Antibody, 28H1, to which the PS, IRb7e700DX, for tPDT is attached. Here we investigated the feasibility of using FAP-tPDT to induce cell death in murine arthritis model.

**Methods:** After conjugation of the IRb7e700DX to 28H1 (28H1-700DX), binding and specificity of the conjugate was determined. Subsequently, tPDT efficiency in vitro was established using a 3T3 fibroblast cell line stably transfected with FAP. Biodistribution using an [111In] In-DTPA-28H1 conjugate with and without IRb7e700DX was performed in healthy C57BL/6 mice as well as in C57BL/6 mice with antigen induced arthritis (AIA). Finally, the potential of FAP-tPDT to induce targeted cell death in the synovial lining was determined by treating knee joints from mice with AIA ex vivo.

**Results:** Conjugation of IRb7e700DX to the antibody did not negatively influence the immunoreactive fraction or binding capacity of the conjugate (94% for 28H1-700DX and 96% for 28H1-IRb7e700DX). Conjugation of FAP-specific cell death in vitro. At 17.6 J/cm² radiant exposure, 89.24% ± 3.67% of fibroblasts died in the group incubated with antibody compared to control incubated with buffer only (p<0.001). Biodistribution of the compound with the PS showed increased accumulation in the liver compared to the antibody without PS (31.46 ± 5.49% injected dose per gram tissue (%ID/g) versus 5.32 ± 1.17 %ID/g for the antibody with or without PS, respectively (p<0.001)). However, despite this increased clearance to the liver, accumulation in the inflamed joints was increased in the group injected with the antibody-PS construct (1.61 ± 0.08 %ID versus 1.13 ± 0.06 %ID for the antibody with or without PS (p<0.001)). Interestingly, ex vivo FAP-tPDT of knee joints of arthritic mice caused significant loss of the PS (19.69 ± 2.02% fluorescent signal remaining versus 96.00 ± 25.86% compared to the unexposed control at baseline, p=0.047). Furthermore FAP-tPDT induced marked apoptosis as was indicated by an increased staining of the markers caspase-3 and yH2AX evident in the synovium of treated knee joints.

**Conclusion:** Here we demonstrated the feasibility of conjugating a PS to an antibody targeting FAP on activated SF without negatively impacting the binding capacity thereof. Furthermore we showed that this construct can then be used to deliver cell specific cytotoxicity through tPDT both in vitro and ex vivo in a mouse model of arthritis. This approach may have therapeutic potential in the treatment of RA.

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**Background:** In rheumatoid arthritis (RA), autoantibodies against citrullinated proteins (ACPAs) have been reported to be associated with bone loss, pain and tenosynovitis prior to disease onset.

**Objectives:** We aimed to investigate if transfer of human ACPAs into mice could reproduce these clinical observations.

**Methods:** Monoclonal ACPA (1325:04C03 and 1325:01B09) and control antibody (1362:01E02) antibodies (mAbs) were generated from synovial plasma or memory B cells of RA patients. 2mg of combination of monoclonal ACPAs or control antibody were injected in BALB/c female mice (12-16 Weeks) with or without a consecutive intra-articular injection of LPS after 8 days. Pain-like behavior was monitored by measuring mechanical hypersensitivity using von Frey filaments every 3 days and estimation by up-down Dixon method. Bone mineral density was measured by micro-CT. Using specially designed mobilization casts, dedicated mouse MRI coils, and gadolinium enhanced contrast medium, the hind limbs of these mice were scanned and evaluated for any signs of soft tissue joint inflammation. Blinded to ACPA and controls, the MRI images were scored for the presence of synovial thickening, effusion and tendon inflammatory changes by 3 readers in consensus.

**Results:** ACPAs (1325:04C03 and 1325:01B09) induced significantly more pronounced pain-like behavior (lasting for at least 4 weeks) and reduction of the trabecular bone thickness in the hind limbs, whereas no such effect was seen with the control mAbs generated in the same way as the monoclonal ACPAs. While no macroscopic sign of joint inflammation could be detected, preliminary MRI data shows that sub-clinical joint inflammation (such as tenosynovitis) in mice injected with ACPAs but not those injected with control mAbs. Intra-articular LPS injection resulted in significantly increased prolonged mechanical hypersensitivity in mice initially receiving sub-optimal doses of monoclonal ACPA as compared to those receiving control mAb. This was associated with higher levels of sub-clinical inflammation (as shown by MRI scans) in ACPA injected mice.

**Conclusion:** We show that ACPA induces pain-like behavior, bone loss and sub-clinical inflammation in mice, a model that mimics the pre-clinical state of ACPA positive RA.
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SAT0055

JOINT SPECIFIC TNF RESPONSE OF SYNOVIAL FIBROBLASTS IN RHEUMATOID ARTHRITIS

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Background: Synovial fibroblasts (SF) in rheumatoid arthritis (RA) play a major role in chronic inflammation and joint destruction. We showed previously that their epigenetic and transcriptional profile as well as their function vary significantly between different joints. However, it is unknown whether there is a joint-specific inflammatory response of SF.

Objectives: To compare transcriptional changes between SF from hand, shoulder and knee joints after stimulation with the pro-inflammatory cytokine TNF.

Methods: We cultured SF from synovial tissues of hand (metacarpophalangeal and proximal interphalangeal joint), shoulder and knee joints of RA patients. After stimulation of the SF with 10 ng/ul TNF for 24h (n=2 for each joint location), RNA was sequenced on the Illumina HiSeq4000 platform. We analyzed differential gene expression with R v3.5.2 and CuffDiff and DESeq2 packages.

Results: As shown in Figure 1, principal component analysis showed evident separation of joint locations and condition (unstimulated vs TNF stimulated). In a sample-to-sample distance matrix, hand samples (TNF stimulated and unstimulated) grouped apart from shoulder and knee samples (Figure 2).

Of the regulated genes, 26% appeared in all three joint locations and 56% overlapped between knee and shoulder, but only 30% overlapped between hand and shoulder and between hand and knee SF. Similarly, also enriched pathways differed particularly between hands and the more proximal joints.

‘Defense response’ (p=8.11x10^{-23}) and ‘cytokine activity’ (p=1.27x10^{-21}) were the most significantly enriched gene ontology terms for genes regulated in TNF stimulated hand SF. These processes were less prominently enriched in stimulated knee (1.58x10^{-15} and 8.84x10^{-09}) and shoulder SF (2.02x10^{-11} and 3.20x10^{-05}), where ‘cell cycle’ (p=2.26x10^{-30} in knee and p=2.18x10^{-32} in shoulder), and ‘DNA packaging complex’ (p=4.63x10^{-39} in knee and p=8.91x10^{-36} in shoulder) were the most significantly enriched gene ontology terms. These processes were not significantly enriched in stimulated hand SF.

Conclusion: SF from different joints in RA react differently to TNF stimulation. In particular hand SF reacted different to TNF stimulation than shoulder and knee SF, which appeared more similar. These qualitative and quantitative differences of the inflammatory response might translate into joint-specific pathotypes of synovitis with distinct therapeutic responses and disease outcomes.

Figure 1. Principal component analysis of TNF stimulated and unstimulated synovial fibroblasts

Figure 2. Sample-to-sample distance matrix of TNF stimulated and unstimulated synovial fibroblasts