti-rabbit total Igg-horseradish peroxidase antibody was obtained from Jackson ImmunoResearch, West Grove, PA, goat anti-mouse IgM-HRP, IgG1-HRP, IgG2a-HRP, IgG2b-HRP, or IgG3-HRP (Southern Biotechnology Associates, Birmingham, AL) were used as secondary antibodies. Serum antibody was detected using ABTS substrate (Roche Molecular Biochemicals, Indianapolis, IN). Absorbance was measured at 414 nm. The serum titer was defined as
the reciprocal of the last dilution, which gave an OD >3 times higher than that of the background.

**Flow Cytometry**

Popliteal lymph nodes were harvested from \( n = 5 \) recipient mice 35 days post T cell transfer and evaluated for \( \text{V}\beta 6 \) cellular expression. Cells were surface-stained according to standard protocols using the biotin-conjugated rat anti-mouse \( \text{V}\beta 6 \) antibody (BD Pharmingen, San Diego, CA), which represents >60% of the TCR in the K/BxN model, along with streptavidin-phycocerythrin-Cy7 (BD Pharmingen, San Diego, CA). All samples were analyzed on a FACSCanto/FACSDiva platform (BD Biosciences, San Diego, CA), which represents data were analyzed using FlowJo software (Tree Star, Inc., Ashland, OR).

**RNA Isolation and Amplification**

Right-sided hind paws, sectioned just proximal to the tibiotalar joint, were individually flash-frozen in liquid nitrogen and stored at \(-80^\circ\text{C}\). Frozen tissue was later powdered using a SPEX 6750 freezer mill and 6753 microvials (Wolf Laboratories, York UK). Total RNA was prepared from each paw using a modified TRIzol (Invitrogen, Carlsbad CA) isolation protocol incorporating additional purification steps. Total RNA integrity (size and quality) was assessed with an Agilent HP2100 Electrophoresis Bioanalyzer (Agilent Technologies, Santa Clara, CA) and quantified using a NanoDrop 1000 UV spectrophotometer by measuring the absorbance at A260 and A280 nm (Thermo Fisher Scientific, Waltham MA). Typical yields ranged from \( \sim 200 \) to \( >800 \) ng/ul. Ten ng of total RNA from each paw was amplified for two rounds using a modified amplification protocol (Message Amp II aRNA Amplification Kit, Ambion, Austin TX).

**Preparation and Analysis of Agilent Whole Mouse Genome Arrays**

Amplified RNA was labeled using a modified protocol with the Kreatech ULS aRNA Fluorescent Cy3/Cy5 labeling kit (Kreatech Diagnostics, Amsterdam, The Netherlands); specifically, 2 \( \mu \)l of labeling buffer and 4 \( \mu \)l of ULS were added to 5 \( \mu \)g of aRNA in a total volume of 20 \( \mu \)l. Once combined, samples were incubated for 15 minutes at \( 85^\circ\text{C} \) and after incubation samples were returned to room temperature and spun down to collect condensate. All samples were labeled with each dye to allow for generation of fluoro-reverse duplicates. Labeled samples were purified using Zymo Research RNA Clean and Concentrate spin columns according to manufacturer’s protocol (Zymo Research Corporation, Orange CA). Samples were eluted in 100 \( \mu \)l of RNase/DNase-free water. Labeled samples were quantified on the Nanodrop ND-1000 spectrophotometer, and 750 ng of each labeled aRNA from a comparison was added to a total volume of 230 \( \mu \)l water and 9 \( \mu \)l of fragmentation buffer (Agilent Technologies, Santa Clara, CA). Hybridization mixtures were then incubated for 30 minutes at \( 60^\circ\text{C} \). Hybridization mixtures were cooled to room temperature and spun down to collect condensate. 240 \( \mu \)l of 2X Hi-RPM hybridization buffer (Agilent Technologies) was added to each mixture. These probes were then hybridized to 4x44K Mouse Whole Genome Microarrays for 18 hours at \( 65^\circ\text{C} \) in Sure-Hyb Hybridization chambers (Agilent Technologies). Chambers were disassembled in a 0.6X SSC 0.005% Triton X-102 wash bath at room temperature and washed for 10 minutes in wash bath 1. Microarrays were transferred to a 0.01X: SSC 0.005% Triton X-102 wash for 5 minutes at 40°C. Microarrays were then dried with HEPA-filtered compressed nitrogen and scanned on the Agilent Technologies DNA Microarray Scanner at 5 \( \mu \)m resolution. Data were extracted from the scanned image using Agilent Technologies Feature Extraction Software version 9.5.

