Autofluorescence of stingray skeletal cartilage: hyperspectral imaging as a tool for histological characterization

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Abstract
Tessellated cartilage is a distinctive composite tissue forming the bulk of the skeleton of cartilaginous fishes (e.g. sharks and rays), built from unmineralized cartilage covered at the surface by a thin layer of mineralized tiles called tesserae. The finescale structure and composition of elasmobranch tessellated cartilage has largely been investigated with electron microscopy, micro-computed tomography and histology, but many aspects of tissue structure and composition remain uncharacterized. In our study, we demonstrate that the tessellated cartilage of a stingray exhibits a strong and diverse autofluorescence, a native property of the tissue which can be harnessed as an effective label-free imaging technique. The autofluorescence signal was excited using a broad range of wavelengths in confocal and light sheet microscopy, comparing several sample preparations (fresh; demineralized and paraffin-embedded; non-demineralized and plastic-embedded) and imaging the tissue at different scales. Autofluorescence varied with sample preparation with the signal in both plastic- and paraffin-embedded samples strong enough to allow visualization of finescale (≥ 1 μm) cellular and matrix structures, such as cell nuclei and current and former mineralization fronts, identifiable by globular mineralized tissue. A defined pericellular matrix (PCM) surrounding chondrocytes was also discernible, described here for the first time in elasmobranchs. The presence of a PCM suggests similarities with mammalian cartilage regarding how chondrocytes interact with their environment, the PCM in mammals acting as a transducer for biomechanical and biochemical signals. A posterior analysis of hyperspectral images by an MCR-ALS unmixing algorithm allowed identification of several distinct fluorescence signatures associated to specific regions in the tissue. Some fluorescence signatures identified could be correlated with collagen type II, the most abundant structural molecule of cartilage. Other fluorescence signatures, however, remained unidentified, spotlighting tissue regions that deserve deeper characterization and suggesting the presence of molecules still unidentified in elasmobranch skeletal cartilage. Our results show that autofluorescence can be a powerful exploratory imaging tool for characterizing less-studied skeletal tissues, such as tessellated cartilage. The images obtained are largely comparable with more commonly used techniques, but without the need for complicated sample preparations or external staining reagents standard in histology and electron microscopy (TEM, SEM).

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Abbreviations
AF  Autofluorescence
PCM  Pericellular matrix
ECM  Extracellular matrix
UC  Unmineralized cartilage
MCR-ALS  Multivariate Curve Resolution Alternating Least Squares
EEM  Excitation–Emission
WLL  White Light Laser
BSEM  Backscatter electron microscopy
EDTA  Ethylenediaminetetraacetic acid
PFA  Paraformaldehyde
PMMA  Polymethyl methacrylate

1 Introduction

Cartilage is a viscoelastic and resilient connective tissue essential to the mechanics and growth of the skeleton of vertebrates. Its gel-like extracellular matrix is composed of water and contains embedded cells (chondrocytes) interspersed within a network of mainly collagen type II fibers and proteoglycans [1, 2]. Variation in the organization, proportion, and composition of extracellular matrix components is reflected in the structural morphology of the tissue, affecting its mechanical properties and its physiological functions [3, 4]. Examinations of cartilage tissue morphology, therefore, have provided great insights into the biomechanics, function and growth of cartilage. Most of this research has focused on mammals, where cartilage (as in most vertebrates) forms the greater part of the embryonic skeleton, providing a tissue scaffold that guides the growth of and eventually gives way to bone [1]. Cartilage is far less predominant in the adult skeleton of most vertebrates, but still performs diverse roles including load-bearing (intervertebral fibrocartilage); shaping of resilient structures like noses, ears and ribs (elastic cartilage); and reduction of frictional forces between bones (articular hyaline cartilage).

Cartilage is, also, far more diverse and ancient than typically appreciated. It is present even in invertebrate taxa [5, 6] and forms an extremely large proportion of the adult skeleton of several fish groups, particularly the clade of sharks and rays (elasmobranch fishes) [7] (Fig. 1). The tissue composition and the cell type of elasmobranch cartilage appear to be similar to the hyaline cartilage in mammalian joints [8–11 and references therein]. However, elasmobranch cartilage performs many functions more akin to those of bone than mammalian hyaline cartilage: providing skeletal shape, body support, and locations for muscle attachment. This multi-functionality is believed to be due to a unique structural feature of elasmobranch cartilage, wherein the skeleton mineralizes only at the surface, forming an ensheathing layer of mineralized tiles called tesserae (Fig. 1B–E).

