Lab on a Chip



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Cite this: Lab Chip, 2021, 21, 4128

Received 19th February 2021, Accepted 7th September 2021

DOI: 10.1039/d1lc00130b

rsc.li/loc

Introduction

In today's ageing society disorders affecting the human musculoskeletal apparatus, also referred to as musculoskeletal diseases (MSD), are of great clinical and scientific interest. In particular, arthritis describes various pathophysiological conditions, including inflammatory arthritis (IA) and osteoarthritis (OA). Types of IA include rheumatoid arthritis (RA) systemic inflammatory autoimmune

Establishment of a human three-dimensional chipbased chondro-synovial coculture joint model for reciprocal cross talk studies in arthritis research[†]

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Rheumatoid arthritis is characterised by a progressive, intermittent inflammation at the synovial membrane, which ultimately leads to the destruction of the synovial joint. The synovial membrane as the joint capsule's inner layer is lined with fibroblast-like synoviocytes that are the key player supporting persistent arthritis leading to bone erosion and cartilage destruction. While microfluidic models that model molecular aspects of bone erosion between bone-derived cells and synoviocytes have been established, RA's synovial-chondral axis has not yet been realised using a microfluidic 3D model based on human patient *in vitro* cultures. Consequently, we established a chip-based three-dimensional tissue coculture model that simulates the reciprocal cross talk between individual synovial and chondral organoids. When co-cultivated with synovial organoids, we could demonstrate that chondral organoids induce a higher degree of cartilage physiology and architecture and show differential cytokine response compared to their respective monocultures highlighting the importance of reciprocal tissue-level cross talk in the modelling of arthritic diseases.

conditions often found in diarthrodial joints.¹ Overall, arthritic diseases frequently lead to swelling, pain and impaired movement of joints, severely affecting the quality of life.^{2,3}

Fibroblast-like synoviocytes (FLS) are the primary synovial cell type studied in RA.⁴ In healthy synovia, FLS lubricate the joint by producing e.g. lubricin or hyaluronan, which ensures homeostasis and function. FLS are also responsible for synovial lining formation and maintaining the inner joint membrane (Fig. 1A, left panel).⁵ In RA, the synovium is the main joint tissue structure infiltrated by immune cells at the synovial sublining layer and is a critical player in osteoclastmediated bone erosion and pannus formation as well as enzymatic cartilage degradation.6,7 During the onset and progression of RA, FLS are stimulated by cytokines secreted from activated and migrating immune cells such as macrophages, B cells and T cells (e.g. IL-1β, TNFa, IL-6, IL-17), and in turn also secrete cytokines (e.g. IL-6, IL-15, IL-23), chemokines (e.g. IL-8. CXCL12) and degradative enzymes (various metalloproteases as well as cathepsins).⁸⁻¹⁵ In this context, IL-6 is a central pro-inflammatory molecule promoting joint inflammation and destruction.16,17

Over the last decade, arthritis research has gradually moved from conventional two-dimensional to more complex

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 $[\]dagger$ Electronic supplementary information (ESI) available. See DOI: 10.1039/ d1lc00130b

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Fig. 1 A) Schematic drawing of the synovial activity influencing cartilage and subchondral bone in synovial knee joints. B) Overview of the organoid-based joint-on-a-chip coculture system comprising chondral and synovial compartments for reciprocal inflammatory cross talk.

three-dimensional cell cultures, which allowed more physiologically relevant interactions mediated by unidirectional as well as reciprocal cell communication and transformation to human arthritic synovial tissue in vitro.¹⁸ Starting in 2006, Kiener et al. introduced a 3D synovial micromass organoid model with FLS embedded in Matrigel to form a synovial construct comprising lining and sub-lining layers spontaneously.⁵ This model has been used to investigate the influence of FLS activity on inflammatory cascades¹⁹ and the contribution of the complement system in FLS metabolism.²⁰ Broeren et al. adapted this Matrigel-based primary model by combining FLS and CD14⁺ monocytes. When stimulated with TNF- α , these micromasses triggered arthritic hallmarks including synovial lining thickening, proinflammatory macrophage phenotype (M1) as well as upregulation of pro-inflammatory cytokines.²¹ Maracle et al. used a 3D drug screening model based on spheroids fabricated from RA-FLS and endothelial cells to recapitulate synovial angiogenesis, which constitutes a detrimental process in RA disease progression.²² Furthermore, Peck et al. successfully engineered a scaffold-free tri-culture drug screening model comprising of FLS (SW982 cell line), lipopolysaccharide (LPS)-activated macrophages and primary isolated chondrocytes from knee-cartilage to investigate cartilage deconstruction and RA reversal in affected porcine joints.²³ Our group used a lab-on-a-chip with non-invasive light scattering sensing strategy to rapidly investigate TNFa induced rheumatoid architectural changes in 3D synovial micromass cultures concurring with synovial reorganisation as well as interleukin (e.g. IL-6, IL-8) and MMP (e.g. MMP-1, MMP-3, MMP-13, etc.) molecule secretion.²⁴ More recently, also microphysiological models and organ-on-a-chip devices have been introduced to recapitulate various aspects of arthritic diseases: Ma et al. established a coculture of human FLS (synovial sarcoma SW982 cell line) with murine preosteoclasts (RAW264.7) and bone-marrow-derived stem cells (BMSCs) to study FLS migration and invasion-mediated bone erosion in RA.25 Nonetheless, to date most systems rely on non-human or immortalised cell lines to establish these complex 3D cocultures which are less predictive to model arthritic processes in human.