Background subtraction was performed and normalization was completed using the LOWESS method. After quality control assessments at the spot, chip, and experiment levels, the fold increase or decrease of mRNA expression compared with normal tissue was calculated. The fold value reported for each oligonucleotide was the average of two arrays with swapped fluorophores for experimental and control RNA. Transcriptional array data have been deposited to Array Express (http://www.ebi.ac.uk/microarray-as/ae/browse.html; accession number E-TABM-966). The oligonucleotides where the values did not flip in the fluoro-reverse were excluded from the analysis. A 1.7-fold cutoff was applied, and genes that did not show differential regulation in at least one condition were removed from the analysis. TaqMan confirmation of reported changes in transcript was completed (data not shown).

**Whole Joint Tissue Protein Analyses**

Protein cytokine analysis was performed using a portion of the same right ankle powdered tissue used for whole genome microarray analysis. Protein was extracted using Pierce T-Per tissue extraction buffer in the presence of HALT protease inhibitors (Thermo Fisher Scientific, Rockford IL). The soluble fraction of paw extracts was analyzed using murine 22-plex cytokine luminex immunoassays (Linco/Millipore, Billerica MA). Cytokine concentrations were normalized to the total protein concentration in the same sample using the bicinchoninic acid total protein determination assay (Thermo Fisher Scientific, Rockford IL).

**Microscopic and Immunohistochemical Analyses**

The left hind paw from each animal was fixed in 10% neutral buffered formalin for 24 hours at 20°C, followed by decalcification in Immunocal (Decal Chemical Corpora- tion, Tallman NY) for 7 days at 20°C. Decalcified joints were then paraffin-embedded with medial aspect down for longitudinal view, sectioned twice (4 \( \mu \)m each), and stained with H&E for general evaluation or toluidine blue for specific assessment of cartilage changes. For F4/80 immu-
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nohistochemistry staining, tissue slides were treated with proteinase K (Dako, Carpinteria CA) for 3 minutes followed by blocking solution (Rodent Block M, Biocare Medical, Concord MA) for 15 minutes. This was followed by 30 minutes incubation with F4/80 working dilution. Rat on mouse Polymer kit was used for detection (Biocare), followed by DAB+ (Dako) and a light hematoxylin counterstain. Matched IgG was used for negative control. For MPO immunohistochemistry staining, a CC1 mild cell conditioner was used for pretreatment, followed by a 60-minute incubation with the anti-MPO antibody (Ventana Medical Systems Inc., Tucson AZ). The OmniMap anti-rabbit kit (Ventana) was used for detection followed by hematoxylin counterstain (Discovery XT automated stainer, Ventana).