The mechanical performance of tessellated cartilage is thought to be related largely to the structure and arrangement of its different tissue components, in particular the associations of soft (unmineralized) and hard (mineralized) materials [12–15]. However, the difficulties associated with characterizing the complex structural arrangements of these heterogeneous tissues has limited the definition of clear composition-form-function rules for tessellates cartilage. In particular, the close association of mineralized and unmineralized tissues, in complex structural arrangements at both large and fine scales, has challenged common visualization techniques, such as conventional microscopy, immuno-histological stainings and electron microscopy [8, 16]. For example, the mineralized matrix of tesserae is perforated by numerous small cavities called lacunae, containing living chondrocytes surrounded by a thin layer of unmineralized matrix, with adjacent lacunae linked by short, matrix-filled passages [17, 18] (Fig. 1C–E). This arrangement results in several tissue type transitions over small spatial scales (1–10 µm), also challenging proper fixation and sectioning of these regions. The lateral margins of tesserae also present a heterogeneous mixture of tissues, associated with the joint region between neighbouring tesserae, which comprises a complex combination of unmineralized matrix, fibers and cells [8] (Fig. 1E). Additionally, the high water content of the unmineralized cartilage, contrasted with the extremely low water content (high mineral content) of some regions in tesserae (e.g. the hypermineralized ‘spokes’ reinforcing contact zones, Fig. 1C, E; [19]) makes tessellated cartilage prone to differential swelling and cracking during standard fixation and dehydration.
protocols [12]. Lastly, the scarcity of antibodies and protocols optimized for tissue components (e.g. cell processes, collagen types) in non-model organisms makes the characterization of tessellated cartilage tissue components both time-consuming and costly [20].

Label-free imaging—in other words, not using exogenous fluorophores—has recently proved successful in the study of cartilage composition and structure, using both confocal (linear excitation) and multiphoton (non-linear excitation) microscopy. In label-free imaging, image contrast is provided by the tissue’s autofluorescence (AF): the excitation of intrinsic endogenous fluorescent molecules (e.g. coenzymes and amino acids, among others) rather than by the addition of external labeling or contrast agents (e.g. fluorescent or non-fluorescent dyes). Endogenous fluorescent molecules contain aromatic rings with Pi-bonds, with electrons that react to specific illumination wavelengths, passing to an excited state before decaying to the ground state and emitting energy, including the autofluorescent emission [21]. As tissue AF can be diagnostic for particular tissue components, AF imaging has been demonstrated to be a powerful resource especially in mammalian systems, for evaluating biological substrate structure and composition, from cells to entire organisms, and can provide diagnostic information in a minimally invasive approach (reviewed by [22]. Nonlinear optical microscopy techniques (NLOM, allowing excitation of multiple endogenous fluorophores with a single laser line) have been particularly successful in the minimally-invasive examination of cartilage, using only endogenous fluorophores (e.g. flavoproteins, NAD(H)P, collagens and elastin) to characterize the organization and orientation of components within

Fig. 1  Tessellated skeleton of sharks and rays (elasmobranchs). A Image of the common stingray Dasyatis pastinaca (left) and a microCT scan image of the skeleton (right), with the propterygium highlighted in red. B MicroCT scan of bisected propterygium, showing the inner cartilaginous core (blue) surrounded by the tessellated layer (gray). C Backscatter Scanning Electron Microscopy (BSEM) image of a planar section through multiple tesserae, showing spokes (white streaks) and lacunae (black spaces) within tesserae. D Schema of the propterygium cross section from B, showing also the outermost perichondrial layer (red). E Schema of zoomed region in D, showing tesserea in vertical section, sandwiched between the perichondrial layer and the cartilaginous matrix. The single tessera in the center is connected to neighboring tesserae by a joint region (marked with a rectangle). Cells are located within tesserae, the cartilaginous matrix and joint regions. Chondrocytes (C), Cartilaginous matrix (CM), Joint (J), Muscle (M), Perichondrium (P), Spokes (Sp), Tesserae (T)
the tissue [23, 24]. By comparison, linear excitation techniques have been less used to study cartilage, although they offer broader excitation and emission spectra than NLOM.