Therefore, herein we present a human joint-on-a-chip model based on co-cultivation of two spatially separated hydrogel-based 3D organoid constructs to understand how reciprocal tissue cross talk between synovial and chondral organoids and *vice versa* stimulates joint (patho)physiology (see Fig. 1B). An essential aspect of our joint-on-a-chip development capable of mimicking the human chondro-

synovial niche was relating to improved reproducibility, which is key to all advanced in vitro cell-based assays. Consequently, a range of parameters known to influence organoid formation is investigated, including passaging of isolated synovial cells in 2D, variations of immune cell composition and cytokine secretion. Another focus is the investigation of the effects of chondro-synovial cross talk on tissue-level behaviour. Also, culture conditions including chondrogenic growth factor (TGF-B3) and fetal calf serum (1% or 10% FCS) supplementation were investigated to assess organoid maturation and soluble cross talk changes. Overall, the proposed chip-based organoid coculture model shows promising results due to а high degree of microenvironmental and cellular control. We also excellent imaging capabilities demonstrate of the architectural context and cellular composition with our proposed biochip features, that are impossible for randomly placed conventional 3D micromass and spheroid cultures established within microtiter plates.

Experimental

Clinical specimens

Human synovial tissues from patients were obtained from RA patients undergoing synovectomy (fulfilling the ACR/EULAR classification criteria²⁶) with written informed consent under the terms of the ethics committee of Medical University of Vienna (EK-No. 1065/2011). The medical background of patients was documented and included age (average age: 55, min.: 27, max.: 71), sex (8 female and 2 male), tissue location (carpals: 4, phalanges: 3, elbow joint: 2, tibiofemoral joint: 1), diagnosis (total: 10, rheumatoid arthritis: 9 patients, control: 1 OA patient) and medication (no treatment: 2, Embrel: 2, Arava: 1, cortisone: 3, Salazopyrin: 1, Arava + cortisone: 1). Primary cell populations were isolated and stored in liquid nitrogen, as described earlier.²⁷

Cell culture

FLS were thawed at 37 °C and immediately plated in a T75 cell culture flask containing pre-warmed DMEM culture medium supplemented with 10% FCS, 1% MEM NEAA and 1% antibiotics. Cells were split upon reaching 90% confluency. Before passaging, 1 ml of culture medium was collected and frozen for supernatant analysis. For passaging experiments, cells were enzymatically detached, pelleted and resuspended in culture medium. An aliquot of 5×10^5 cells was weekly transferred to a new flask, whereas the residual cells were used for flow cytometry analysis and 3D cultivation. For cross talk experiments, expanded FLS were used between P5 and P7 if not indicated otherwise.

Commercial human primary chondrocytes (C-12710; female donor; PromoCell) were maintained in chondrocyte growth medium (411–500; CellApplications Inc.), which was exchanged every other day. Cells were expanded until passage 5 and frozen as aliquots for on-chip cross talk studies. Well plates were coated with 0.03 w/v% ethanolic poly(2-hydroxyethyl methacrylate) (poly-Hema) as described initially for synovial micromass cultures.⁵ Briefly, after pelleting, the primary FLS cells at a passage number between 1 and 8 were resuspended in ice-cold Matrigel at a concentration of 3000 cells per μ l Matrigel. To initiate micromass cultures, a 20 μ l drop of the cell-laden Matrigel suspension was pipetted into the centre of a coated well. After 40 min of polymerisation, cell/Matrigel drops were overlaid with 2 ml fully supplemented organoid culture medium containing 1% insulin-transferrin-selenium (ITS) and 0.032 μ g ml⁻¹ L-ascorbic acid 2-phosphate with a weekly medium exchange.

Microfabrication

The two multi-layered designs for either synovial monoculture or chondro-synovial coculture illustrated in ESI[†] Fig. S1 were fabricated using a CAMM-1 GS-24 vinyl cutter (Roland, Germany). The devices were assembled layer by layer by xurography from two-dimensional CAD templates from 500 μ m thick transparent silicone sheets (MVQ Silicone) as described previously.^{28,29} The basic design principles were adapted from an earlier vascular organoid study,³⁰ and circular medium reservoirs were added to contain a total medium volume of 300 μ l. For height variation, the layer numbers of the circular hydrogel compartment were adjusted in the range of 1–4 mm. To prevent cellular outgrowth from within organoids, chip surfaces were coated with 0.5% ethanolic Lipidure® (Amsbio, UK) overnight before UV light sterilisation for 1 hour.

Two basic designs with five structured PDMS layers were assembled on object slide base layers to form a joint-on-achip (see ESI[†] Fig. S1A): the monoculture design (left column layers I-V) comprised of a single organoid chamber (Ø 5 mm). In contrast, two opposing organoid chambers allow for soluble tissue-level communication via reciprocal signalling (right column).^{30,31} To improve medium replacement and sample withdrawal of the optimised design, the medium supply channels interconnect two circular medium reservoirs. To enable reproducible and facile loading of hydrogels, structured layers I and II form a step-like hydrogel stopper feature that prevents hydrogel bleeding into the medium supply channel. Additionally, structure layers III-V form a chamber geometry favouring controlled organoid formation by an anchoring organoid condensation guide (see ESI[†] Fig. S1B and C for detailed functional and dimensional elaborations on both features).

On-chip organoid culture

For the generation of synovial organoids, a hydrogel suspension containing FLS was loaded *via* a loading vent into each cell cultivation chamber. To start organoid formation, a total of 45 μ l FLS cells in Matrigel were slowly injected at a concentration of 3000 cells per μ l through the hydrogel port and was sealed by a small piece of PCR sealing film on ice.