Joints were evaluated microscopically by a board-certified veterinary pathologist (TPL) and scored for inflammation, cartilage damage, and bone resorption according to previously published criteria.32 For inflammation, scores were as follows: 0 = normal; 1 = minimal infiltration of inflammatory cells in the synovial and/or peri-articular tissues; 2 = mild infiltration with mild edema; 3 = moderate infiltration (including joint space) with moderate edema; 4 = marked infiltration with marked edema; and 5 = severe infiltration with severe edema. For cartilage damage, scores were as follows: 0 = normal; 1 = loss of toluidine blue staining with no chondrocyte degeneration/loss and/or matrix disruption; 2 = loss of toluidine blue staining with minimal chondrocyte degeneration/loss and/or mild matrix disruption in some affected joints; 3 = loss of toluidine blue staining with moderate chondrocyte loss and obvious (depth to deep zone) matrix loss in affected joints; 4 = loss of toluidine blue staining with marked (depth to tide mark) chondrocyte and matrix loss; 5 = loss of toluidine blue staining with severe (depth to subchondral bone) chondrocyte loss and matrix loss in affected joints. For bone resorption, scores were as follows: 0 = normal; 1 = minimal (small areas of resorption in the medullary trabecular or cortical bone, not readily apparent on low magnification, and rare osteoclasts); 2 = mild (increasing areas of resorption in medullary trabecular or cortical bone, not readily apparent on low magnification, with osteoclasts more numerous); 3 = moderate (obvious resorption of the medullary trabecular and cortical bone, without full-thickness defects, lesion apparent on low magnification, and osteoclasts more numerous); 4 = marked (full-thickness defects in the cortical bone, marked loss of medullary trabecular bone, numerous osteoclasts); and 5 = severe (full-thickness defects in the cortical bone, severe loss of medullary trabecular bone). For F4/80 and MPO immunohistochemistry, scores were as follows: 0 = normal; 1 = minimal (small numbers of cells within the synovial epithelium and in proximity to blood vessels within the subsynovium); 2 = mild (increasing numbers of cells evenly distributed within synovial tissues); 3 = moderate (moderate numbers of cells within the synovial tissues and joint space, low numbers of cells within the local peri-articular soft tissues); 4 = marked (obvious cell population within the synovial tissues, joint space, and peri-articular soft tissues with extension to the subcutis and skin); and 5 = severe (high numbers of cells within the synovial tissues, joint space, peri-articular soft tissues, and skin).

Statistical Analyses

Prism 5.01 (GraphPad Software, Inc. La Jolla, CA) was used for all graphical representation of data, including calculation of mean and SEM.

Results

Naive KRN T Cells Induce a Chronic, Destructive Arthritis in T Cell-Deficient Mice

To study the role of T cells in the development of arthritis we transferred naive KRN T cells into T cell deficient B6.TCR.Cα−/−H-2b/g7 mice. These mice expressed the autoantigen GPI on I-Aβ7 and were histocompatible with the transferred KRN T cells (H-2b). On intravenous transfer of purified naive KRN T cells, 100% of the recipient male mice developed arthritis; female mice were also susceptible to disease induction (data not shown). There was a visible erythema and increase in the thickness of the ankles, and the mean thickness of both rear ankles as measured via caliper (mean ± SEM, n = 9) peaked 14 to 21 days post T cell transfer (Figure 1A). A decrease in ankle thickness was present 24 days post T cell transfer and beyond, indicating that the inflammatory response to the cartilage damage shifted to a chronic (decreasing edema) stage. This was consistent with the microscopic presentation (Figures 1, B–F). For instance, Figure 1B (a semiquantitative histopathology summary of general inflammation in the ankle over time) demonstrates a decrease in edema and cell influx after 14 days post transfer. Figures 1, C–F provide representative photomicrographs (H&E, 2× magnification) that highlight the progression from normal (Figure 1C), to acute severe inflammatory cell influx 14 days post T cell transfer (Figure 1D), to severe articular cartilage damage with cartilaginous metaplasia and periosteal new bone formation 29 days post T cell transfer (Figure 1E), to joint ankylosis with severe bone remodeling, ulcerated articular cartilage, extra-articular fibroplasia and few remaining inflammatory cells 42 days post T cell transfer (Figure 1F).

The development of arthritis in the K/BxN mouse model is dependent on antibodies directed against the ubiquitously expressed protein GPI. Therefore, we assayed for total anti-GPI IgG by enzyme-linked immunosorbent assay (mean ± SEM, n = 9). Anti-GPI antibody levels preceded the increase in ankle thickness, yet peaked at the same time (14 days post T cell transfer) (Figure 1G). Antibody levels in KRN-CTM mice persisted for at least 7 weeks, and when their serum was injected into BALB/c mice disease was transferred (data not shown), thereby demonstrating the pathogenic relevance of autoantibodies in this new model. The role of popliteal lymph node lymphocytes in facilitating the production of arthritogenic autoantibodies in K/BxN mice has previously been described.33 In this study, popliteal lymph node cellular expression of Vβ6
neutrophils are found in the joints of RA patients and myeloid cells such as monocytes, macrophages and dendritic cells. Infiltration in the Ankles of KRN-CTM Mice