In this study, we explore the efficacy of linear, one-photon excitation methods in the imaging of tessellated cartilage, examining also whether these effects are altered by standard sample preparation techniques. AF imaging techniques have never been applied as a tool to study this tissue, however there is some evidence that particular regions of tesserae exhibit a characteristic AF [11]. Previous studies have shown that tesserae have distinct and consistent structural features and, thus, can be considered as a useful and repeatable unit of study, particularly since every skeletal cross-section contains multiple sectioned tesserae for examination [18, 25]. Additionally, since tesserae have been shown to grow by accreting mineralized material on all their edges and do not appear to remodel their tissue at all, every tessera contains a permanent structural record of its growth [25–27]. Here, using common imaging techniques (e.g. histology, electron microscopy and microCT) and recent ultrastructural characterizations of tesserae as reference, we demonstrate that specific tissue components have consistent and characteristic AF signals, using confocal laser scanning microscopy of thin sections and light sheet microscopy of larger sample blocks to provide a more macroscopic perspective. In the process, we explore the potential of AF imaging for ultrastructural examination of elasmobranch skeletal tissue, especially to clarify aspects of fibrous tissue architecture.

2 Methods/Experimental

2.1 Specimens and sample preparation

Skeletal samples were collected in May 2019 from two common stingrays (Dasyatis pastinaca: Dasyatidae, Batoidea), a male of 21.3 cm disc width (DW) and a female of 32.7 cm DW. Specimens were by-catch captures from trammel net fishery in Alicante and Valencia (Valencian Community, Spain), respectively. Samples were harvested from both specimens less than 1 h after death.

We compared tissue autofluorescence (see below) for three sample preparation types that have been used previously in the study of tessellated cartilage (Fig. 2A): (1) “thawed samples”: non-fixed, non-demineralized and non-embedded; (2) “paraffin samples”: fixed, demineralized and paraffin-embedded; and (3) “PMMA samples”: fixed, non-demineralized and embedded in plastic (PMMA). For the thawed samples, the whole head of the second specimen was frozen and sent to the Institute of Photonic Sciences’ facilities (Castelldefels, Spain), where all imaging was performed. Immediately prior to imaging, the head was partially thawed in cool water and a scalpel was used to remove thin slices (~0.5–1.0 mm) of the proptrygium (largest skeletal piece supporting the wing). Slices were mounted for confocal imaging on a glass bottom MatTek dish in solidified 1%, low-melting agarose, to maintain position and reduce dehydration.

Samples for paraffin and PMMA were obtained from the first animal specimen. The proptrygium was cut in small pieces and fixed with 4% PFA in PBS 0.1 M at the harbor and within 20 min of the animal's death to avoid tissue degradation. Samples were kept in the fixative at 4 °C overnight and stored in PBS 0.1 M at 4 °C before processing for paraffin embedding, PMMA embedding or clearing for SPIM imaging (see next section).

For paraffin embedding, fixed samples were rinsed in PBS, decalcified with ethylenediaminetetraacetic acid (EDTA) at 20 wt-% concentration with pH 7.4 for one week, dehydrated with a graded isopropanol and xylene series, and embedded in paraffin using the reagents and protocol provided by the manufacturer. All embedding steps were performed at 4 °C. Once the embedding media was polymerized, samples were microtomed in 10 µm slices, deplastified using 2-methoxyethyl acetate, and mounted on DakoFlex® slides.

For PMMA (Polymethylmethacrylate) embedding, fixed samples were dehydrated through increasing ethanol concentration (70%, 80%, 96%, 100%), then maintained in xylol for 3 h, and finally embedded in PMMA (Morphisto Technovit® 9100) using the reagents and protocol provided by the manufacturer. All embedding steps were performed at 4 °C. Once the embedding media was polymerized, samples were microtomed in 10 µm slices, deplastified using 2-methoxyethyl acetate, and mounted on DakoFlex® slides.