After 40 min of polymerisation at 37 °C, 400 µL organoid chip medium was added to start the organoid culture. A previously established cartilage organoid based on TISSEEL fibrin hydrogel (Baxter, Austria) was injected in the chondral compartment as previously described to establish on-chip cocultures.^{31,32} Briefly, 30 µl of fibrinogen solution at a concentration of 100 mg ml⁻¹ containing 2000 cells µl⁻¹ were mixed with 30 μ l 4 U ml⁻¹ thrombin solution supplemented with 40 µM CaCl₂ solution and was polymerised for 30 min at 37 °C. For medium composition experiments, TGF-β3 was added to the chip medium at 1 and 10 ng ml⁻¹ in either 1% or 10% FCS-containing organoid culture medium and compared to commercial chondrogenic differentiation medium (411D-250; CELL Applications Inc., USA). For medium replacement, the complete chip reservoir medium of 300 µL was replaced with fresh organoid medium and supernatant was frozen for off-chip supernatant analysis. After cultivation, the top ceiling of the organoid chambers was removed with a scalpel, and the individual organoids were placed in 7% paraformaldehyde for histological preparation or cytometry buffer for flow cytometry analysis as described below.

Flow cytometry

Two different staining mixes were prepared on ice, as shown in ESI[†] Table S1. Aliquots of 1.5×10^5 cells were washed with flow cytometry buffer (PBS with 2% FCS) and pelleted at 4 °C and 1350 rpm for 8 min. After pelleting, cells were resuspended in 50 µl of the staining mix solution, incubated for 20 min, washed and resuspended in 150 µl flow cytometry buffer. Measurements were performed on a BD Canto II flow cytometer using BD FACS VerseTM (note: one measurement cycle lasted either 100 000 cells or 2 min). Gating of cell populations was performed as described elsewhere.^{33,34} Cell populations from organoids were harvested by centrifugal removal after Matrigel liquification for 2 h on ice.

Confocal microscopy

Imaging was performed using an upright Leica TSC SP5 Microscope (Leica Microsystems) with 20× and 40× water immersion objectives. Imaris software (Bitplane) was used for image processing. Live-cell staining was performed with fluorescent cell tracker dyes (Green CMFDA/Orange CMRA and Deep Red; Thermo Fisher) before pelleting and hydrogel mixing according to the manufacturer's staining protocol and ESI† Table S2.

Histology

Tissue and organoid specimens were fixed in 7% paraformaldehyde, dehydrated in ethanol and paraffinembedded according to standard in-house procedures.³⁵ Paraffin sections of approx. 2.5 μ m thickness were stained with hematoxylin and eosin (HE) for morphological evaluations.

Morphometric analysis

For morphometric analysis, manual tracing of the organoid boundaries of phase-contrast images of organoids taken at selected time points was used in ImageJ (version 2.0.0.rc.69/ 1.52.p) in combination with the shape descriptors option of the 'measure' as well as the 'analyse particles' functions as previously described.³⁶ For synovial network parameter analysis AngioTool (version 0.6a) was used with optimised analysis parameters (vessel diameter: 5; intensity range: 15–100; remove small particles option set to 240).

Supernatant protein analysis

For AlphaLISA analysis of IL-6, IL-8, MMP-1 and MMP-3 (AL223C, AL224C, AL284C, AL242C, Perkin Elmer), undiluted supernatants of 2D cultured FLS were analysed according to the manufacturer's protocol using an EnSpire® Multimode plate reader (Perkin Elmer).

For cytokine analysis of IL-6, IL-8, MMP-13 and VEGF using human Magnetic Luminex Assay (R&D Systems), supernatants of synovial and chondral organoids in mono and coculture were diluted 1:10 in dilution reagent and analysed according to the manufacturer's protocol using a LuminexTM MAGPIXTM Instrument System (Invitrogen). After medium background subtraction, molecule secretion of mono- and cocultures were normalised to 10^4 cells per mL and day to compensate for differences in total cell numbers between synovial and chondral monocultures as well as cocultures. To reduce overall sample size, occasionally supernatants from n = 3-4 organoids were pooled for analysis of medium compositions (TGF/FCS content) as well as monoculture organoids in the comparative study on mono- and co-culture organoids.

RT-qPCR preparation and analysis

mRNA of snap-frozen patient tissue and organoids using an optimised extraction protocol (Qiagen Easy prep) using Precellys lysing kits (CK25; 3×20 s, 5000 rpm, 3 min cooling break) in combination with frozen lysis buffer beads (innuPREP RNA Mini Kit, Analytik Jena) supplemented with 1% beta-mercaptoethanol on a Precellys 24 system. RNA was reverse transcribed into cDNA (Firma und Kit einfügen) and subsequent RT-qPCR was performed on a QuantStudio3 QPCR system (Applied Biosystems, Thermo Fisher) using SensiMix SYBR Green Master Mix as previously described.³⁵ Primer sequences are described in ESI[†] Table S3.

Statistical analysis

Prism software 7 (version 7.0a; GraphPad, USA) was used for statistical analysis of data sets for either *t*-test analysis (paired and unpaired) or 2-way ANOVA combined with appropriate *post-hoc* analysis (Tukey, Sidak or Dunnet's test as recommended by the software). Shapiro–Wilk normality test for normal distribution was performed. *P* values < 0.03

were considered as statistically significant (*p < 0.03, **p < 0.002, ****p < 0.002, ****p < 0.0001).