T Cells Induce Monocyte/Macrophage Infiltration in the Ankles of KRN-CTM Mice

Myeloid cells such as monocytes, macrophages and neutrophils are found in the joints of RA patients and have been previously shown to be important in animal models of arthritis, including the K/BxN model. To determine the role of monocytes/macrophages in KRN-CTM, microarray analysis of macrophage and monocyte transcripts was performed on ankle tissues. Gene transcripts selective for macrophages illustrated a peak 9 days post T cell transfer (Figure 2A), while in Figure 2B, transcripts for monocyte/macrophage CCL cytokines peaked 14 days post T cell transfer with levels maintained through 35 days post T cell transfer. In Figure 2C, semiquantitative scoring of immunohistochemical staining for F4/80 (marker for murine monocytes, macrophages, and dendritic cells) correlated with transcriptional profiling in the illustration of an early influx of cells (peak 14 days post T cell transfer) followed by sustained levels of cells through 42 days post T cell transfer; data points are mean ± SEM (n = 4). In Figure 2, D–F, ankle tissue levels of several protein cytokines and chemokines (interleukin-6, monocyte chemotactic protein (MCP)-1, and macrophage inflammatory protein [MIP]-1α) peaked 7 to 9 days post T cell transfer, with strong correlation between the protein and gene transcript. In Figure 2, G–H, (×2 and ×20 magnification, respectively), photomicrographs revealed a uniform distribution of F4/80-positive cells within the joint space, the synovial tissues, the subcutis and skin, while in Figure 2I many large, activated macrophages are present in the tibiotarsal joint space against a backdrop of (unstained) neutrophils (×20 magnification).

Well-described protein chemoattractants of neutrophils granulocyte-colony stimulating factor, IL-1β, and KC peaked in ankle tissue 7 to 9 days post T cell transfer and were strongly correlated with transcript (Figure 3, A–C). Semiquantitative microscopic scoring of immunohistochemical staining for the neutrophil-specific azurophilic granule enzyme MPO correlated with cytokine levels in identifying an early influx of cells (peak 14 days post T cell transfer), followed by a rapid loss of cell numbers through 42 days post T cell transfer (Figure 3D). During the early influx period large numbers of MPO-positive cells were present within the tibiotarsal joint space with fewer numbers within the subsynovial tissues (Figure 3E, ×10 magnification), while higher power examination of the same image revealed particularly concentrated MPO staining due to phagocytosis of apoptotic neutrophils by activated macrophages (Figure 3F, ×100 magnification).

There was no microscopic evidence of appreciable numbers of lymphocytes in ankle tissues. Also, evidence of an increased lymphoid (T cell and B cell) signature in ankle tissues, the synovial tissues, the subcutis and skin, while in Figure 2I many large, activated macrophages are present in the tibiotarsal joint space against a backdrop of (unstained) neutrophils (×20 magnification).
by neutrophils and macrophages), peaked 21 days post T cell transfer then decreased significantly, presumably as the loss of chondrocytes outpaced the repair response (Figure 4A). Ultimately, this resulted in ulceration of articular cartilage 28 to 42 days post T cell transfer (Figure 4B). In Figure 4C, transcripts in the ankle selective for osteoclasts increased after 14 days and were sustained through 42 days post T cell transfer, correlating with the microscopic observations of active bone resorption by multinucleated osteoclasts (Figure 4D) and invasion by fibrovascular “pannus” tissue (Figure 4E, ×10 magnification).

Discussion

Using clinical, gene transcriptional profiling, protein cytokine and chemokine, and microscopic and immunohistochemical endpoints this study provides a temporal account of how transferring 3.5 × 10^5 purified KRN splenic CD4^+ T cells (H-2^d) into B6.TCR.Cα^−/− H-2^b/q7 recipient male mice resulted in 100% arthritis incidence, with a high degree of disease synchronization through 42 days post T cell transfer. Korganow et al., in a limited study, observed similar arthritis induction with the transfer of KRN spleen cells into a Cα^−/− host, but did not characterize the disease pathology or progression. In this study, arthritis was more severe in the ankle and wrist than in either the knee, elbow, or shoulder joints (data not shown), and progressed from an acute inflammatory cell influx with edema (initiated 7 to 9 days post T cell transfer, peaked 12 to 14 days) to articular cartilage damage with periosteal new bone formation (14 to 21 days post T cell transfer). Bone resorption, with loss of articular cartilage and sustained low numbers of inflammatory cells were present 21 to 42 days post T cell transfer.