To compare AF images with those obtained using more traditional methods and to facilitate location of structures of interest, tessellated cartilage samples were also examined using histology and backscatter electron microscopy (BSEM). The sample preparation protocols for each technique can be found in section 1 of Supplementary Materials.
2.2 Confocal imaging of thawed, paraffin and PMMA samples

Cartilage samples from all three preparation types described above were examined using a Leica TCS SP8 STED 3x (Leica, Mannheim, Germany) Laser Scanning Confocal Microscope (LSCM) with an HC PL APO CS2 63x/1.40 oil objective. Samples were excited using two lasers and imaged in a wide spectral range that allowed a broad spectral analysis of autofluorescence signal (Fig. 2B). Three types of images (4D, 3D and 2D images) were generated in each imaged region of interest, as described below.

To generate 4D images, samples were excited with the supercontinuum White Light Laser (WLL) in xyλλ (lambda-lambda scan) acquisition mode in a range of $\lambda_{ex} = 470$ to 670 nm in 21 excitation steps with 10 nm sampling interval (400 Hz scan speed, frame and line accumulations = 1). At each step, fluorescence spectra were collected using a HYD SMD detector in a range of $\lambda_{em} = 490$ to 775 nm in 20 emission steps with a 14.25 nm sampling interval. In this way, we obtained a fluorescence emission hyperspectral image for every excitation step, resulting in a 4D excitation-emission fluorescence hyperspectral image (EE-HSI) with two spatial dimensions (x- and y- pixel coordinates) and two spectral dimensions (excitation and emission), where each pixel was associated with a 2D excitation-emission fluorescence landscape (i.e. quantifying the relationship between excitation and emission wavelengths over a broad spectral range) (Supp. Figure 1B).

Additionally, to generate 3D images, samples were excited with a 405 nm laser in xyλ (lambda scan) acquisition mode, with fluorescence spectra collected using a HYD SMD detector in a range of $\lambda_{em} = 422.5$ to 702.5 nm with a 14.74 nm sampling interval (400 Hz scan speed, frame and line accumulations = 1). Resulting 3D images had two spatial dimensions (x- and y- pixel coordinates) and one spectral (λ) dimension, with each pixel associated with an emission fluorescence spectrum (i.e. quantifying the autofluorescence produced by the 405 nm laser) (Supp. Figure 1A).

Lastly, we acquired two types of 2D images as reference images. Firstly, an individual, high-resolution optical image (pixel size = 1.71 nm) of each region of interest (see below) was acquired using the excitation wavelength found experimentally to produce the strongest AF (405 nm laser with 200 Hz scan speed, two frame and line average, 20% intensity) and a wide detector range ($\lambda_{em} = 459$ to 602 nm, with the HYD SMD detector). The resultant image
was two-dimensional, with x- and y-pixel coordinates and an integrated global AF value assigned to each pixel. In order to prevent loss of signal by bleaching and longer dwell-times, analyses were always performed in the following order: WLL lambda-lambda scan, followed by the 405 nm lambda scan, and then 405 nm high-resolution images. Additionally, 2D transmitted light images were acquired simultaneously with the confocal images using the same illumination beam, to ensure that images were in registration.

2D, 3D and 4D images were acquired in three specific regions of interest (ROIs): the tesserae, the joints, and the unmineralized cartilage. Additional 2D, 3D and 4D images were made of the embedding media (paraffin, PMMA) without samples, to determine if these media contributed any AF. We replicated the analyses of each ROI type three times for each sample preparation type (i.e. for PMMA, paraffin and thawed samples, scanning three different tesserae, joints, and unmineralized cartilage regions each).

It was challenging in all sample preparations to find a flat region of interest for high magnification imaging (63), especially at the interface between tesserae and unmineralized cartilage. Unembedded thawed samples were particularly difficult to prepare and image. As this tissue was not embedded, non-uniform sections were cut manually by razor blade and therefore were not uniform. Also, evaporation of water from the tissue (e.g. the unmineralized cartilage, which has a high water content) caused large shifts in the position and morphology of the tissue, making imaging almost impossible at high magnification. Thus, the images of thawed tissue were only taken at magnification of 10x, with the exception of one tessera sectioned flat enough to be imaged at 63x.