Results & discussion

Characterisation of the chip-based three-dimensional synovial organoid model

Initially, individual hydrogel compartments were studied separately to investigate each organoid unit's formation and maturation. In synovial organoid cultures, RA patient-derived primary FLS were mixed into Matrigel before injection and cultured for up to four weeks. It is important to note that during synovial organoid formation, significant hydrogel condensation and remodelling takes place. For instance, Fig. 2A shows the influence of increasing hydrogel volume (note: increasing chamber heights from 1 to 4 mm results in a volume increase from 19.6 to 78.5 µl) on condensation behaviour, where RA-FLS organoid volume gradually declined to between 18-36% after 28 days of organoid cultivation. Interestingly, a chamber height of 1 mm (19.6 µL) was frequently insufficient to generate a single synovial organoid; instead, fragmentation of the organoids and cell dispersion as well as a significantly different condensation behaviour starting at day 6 (p < 0.0001; Tukey's multiple comparisons test) occurred (see also images of ESI† Fig. S2) in comparison to the other hydrogel volumes. In contrast, chamber heights of 1.5, 2.0 and 2.5 mm displayed uniform organoid formation exhibiting an area of $3.8 \pm 0.27 \text{ mm}^2$ at day 28. Larger volumes (e.g. chamber heights of 3 and 4 mm) produced bigger synovial constructs with 5.7 \pm 0.12 mm² and 7.1 \pm 0.09 mm², respectively. A more detailed morphology analysis revealed that most homogenous synovial organoid formation is achieved using a cultivation chamber with 2 mm height, where almost spherical structures are obtained. Fig. 2B illustrates the time-dependent change of synovial organoid shape from rectangular to spherical throughout 28 days of the cultivation period, which points at increase of organoid sphericity over time as the FLS organoids mature. During early biochip design iterations we have frequently observed that when organoids came in closer contact, they formed direct connections and allowed FLS to migrate towards the chondral unit. As more reproducible spatial control over organoid positioning and maturation was the aim of the current biochip design to enable reciprocal molecular cross talk in the absence of direct cell-cell interactions, it was imperative to guide and spatially control formation of the



Fig. 2 Performance evaluation of the synovial on-chip organoid technology to form organoids from RA patient-derived fibroblast-like synoviocytes. A) Influence of organoid chamber height on organoid area throughout 28 days of cultivation with representative phase contrast images at day 28 post-seeding. Data is expressed as mean \pm standard deviation for n = 3 organoids of n = 1 donor. Two-way ANOVA with Dunnett's multiple comparisons test was performed (**p < 0.002, ****p < 0.0001). B) Change of synovial organoid shape from rectangular to spherical throughout 28 days of cultivation in 2 mm high organoid compartments using manual z focus analysis of the organoid surface. C) Evaluation of precision of organoid positioning of synovial organoids relative to the hydrogel anchor (upper graph) and chamber centre (lower graph). Dashed line indicates theoretically best geometrical values. Data is expressed as mean \pm standard deviation for n = 32 organoids from n = 3 donors (overall sample population includes organoids from cell passages up to p4 and generated by height variations).

two organoid types on the biochip using the proposed features. A detailed investigation of overall relative and absolute positioning precision in relation to the hydrogel anchor and the waveguide features was performed as shown in Fig. 2C (for details see also ESI† Table S4). The on-chip positioning features improves the hydrogel positioning of the tissue organoids during the first week of the initial organoid remodelling phase with a lower relative standard deviation of 6.6% compared to 11.9% at day 7 and 32% at day 21 post seeding, which indicates that the integrated structures improve the initial positioning of the hydrogel. During three weeks of cultivation, both positioning parameters aggravated by manual user manipulation (*i.e.*, sample withdrawal and medium exchange). However, positioning accuracy of organoids was not significantly affected by height of the hydrogel compartment (p > 0.1, Tukey's multiple comparisons *post hoc* test). Next, on-chip synovial condensation dynamics were compared to polyHEMA-coated anti-adhesion well plates (MMC). Results of this comparative study shown in Fig. 3A revealed similar overall architectural features with a significantly higher reproducibility exhibiting a 3.5-fold lower relative standard deviation of 4.4% of organoid biochips (n = 5) compared to 15.5% (n = 3) for MMC cultures generated in anti-adhesion well plates.



Fig. 3 Morphology analysis of FLS micromass organoids floating in wellplates in comparison to FLS on-chip organoids. A) Analysis of organoid area and condensation dynamics over 28 days and initial synovial network formation up to day 28 for on-chip organoids and wellplate micromass organoid cultures. Data is expressed as mean \pm standard deviation for n = 3 organoids generated from n = 1 donor. Paired *t*-test was performed (n.s., not significant; p > 0.1). B) Confocal images of CMFDA-stained FLS network maturation over the initial culture period of 7 days for micromasses (top row) and on-chip organoids (bottom row) with C) AngioTool analysis software analysis overlays. D) Comparative analysis of synovial network parameters including network area, junction number and synovial network protrusion length for micromasses (n = 2) and on-chip organoids (n = 3). Unpaired *t*-test was performed. Data is expressed as mean \pm standard deviation n = 3 donors and n = 2 organoids.

Moreover, morphometric AngioTool image analysis revealed that synovial network morphology (see Fig. 3B–D) was similar for synovial micromasse and biochip organoids with gradually increasing parameters. The total increase in vessel area 21.45 \pm 4.7% and 29.2 \pm 0.4%, protrusion length of 119.8 \pm 29 μ m and 135.6 \pm 14 μ m, as well as number of junctions of 627 \pm 201 and 970 \pm 55 junctions per mm² was observable for biochip in comparison to micromass organoids over the first 7 days of network maturation.