Demonstrating CD4^+ T cell-induced myeloid cell recruitment into the joints, immunohistochemistry revealed uniform distribution of F4/80-positive macrophages and MPO-positive neutrophils within the joint space, monocytes/macrophages and neutrophils within the vascular-laden subsynovial tissue, and F4/80-positive dendritic cells within both the superficial and deep layers of the skin. However, appreciable numbers of T cells were not observed in the joint or surrounding tissues via micros-
copy, a finding supported by transcriptional array analysis performed on ankle tissues (no increased expression of CD4, CD3d, CD3e, CD3g, or CD28 relative to normal mice; Supplemental Figure S1 at http://ajp.anipathol.org). Instead, the T cells orchestrating this response likely resided in the popliteal lymph nodes, where Vβ8 expression was strong. Demonstrating that transferred T cells provided B cell help in autoantibody production, KRN-CTM mice produced high titer, arthritogenic anti-GPI antibodies that peaked at the same time as tissue inflammation.

There was a significant increase in macrophage-specific gene transcription markers 9 to 14 days post T cell transfer, which is 3 to 5 days before the observed increase in macrophage numbers via microscopy. This burst of transcription is likely due to the differentiation of recruited monocytes into macrophages, and the expression continued as newly recruited monocytes presumably replaced exiting macrophages. Also, several protein cytokines tracked well with both the microscopic and transcriptional evidence of monocyte/macrophage infiltration into the joint including, but not limited to, CCL3 (MIP-1α), CCL4 (MIP-1β), CCL8 (MCP-2), and CCCL12 (MCP-5). Overall, these findings indicate that the KRN-CTM recapitulated many features of the K/BxN model.18

The KRN-CTM has several key advantages over other murine models of RA, particularly when studying the role of T cells in disease pathology. For instance, one can
manipulate the adoptively transferred T cells (use of knockouts, transgenics, or transfer of either polarized T cell subsets or naive T cells subjected to selective gene knockdown) to characterize specific genes and/or Th1 vs. Th2 vs. Th17 cells in the recruitment and activation of inflammatory cells. Second, given the mouse CIA model is more strain specific (DBA) in terms of disease penetrance and severity, and given that a multitude of knockouts and transgenics are available with the (recipient mice) B6 background the KRN-CTM may be a more flexible platform than the mouse CIA. Third, KRN-CTM arthritis is synchronized, completely penetrant, involves all four paws, and persists to at least 42 days post T cell transfer, thereby requiring substantially fewer animals to run a disease characterization or efficacy study than mouse CIA. The transfer of autoreactive T cells into lymphopenic hosts can generate an exuberant autoimmune response, which may not accurately phenocopy the spontaneous diseases, such as in diabetes models34, however, in this study the KRN-CTM’s presentation of disease was very similar to the spontaneous disease in K/BxN mice and limited to distal joints.

Based on our previous studies33,35 and the results presented here, we propose the following events in KRN-CTM arthritis pathogenesis: 1) transferred KRN T cells expand and are activated in peripheral lymphoid organs; 2) activated KRN T cells provide B cell help inducing autoantibody production; and, 3) recognition of GPI on the articular cartilage surface induces activation of the alternative complement pathway as well as the migration of monocytes/macrophages and neutrophils to the joints. Although the utility of the KRN-CTM model resides in its development of a chronic disease phenotype, it is important to further validate this new model in the context of both prophylactic and therapeutic efficacy studies and compare the results against the models that are currently used most frequently (mouse CIA and rat adjuvant-induced arthritis). The effort described here is the union of complementary scientific disciplines, including transcriptional profiling, chemokine and cytokine protein level analysis and histopathology, and provides a more complete characterization and understanding of the development and progression of disease in the KRN-CTM than any one approach could produce. Based on these results, we propose that the KRN-CTM not only retains the utility of the K/BxN serum transfer and mouse CIA models, but also adds significant logistical and platform advantages. The KRN-CTM presents an opportunity for investigators to study pathways and mechanisms involved in human RA pathogenesis and progression, enabling the validation of druggable targets and biopharmaceuticals.

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