### 2.3 Analysis of hyperspectral images. Underlying image analysis model and unmixing procedure

Unlike 2D images, where features can be interpreted directly by their global fluorescence intensity, 3D and 4D images require dedicated data treatments by unmixing methods to disentangle the contributions of the distinct fluorescence signatures in a sample. Indeed, natural fluorophores can exist in different concentrations and locations within an analysed tissue, which can result in spatially complex variation and/or mixing (co-localization) of fluorescence signals, often hindering the interpretation of results. The ability to unmix multivariate signals in fluorescence images is particularly important for 3D images and, most importantly, for 4D images, which have the most potential to detect and distinguish biological regions with distinct fluorescence signatures, due to the wider excitation range used and because differences in both excitation and emission characteristics are considered.

The unmixing algorithm used in this work is Multivariate Curve Resolution-Alternating Least Squares (MCR-ALS), apt for single 3D and 4D images, but also for the treatment of several related images analysed together (multiset analysis) [28, 29, 80, 81]. The results provided by MCR-ALS are distinct fluorescence signatures (emission spectra or excitation/emission fluorescence landscapes for 3D and 4D images, respectively), from either an individual fluorophore (e.g. collagens, elastin, flavins) or a mixture of fluorophores found consistently together (e.g. in a particular tissue type). The resultant distribution maps produced by the MCR-ALS show the spatial distribution of the fluorescence signatures in the 3D/4D source images and therefore their associations with tissues and other biological entities [30]. In this work, all MCR-ALS analyses were performed using in-house implemented routines, programmed in the MATLAB environment (v. 2019) [31].

A detailed description of the function and output of the MCR-ALS is outlined in Section 2 of Supplementary Materials.

### 2.4 Light sheet microscopy of fixed samples

To examine tessellated cartilage in larger samples without the need for thin sectioning, we also employed light-sheet fluorescence microscopy (LSFM), also known as Single-Plane Illumination microscopy (SPIM) [32], using AF as a contrast mechanism. In contrast to the point-laser illumination of the confocal microscope, SPIM relies on the creation of an illumination plane, with the laser beam shaped into a rectangle and focused in a thin “sheet of light” using a cylindrical lens [32, 33]. The uniqueness of its optical configuration, with uncoupled perpendicular illumination and detection axis, combined with fast camera-based detection, allows acquisition of 3D images of samples at unprecedented speeds.

We analyzed the AF signal of *D. pastinaca* tessellated cartilage using a custom made SPIM microscope [34]. The custom microscope achieves magnifications between 2× and 4× (field of view 6.65 × 6.65 mm² and 3.33 × 3.33 mm², respectively), allowing the imaging of samples several millimeters thick. This can be combined with image stitching to allow volumetric imaging of samples several centimeters thick in each linear dimension. In our study, the examined samples were thick cubes of propterygium (5–6 mm thick), fixed as described in the previous section. To facilitate the imaging of larger sample blocks of tessellated cartilage, samples needed to be clarified (i.e. rendered transparent) to equalize the refractive index throughout the tissue block. No protocols exist for clarification of tessellated cartilage; we used ECI (ethyl
cinnamate), which has proved effective for clearing hard tissues in calvaria and long bones [e.g. 35], while also being fast, affordable and non-toxic. For clearing, fixed transverse sections were dehydrated with an ethanol series (pH 9.0) 50–70–100% (2X), 12 h per step, at 4ºC. After dehydration, samples were transferred to ECi (W243000 Sigma-Aldrich) and incubated while gently shaking at room temperature until they became transparent. The excitation and emission spectra used in SPIM made it compatible with our analyses made with confocal. Samples were excited with several excitation wavelengths (λex = 405, 488 and 561 nm) and the emitted fluorescence collected with filters of 482/25, 520/50 nm and 593 long pass, respectively.

3 Results

3.1 General observations of AF signal

Using AF, it was possible to obtain high resolution optical images of different tissue features of tessellated cartilage, including both mineralized (tesserae) and unmineralized elements (perichondrium, cells, the unmineralized cartilage matrix), although the degree of detail in the images varied with sample preparation (see below). The 405 excitation laser with a broad spectrum of acquired emission wavelengths (459–602 nm) produced the clearest imaging results with the strongest signal, comparable to other techniques commonly used to image tessellated cartilage, such as histology and BSEM (Fig. 3).

3.2 Effects of sample preparation

The structures and tissues visible from their AF in tessellated cartilage differed among PMMA samples, paraffin samples, and non-embedded thawed samples (Figs. 4, 5). Examination of the AF signal of the embedding media themselves, showed negligible AF (PMMA) or none at all (paraffin), and so did not contribute to tissue AF signals. The discernibility of particular tessellated cartilage features is discussed in detail in the following section, but some general effects of sample preparation are worth noting.