Influence of passaging and cultivation on the heterogeneity of patient-specific synovial cell populations

Arthritic synovial tissue comprises, aside from synovial fibroblasts (FLS), a diverse set of immune cells such as macrophages, B cells and T cells. While standard 2D monoculture conditions is known to rapidly deplete immune cells resulting in pure synovial fibroblasts (FLS) using DMEM medium,²¹ less is known about the effect of this immune cell depletion on the formation of FLS organoids. To evaluate the long-term effect of co-cultivating isolated patient derived FLS and immune cells on synovial tissue heterogeneity, the samples of seven RA donor tissues were characterised using flow cytometry and supernatant analysis. Results shown in ESI† Fig. S3A and B indicate a gradual depletion of leucocytes and selective growth of synovial fibroblast over 5 weeks (5 passages). A high degree of interpatient immune cell variability was evident for CD45⁺ leucocytes ranging from 22% to 1.1% between individual patient samples. Further analysis showed variations of sub-population of 4.1% to 0.1% CD3⁺/CD14⁻ T cells, and 3.9% to 0.01% CD19⁺/CD14⁻ B cells as well as a predominant population of 8.9% to 0.6% CD14⁺/ CD3⁻ macrophages. Similarly, a gradual decrease of proinflammatory cytokines (e.g. IL-6 and IL-8) and degradative matrix metalloproteinases (e.g. MMP-1 and MMP-3) in cell culture supernatant was detected over the five-week cultivation period with weekly subcultivations (see ESI† Fig. S3C).

Next, the impact of medium composition for on-chip organoid cultivation and 2D/3D cultivation on immune cell survival was investigated in more detail. Initial characterisation of CD45 positive cells of the freshly isolated synovial cell suspensions from donor tissue (n = 4 biological donors) displayed 80.2 ± 3.1% podoplanin positive FLS and 5.7 \pm 4.7% CD45 positive leucocytes (*i.e.*, T cells, B cells and macrophages), which was an equal population distribution also found in p1 subcultivations (see ESI† Fig. S3A). Results shown in Fig. 4A revealed a higher degree of leucocyte survival when substituting the DMEM culture medium with leucocyte medium (e.g. RPMI 1640), leading to 4.9, 2.4 and 1.8-fold higher T cell, macrophage and B cell subpopulations from week one to two post-seeding (**p < 0.01; 2-way ANOVA with Sidak's multiple comparisons test). In turn, synovial organoid on-chip cultivation further improved leucocyte survival for both medium types (n = 2; see Fig. 4B) with 11and a 20-fold increase in leucocyte populations compared to



Fig. 4 A) Impact of medium type on leucocyte populations within synovial RA-FLS organoids. Data is expressed as mean \pm standard deviation in % of total CD 45 positive cell populations for n = 2 organoids for n = 4 donors. B) Comparison of CD45 positive leucocyte populations from isolated RA synovia in p0 cultured as 2D cultures or 3D organoids in DMEM and RPMI 1640 medium using flow cytometry analysis. Data is expressed as mean \pm standard deviation in % for n = 2 donors and n = 2 organoids. Two-way ANOVA with Sidak's multiple comparisons test was performed. (***p < 0.002).

conventional 2D cultivation, which showed a declining population trend. First of all, this means that both the amount and type of immune cells present in rheumatoid arthritis patients vary significantly and need to be accounted for in an *in vitro* arthritis model. For FLS 2D cultures, immune cell depletion decreases the arthritic sample heterogeneity regarding cell composition and cytokine secretion and favours FLS growth. Furthermore, our chipbased synovial organoid system that features a threedimensional tissue-like microenvironment protects synovial leucocyte populations from depletion even in unfavourable culture media (*e.g.* DMEM).

Impact of passaging on the formation of chip-based synovial organoids

To further understand the effects of synovial cell passaging on chip-based 3D synovial cultures, organoid formation and secretion of cytokines were investigated in ensuing experiments. On-chip generated synovial organoids of increasing passages (p1–5) were cultivated over five weeks, and vital organoid parameters including area, solidity, perimeter, roundness and organoid numbers per unit were analysed. Fig. 5A and ESI[†] Fig. S4 show morphometric analyses at day 28 post-seeding and point at stable organoid formation peaking at 5.5 \pm 0.6 mm² and 5.8 \pm 0.06 mm² of organoid area for later passages in contrast to freshly isolated FLS populations. Early passages (p1) showed a tendency to form smaller cell aggregates ranging from 1.8–4.1 mm² with higher variability (0.42% RSD). In turn, FLS organoids formed from p5 cells yielded reproducible organoid areas of 5.6 \pm 0.67 mm² (0.01% RSD). The remaining morphometric parameters, including perimeter (0.04% RSD), roundness (0.05% RSD) and solidity (0.01% RSD), further pointed at the highest reproducibility in the presence of higher passage numbers (p3–4). Furthermore, the system's reproducibility to



Fig. 5 A) General morphometric analysis of synovial organoids produced with fibroblast-like synoviocytes from increasing passage numbers after 28 days of cultivation in mono organoid biochips. B–F) Donor specific influence of fibroblast-like synoviocyte passage number and cell origin on morphology parameters including B) area, C) solidity, D) perimeter and E) roundness, and F) number of organoids per compartment. Data is expressed as mean \pm standard deviation for n = 3 donors and n = 3-4 organoids.

form a single organoid in each organoid compartment scored best for organoids created from cells after 4-5 weeks of passaging. Investigating patient heterogeneity for organoids produced from individual patient-derived synovial tissues (Fig. 5B-F) the overall effect of FLS passaging on organoid formation was still observable. While most parameters including roundness, perimeter and solidity were not affected by patient origin (p > 0.1, 2-way ANOVA with Tukey's posthoc test), organoids from three different RA patient tissue biopsies showed a significant difference in both organoid areas (p < 0.0002) and the number of formed organoids (p < 0.03). These indicators still point out that passaging of FLS populations improves the reproducibility of organoid formation. On-chip cultivation of primary cell in a 3D environment is known for more native tissue-like molecule retention and secretion.37,38 Therefore, next the secretory behaviour for FLS organoids over increasing passages and cultivation time points were monitored as shown for IL-6, IL-8, MMP-13 and VEGF secretion up to passage p5 and week three post-seeding in Fig. 6A-D.

FLS organoids overall showed initially higher overall secretory capacity in week 1, which declined significantly over three weeks of cultivation independent of passage number or molecule type (p < 0.002). Considering the passaging effect, RA-FLS organoids showed a considerably higher secretion performance of MMP-13 and VEGF (p < 0.03, 2-way ANOVA

with Sidak's posthoc test) with an increase for later passages, which again equalized over three weeks of culture. The passage dependent differences in the early secretory activity of FLS organoids are most probably not caused by immune cell depletion alone as no direct correlation between the number of immune cells, specific secretory molecules (*i.e.*, interleukins or MMPs) were discernible. Furthermore, activated FLS are known for their high secretory capacity.⁴ This means that other factors such as inflammatory memory or reprogramming²⁰ before isolation are more likely to influence how RA-FLS react and mature during organoid formation in 3D.

Overall, by intentionally reducing cell heterogeneity in the synovial organoid constructs by subcultivation for 5 passages we could increase reproducibility over the cultivation duration by limiting initial cellular variations. Consequently, RA-FLS cells from p5 were used in all subsequent experiments on cross talk studies to reduce sample variability and improve organoid yield and homogeneity.

Evaluation of the chondral organoid model in monoculture biochips

Recently our group has studied the impact of biochip culture³¹ as well as matrix stiffness/composition³² on chondrocyte phenotype. Previous results already indicated that primary chondrocytes embedded in 50 mg ml⁻¹ fibrin



Fig. 6 Secretion analysis of synovial organoids at day 7 and 28 post-seeding produced with fibroblast-like synoviocytes from increasing passages for A) IL-6, B) IL-8, C) MMP-13, and D) VEGF using Luminex multiplex assay. Data is expressed as mean \pm standard deviation for organoids generated from n = 3 donors. Two-way ANOVA with Tukey's multiple comparisons test was performed. (*p < 0.03).

hydrogel show good chondrocyte viability, morphology and cartilage-specific functions (i.e., collagen II, cggrecan, etc.) independently of culture technique (i.e., biochip organoid vs. micromass). Before co-cultivation and cross talk experiments chondro-synovial organoids, chondral on organoids comprising of human articular chondrocytes suspended in 50 mg mL⁻¹ fibrin were re-evaluated to demonstrate that the current biochip is in agreement with previous findings. Confocal imaging confirmed the typical round morphology of articular chondrocytes within the Fibrin construct after seeding (see Fig. 7A and ESI[†] Video S1) with 83.46 ± 16.5% chondrocyte viability at day 21 post-seeding (see Fig. 7B and C). Further histological comparison (see Fig. 7D) revealed that the chip-based chondral organoids can recapitulate the morphological and cell density characteristics of human articular cartilage of superficial tissue zones, which are primarily affected by mechanical wear as well as protease degradation. Furthermore, the chondral organoid derived from a single healthy donor sample indicated comparable ratios of cartilage health indicators Col II/Col I and AGC/Col I ratio similar to values found in native human cartilage (n = 3 donors; see Fig. 7E). Overall these

results indicate good agreement with previous studies in terms of native tissue-like properties of fibrin-embedded chondrocytes *in vitro*.

Establishment of a joint-on-chip organoid coculture model for chondro-synovial cross talk studies

Another critical aspect in establishing a functional in vitro arthritis model of the joint deals with proper tissue to tissue interactions between the synovium as inflammatory tissue and the cartilage as target tissue during the onset and progression of the disease. High serum content is a crucial requirement for FLS since it provides and maintains the proper synovial cell function and architecture. However, redifferentiation of expanded chondrocytes is conducted frequently using differentiation media that contain no or low serum concentrations, but TGF-ß3 and other chondrogenic supplements to promote a chondrogenic niche.39 Consequently, the impact of TGF-ß3 in addition to serum content (e.g., 1 ng ml⁻¹ vs. 10 ng ml⁻¹, 1 vs. 10% FCS) was investigated against an unknown commercial chondrogenic medium formulation to identify suitable coculture media



Fig. 7 A) Fluorescence images of CMFDA-stained primary chondrocytes in the chondral fibrin-based construct day 1 post seeding. B) Viability and C) representative phase-contrast and fluorescence images of CMFDA (live) and ethidium bromide-stained (dead) human articular chondrocyte organoids at day 21 post-seeding (data is expressed as mean \pm sdev for n = 4 FLS organoids). D) Histological comparison of on-chip chondral organoids (left image) and the superficial region of human articular cartilage tissue from a human tibia plateau. E) Comparison of Col II-to-Col I and AGC-to-Col I ratio of gene expression of chondral organoids generated from primary chondrocytes at day 21 post-seeding with native human cartilage tissues.



Fig. 8 Impact of medium type and supplementation on organoid formation and secretion profile of chondro-synovial organoid cultures. A) Progression of organoid area and representative microscopic images of chondro-synovial organoid cocultures (n = 3-4), and B) secretion analysis of IL-, VEGF and MMP-13 (n = 1) of cocultured chondro-synovial organoids for TGF- β 3 supplemented DMEM as well as chondrogenic differentiation medium at day 3, 8, 16 and 21 post-seeding using Luminex multiplex analysis. Data is expressed as mean ± standard for organoid cocultures generated from individual synoviocyte and chondrocyte donor samples. DM, chondrogenic differentiation medium; Ctrl, control DMEM organoid medium supplemented with 10% FCS.

formulations for chondro-synovial cross talk studies. Results in Fig. 8A show that the chondrogenic medium significantly improves stable chondral organoid formation retaining over 90% of the initial organoid size up to day 8 with a gradual decline to 60.9 ± 8.8% at day 21 in comparison to the control organoids (p < 0.0001). Even though the presence of 1% FCS combined with 10 ng ml⁻¹ TGF- β 3 exhibited comparable values up to day 8, keeping 85 ± 1.28% of the initial chondral organoid area, variations in serum content and TGF- β 3 showed no significant difference to untreated control organoids at day 21 (p > 0.1, 2-way ANOVA with Dunnett's *post hoc* test) and were unable to outperform the commercial formulations.