For unembedded thawed samples, the propterygium was thawed a first time to perform a preliminary test of the sectioning and imaging methods, and then a second time months later for the final experimental analysis. The preliminary images showed AF signal in perichondrium, tesserae, and unmineralized cartilage (both cells and matrix) (Fig. 4Aa, [d–g], C). In contrast, in the later experimental analysis, only a weak AF signal was visible in the tesserae, being entirely lost in the soft tissues (Fig. 4A[b, c], C).

PMMA samples were non-demineralized and more cleanly sectioned than thawed samples and therefore showed both hard and soft tissues in conditions perhaps closest to their native configurations (Figs. 5, 6, 7). Tesserae in PMMA samples, however, often exhibited cracks (Figs. 5D; 6A), making it difficult to obtain large, intact regions of the tesserae layer. The cracks were produced during the preparation of the samples, probably during microtome cutting, oven-curing,
The features of joints (i.e. the structures and tissues between tesserae) were not visible in thawed samples, likely associated with the difficulty of producing samples flat enough for high magnification imaging (Fig. 4A). In PMMA, paraffin and clarified samples, however, the ultrastructures of multiple joint-associated features could be identified using AF (Fig. 5). Depending on the particular plane in which neighboring tesserae were sectioned, two local morphologies were possible, often with both present in a single joint ROI: (1) the tesserae margins separated by a space containing unmineralized matrix, cells and fibers (Figs. 5F; 6A, 8B) or (2) the tesserae margins in direct contact (Fig. 5G). When tesserae were in direct contact, laminated, highly mineralized features called spokes [19, 25] were visible in PMMA samples, radiating or mounting of slices. Similar cracking has been seen in several previous studies of tesserae, particularly associated with high mineral density regions (e.g. tesseral spokes; see below) [25]. In contrast, in paraffin sections, demineralization resulted in more intact tesserae, not disrupted by preparation artifacts, yet needless to say, mineralized tissue details were lacking (Fig. 5).

3.3 Observations by individual feature

1. Tesserae

The tesserae layer could be imaged in all sample preparation methods, with the intensity of the signal always higher than in the unmineralized cartilage (Figs. 4, 5, 6, 7, 8). However, the visible features and morphology varied. PMMA samples, for example, showed the most clearly defined tesserae morphology and mineralized structures (Figs. 5, 6). Within the tesserae, a distinct central region delineated by a bright autofluorescent border showed a stronger AF signal than the peripheral region at the margin of tesserae (Figs. 5C, D; 6A). These three AF zones—center, center border and periphery—were also observed in clarified samples imaged with SPIM (Fig. 8), yet were not visible in histology samples (Figs. 3B; 5B). In the central region, cell lacunae were surrounded by wavy, concentric lines in the mineralized matrix (Fig. 6A, D, E). These lines were identified from our histology and BSEM images as Liesegang lines (lines of accretive tissue growth; [25]), and were also observed in other tesserae regions (e.g. just interior to and tracking the contours of the outer margins of tesserae; Fig. 6C), albeit with lower signal intensity (Fig. 6D, E).

Cells within tesserae are enclosed in lacunae and surrounded by unmineralized matrix, with lacunae connected by small passages (canaliculi). In our examinations, cells within lacunae were clearly visible in PMMA and paraffin samples, however, in contrast with cells in the unmineralized cartilage and joints, appeared more shrunken (e.g. damaged during fixation), and therefore their morphologies could not be properly visualized. A brighter haze of AF was often seen surrounding and connecting the cells within tesserae (Fig. 6B), but it was unclear whether this corresponded to the unmineralized matrix filling the cavities surrounding and linking cells (lacunae and canaliculi, respectively) or to the mineralized tissue that bordered these cavities, or some combination of the two. Similarly, the interface between tesserae and unmineralized cartilage (i.e. at the lower/chondral margins of tesserae) was difficult to detect in paraffin (demineralized) samples as the two materials had nearly the same gray values in the 405 images (Fig. 6F). In contrast, in PMMA samples, the interface was typically a thin, high-brightness rim marking the mineralization front, which made it easier to distinguish (Fig. 6D, E).