In contrast to chondral organoids, where chondrogenic differentiation medium showed a positive effect on the

condensation of chondral organoids in coculture, the synovial organoid formation was severely affected by TGF- β 3 in combination with high serum content as well as chondrogenic medium formulation. For instance, 1 and 10 ng ml⁻¹ TGF- β 3 with 10% serum significantly reduced initial organoid compaction and cell-cell interaction at day 4 post-seeding generating deformed and often scattered organoids with 42% (p < 0.0002, 2-way ANOVA with Dunnett's *post hoc* test) and 55% lower initial organoid area (p < 0.0001), respectively. Moreover, commercial chondrocyte medium also significantly impeded proper organoid formation by 32% (p < 0.0001), as RA-FLS formed not even spherical aggregates but a network of tubular structures. This significant effect of serum and growth factor supplementation on organoid formation was observable for the first week of coculture.

However, it disappeared over the next two weeks and only synovial organoids in the commercial chondrogenic medium were still significantly impeded (p < 0.0001). Interestingly, the combination of 1% FCS and 10 ng ml⁻¹ TGF- β 3 led to a

significant increase in synovial organoid condensation (p < 0.0002) similar to their neighbouring chondral counterparts. To find a tentative explanation, secretion analysis (see Fig. 8B) was performed on coculture supernatants throughout



Fig. 9 Impact of organoid coculture on organoid formation dynamics over three weeks of co-cultivation. A) Representative images and B) organoid area analysis of on-chip monocultures in comparison to chondro-synovial organoid cocultures (scale bar = 2 mm. Data is expressed as mean \pm standard deviation for n = 2 organoids from single RA synoviocyte and 'healthy' chondrocyte donor samples). C) Confocal live/dead images of calcein-AM/ethidium homodimer-2 labelled mono- and cocultured chondral and synovial organoids at day 21 post-seeding. D) Fluorescence images and corresponding count analysis of CMTPX cell tracker labelled mono- and cocultured chondral and synovial on-chip organoids at day 4 and 21 post-seeding (n = 2 organoid cocultures from n = 1 synoviocyte and chondrocyte donor samples). E) Corresponding molecule secretion analysis over three week of chip cultivation for IL-6, IL-8, MMP-13, and VEGF at day 3, 6, 16 and 21 post-seeding. Normalised data is expressed as mean \pm standard deviation for n = 2 cocultures from n = 1 synoviocyte and chondrocyte donor samples.

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three weeks and showed no significant differences in secretion levels of IL-6, MMP-13 or VEGF except for the fact that chondrogenic differentiation medium showed reduced MMP-13 secretion levels and elevated IL-6 and VEGF levels, which were probably caused by the presence of proinflammatory cvtokine supplements (i.e., IL-6) or corticosteroids in the differentiation kit itself, which are known to boost chondrogenic differentiation further.⁴⁰ Also, TGF-β3 is frequently used as a trigger in synovial fibrosis models,²¹ and the effect of enhanced organoid instability with an increase in serum are in line with previous studies on synovial fibroblast models, showing that high serum conditions further exacerbate the aggravating effect of TGF- β 3.⁴¹ As our results indicated that there is a strong negative synovial performance when using media formulations and additives that are known to improve chondrogenic performance, further experiments concerning chondrosynovial cross talk were conducted in the absence of additives to investigate the baseline communication of primary synovial organoids generated from purified patient FLS with chondral organoids under very basic non-compromising conditions.

As a final experimental setup, our cocultures of synovial organoids made from patient-derived RA-FLS cultures were interfaced in coculture biochips with chondral organoids from primary human chondrocytes to establish and investigate the molecular effects of chondro-synovial cross talk (see Fig. 9). Chondral organoid monocultures condensed from 18.56 \pm 0.45 mm² to 0.65 \pm 0.33 mm² (2.0 \pm 4.3% of the initial organoid area), whereas synovial monocultures retained approx. 44% of the initial organoid area at day 2 from an initial organoid area of 8.72 \pm 0.88 mm^2 to 3.85 \pm 0.03 mm² throughout 21 days of monoculture (see Fig. 9A and B). Relative to the respective monoculture counterparts, synovial organoids in co-culture showed similar condensation dynamics with an approx. 4-fold decrease of the initial organoid area (p > 0.1, 2-way ANOVA with Sidak's post hoc test) from 9.7 \pm 0.6 mm² at day 2 to 4.14 \pm 0.03 mm² at day 28 post-seeding. Interestingly, the chondral compartment was more affected by tissue-tissue cross talk and showed a tendency to gain approx. 22% organoid area when cocultured with an organoid area of $2.9 \pm 1.3 \text{ mm}^2$ for cocultured in comparison to $0.65 \pm 0.33 \text{ mm}^2$ for monoculture chondral organoids at day 21. To investigate whether this effect was caused by alterations in cell viability, a live/dead dye exclusion assay as shown in Fig. 9C was conducted and showed comparable high viability of above 91% for chondral and 84% for synovial organoids independent of the cultivation scenario after three weeks of on-chip cultivation (see ESI† Fig. S5A). While epifluorescence analysis of calcein-AM stained monoculture chondral organoids revealed an overall homogenous population of chondrocytes (see Fig. 7C as reference), higher resolution confocal microscopy at a higher resolution for superficial areas near the organoid surface revealed higher round chondrocyte numbers for cocultured chondral organoids

compared to monoculture organoids, that also showed slightly elongated cell subpopulations. To better understand the correlation of this cellular difference with organoid condensation, mono and cocultured organoids were stained with CMTPX cell tracker before hydrogel loading at day 0 and organoids were investigated over 3 weeks. This specific cell tracker dye creates cell-impermeant and red-fluorescent reaction products by intracellular esterase hydrolysis. It provides strong and stable staining for chondrocytes over weeks and months. In contrast, highly proliferative cells such as dermal or synovial fibroblasts in our experience lose their fluorescence signal within a few days, especially when using standard epifluorescence microscopy analysis. Consequently, the loss of fluorescent signal of chondrocytes was investigated to quantify the change in chondrocyte morphology over time directly. Representative epifluorescence micrographs of chondral organoids in monoand coculture for 21 days (see Fig. 9D) revealed that the initial number of chondrocytes with round morphology at day 4 gradually declined for chondral monocultures, while round chondrocyte subpopulations were significantly more retained in the coculture with synovial organoids (p < 0.005, paired t-test) by 23 ± 2.3%. As shown in ESI† Fig. S5B, dedifferentiating chondrocytes in 2D cultures show approx. 300% increase in resazurin to resorufin conversion within 7 days of culture caused by increased esterase activity and cell proliferation. Chondral organoids showed a 2-fold lower conversion rate over the first seven days of culture, pointing at low proliferation and is in line with previous work comparing chondrocyte culture conditions.³¹ To investigate the influence of chondro-synovial cross talk on molecule secretion, Luminex analysis was used to compare monoculture with coculture organoids. As shown in Fig. 9E, the chondro-synovial organoid coculture system showed no significant changes in cytokine secretion levels of IL-6, IL-8 and MMP-13 (p > 0.1, 2-way ANOVA with Tukey's post hoc test) at day three with time-dependent a comparable decline similar to previous secretion curves of FLS organoids. However, chondral organoids in coculture, which were characterised before by a more chondrogenic environment with a higher degree of organoid area and chondrocyte roundness as well as lower metabolic actividisplayed significantly reduced VEGF secretion levels compared to chondral monocultures (p < 0.03, 2-way ANOVA with Sidak's post hoc test). This is in line with previous studies on VEGF secretion increase of chondrocytes cultivated under nonphysiological conditions.35,36,42

Conclusions

In the current study, we have, for the first time to our knowledge, established a chip-based chondro-synovial dual organoid model based on self-organising patient-derived cells that allow investigating reciprocal tissue-tissue cross talk in orthopaedic/arthritic research. Due to the intrinsic microfluidic design features, the current microfluidic joint-

on-a-chip design enables the reliable formation of organoids with high reproducibility and positioning precision, which is crucial for any organ-on-a-chip analysis system used in biomedical and biopharmaceutical research. We also demonstrated that expanded synovial fibroblast cultures generate organoids of higher precision and reproducibility compared to freshly isolated cell populations. This passaging effect that reduces the epigenetic functional characteristics of synovial subpopulations and results in synovial cell assimilation in vitro has been reported for the loss of lining versus sublining characteristics of FLS.⁴³⁻⁴⁵ We furthermore investigated chondrogenic and fibroblast medium conditions to rule out the effects of medium type on chondro-synovial cross talk as cocultures require a typical medium formulation. For instance, primary chondrocytes perform best in culture medium supplemented with TGF-\u03b33 at low or no serum conditions as it enhances chondrocyte redifferentiation⁴⁶ and stimulates extracellular matrix production.47 At the same time, proper cultivation environment for FLS and FLS micromasses encompasses high serum conditions with vitamin C supplementation. However, these chondrogenic factors severely influence synovial fitness (i.e., proliferation) in low concentrations of FBS and high TGF-\u00b33 concentrations and fail to recapitulate healthy synovial tissue function.⁴¹ In our current study, we followed a basal medium formulation with 10% FCS for the present proof-of-concept study; however, a more optimised medium formulation, most probably using xeno-free formulations, needs to be investigated to reduce the effects of molecules from the medium formulation for chondro-synovial cross talk. Finally, we could demonstrate that co-cultivation of chondral and synovial organoids improved the phenotype of chondrocytes within chondral organoids in comparison to chondral monocultures in the absence of synovial fibroblasts. We plan to investigate this molecular aspect and more disease-relevant mechanisms (i.e. induction of synovitis and pathway inhibition) in the future and believe that our current dual organoid-based joint-on-achip approach highlights the importance of tissue-level cross talk in the context of musculoskeletal diseases. For the current proof-of-principle study, we envision for the future of the chondro-synovial arthritis model to elaborate on the exact molecular cross talk mechanism between synovial and chondral organoids with a higher, statistically more significant patient sample size in combination with genomic and proteomic analysis.

Author contributions

AF, EIR, IOC, and SSch: experiments and data analysis; editing and reviewing – original draft. FS, JH and RW: surgical preparation of primary patient tissue; reviewing – original draft. SS, BB, WH and HR: assistance with fibrinbased organoids, editing and reviewing – original draft. HPK: reviewing and editing – original draft. RAB: experiments; data analysis; reviewing and editing – original draft. MR: supervision; experiments; data analysis; methodology; supervision, writing – original draft, review and editing. ST: writing – original draft, review and editing. PE: supervision; writing, review and editing – original draft; funding acquisition.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This project was supported by the Austrian Federal Ministry of Education, Science and Research (BMBWF; grant agreement number 1612889) and the Vienna Science and Technology Fund (WWTF, project number LS13-092). Figure schemes were created with BioRender.com under paid subscription. The authors acknowledge the TU Wien University Library for financial support through its Open Access Funding Program.

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