Small autofluorescent globular features (~1–2 μm) were often visible in PMMA samples at interfacial locations in tesserae: at the border between the central and peripheral regions of tesserae (Fig. 6A), along the interior walls of cell lacunae (Fig. 6B) and surrounding cells that were in the process of being engulfed by tesserae at the interface with the unmineralized matrix (Fig. 6D). These globules corresponded in size and location to mineralized globular features observed in BSEM, but were not seen in other techniques where tissue had been demineralized (e.g. paraffin).

2. Joints

The features of joints (i.e. the structures and tissues between tesserae) were not visible in thawed samples, likely associated with the difficulty of producing samples flat enough for high magnification imaging (Fig. 4A). In PMMA, paraffin and clarified samples, however, the ultrastructures of multiple joint-associated features could be identified using AF (Fig. 5). Depending on the particular plane in which neighboring tesserae were sectioned, two local morphologies were possible, often with both present in a single joint ROI: (1) the tesserae margins separated by a space containing unmineralized matrix, cells and fibers (Figs. 5F; 6A, 8B) or (2) the tesserae margins in direct contact (Fig. 5G). When tesserae were in direct contact, laminated, highly mineralized features called spokes [19, 25] were visible in PMMA samples, radiating...
A) Preliminary analysis

Tesseract

P, T, CM

20 µm

Experimental analysis

B) Preliminary analysis

CM

BF

λ_ex = 405 nm / λ_em = 418 - 585 nm

λ_ex = 470 nm / λ_em = 482 - 690 nm

λ_ex = 520 nm / λ_em = 535 - 646 nm

C) Preliminary analysis

λ_ex = 470 nm

Fluo. 1

Fluo. 2

Normalized emission

Wavelength (nm)

420 450 500 540 580

0.4 0.3 0.2 0.1

Fluo. 3

Fluo. 4

Normalized emission

Wavelength (nm)

450 550 650 540 580 620

0.4 0.3 0.2 0.1

Fluorescence Emission

Emission wavelength (nm)

500 600 700

0 40 80

500 600 700

0 40 80
from contact points between neighbouring tesserae. These structures exhibited high AF signal and showed a similar morphology to spokes seen in BSE images (Fig. 5G). In both histology and AF of paraffin samples, spokes were only represented by gaps in the tissue where the high mineral density laminae had been prior to demineralization (Fig. 5C, E).

In zones where neighboring tesserae were separated by soft material, cells and fibers connecting tesserae were typically visible. Cells located in the joint space were flattened and arranged in series, passing between neighboring tesserae (Figs. 5E, F; 6A; 8B). In our histology, these cell arrays occupied horizontal gaps between the fiber bundles linking tesserae; using the AF signal, these fibers were often visible as parallel streaks, connecting tesserae and framing cell arrays (Fig. 6A; 8B).

3. Unmineralized cartilage

The unmineralized cartilage comprises cells embedded in copious amounts of extracellular matrix, as visible in our histological samples (Figs. 3; 5H). This basic tissue arrangement was also visible in AF imaging of all sample preparations, where cells emitted a brighter AF signal than the surrounding matrix (Figs. 3D; 5I, J; 7A, C). Cells in the unmineralized cartilage also appeared more intact than cells within tesserae (see above), allowing observation of some local variation in cell morphology and arrangement across different zones (Fig. 7). In the unmineralized matrix directly beneath tesserae, cells were either well separated or formed twins (closely-associated, isogenous pairs, indicating recent separation by mitosis; Fig. 7D–F). In this zone, the cells exhibited a variety of cross-sectional morphologies, spheroidal (typical of chondrocytes), but also flattened or elongated (Figs. 5I, J; 7F). In contrast, further from the interface with tesserae (i.e. deeper into the unmineralized matrix), cells always presented spheroidal morphologies and often formed isogenous groups (clusters of sister cells) of two or four chondrocytes (Figs. 5H–J; 7D–F; 8C).

Among the sample preparation methods, cells differed in their morphology and the fluorescence signal emitted. In the preliminary analysis of thawed samples, cells emitted a strong AF signal, in comparison to the very low signal of the surrounding matrix (Fig. 4A); both sources of AF were completely lost in the final experimental analysis. Cells in PMMA and paraffin samples exhibited different degrees of shrunkeness, where the cell appeared collapsed away from the walls of its lacuna and surrounded by an empty (black) space [36] (Fig. 7A). Better preserved cells exhibited different appearance and cell structure depending on the sample preparation method. In paraffin samples, these cells were bright ovoids with a darker rounded nucleus and relatively homogeneous cytoplasm (Fig. 7C). In PMMA samples, these cells showed a thin, bright surrounding border and a bright nucleus, but the intracellular compartment had a loose foamy appearance (Fig. 7B, E). Outside the cell, a distinct pericellular region was often visible, typically with a lower grayscale value and different matrix texture (Fig. 7B, E).

3.4 Analysis of hyperspectral images

For both 3D and 4D fluorescence hyperspectral images, three separate multisets were created (one per sample preparation method), formed by combining preprocessed fluorescence images from all ROIs (tesserae, joints and UC) for a given sample preparation method (see Fig. 9, 10, 11; Suppl. Figure 1–4). MCR–ALS was then applied to the 3D and 4D multisets to elucidate the sample constituents present in the analyzed sections of cartilage tissue. The primary parameters and model results of MCR–ALS, as applied to the different multisets analysed, are summarized in Table 1, including the number of images analyzed per multiset, the number of fluorescence signatures resolved by MCR-ALS, and estimates of model quality parameters: lack of fit (LOF) and explained variance. The variance explained by the MCR-ALS models is...
satisfactory considering the noise level of the spectra analysed. The lower variance explained by 4D images compared with 3D images is explained by the lower signal-to-noise ratio of these data. The unmixed fluorescence signatures detected in our samples by the MCR-ALS were characterized by distinct pure excitation-emission spectra and distribution in the tissue. To facilitate description and reference in our analysis, we designate the observed unmixed fluorescence signatures “Fluorophores”, however, it is important to note that these AF signatures can be produced either by a single fluorophore (e.g. from elastin or a particular collagen or flavin) or a co-occurring combination of them. We number the fluorophores arbitrarily for reference (e.g. Fluorophore 1, 2, 3) and in the text below provide general summaries of their excitation/emission ranges; these data are listed in more spectral detail and according to tissue associations, sample preparation and imaging multiset (3D or 4D) in Table 2.

4D images showed a higher number of distinct signatures than 3D images, likely due to the wider excitation range. Regarding the preparation method, PMMA samples showed a higher number of distinct signatures, followed by paraffin. Thawed samples presented a weak AF signal and only preliminary samples exhibited a signal-to-noise ratio that allowed identification of distinct AF signatures.

### 3.5 Paraffin sample analysis

In paraffin samples, four fluorescence signatures were detected with MCR-ALS, one with 3D images (Fluorophore 1) and three with 4D images (Fluorophore 2, 3 and 4). Fluorophore 1 was excited by the 405 laser (3D multiset: Table 2), observed
in the perichondrium, tesserae, joints and cartilage matrix, although with intensity variations across the regions (Fig. 9A). The highest intensities were seen in the perichondrium and within the tesserae.

The tesserae—with this fluorophore, but also others below—presented distinct spatial variation in signal intensity, allowing three distinct tesseral regions to be easily discerned from each other: a center region, a peripheral region surrounding the center concentrically, and a thin edge at the extreme outer border of the tesserae (~ 1 μm thick). These three regions—center, center border, and periphery—were similar in general appearance to those identified in 2D AF images and SPIM (Figs. 8C, 9). The AF of the central region and the tesseral outer edge (especially where it bordered the joint region) was more intense than in the peripheral region. In the joints and unmineralized cartilage, this AF signature was weak in the extracellular matrix, but stronger within the cells. Some cells presented local areas of higher intensity, apparently in the cytoplasm.

Fluorophores 2, 3 and 4 (4D fluorescence images) were excited with the WLL in a broad excitation range from 470 to 670 nm. Fluorophore 2 and Fluorophore 3 presented similar excitation-emission profiles, however, the MCR-ALS algorithm could unmix them based on their spectral shape differences (Table 2; Fig. 9B).

Fluorophore 2 was associated with soft tissues, in the extracellular matrices and cells of joints and unmineralized cartilage, but absent in tesserae and perichondrium. The highest signal intensities were seen within the cells of the joints, followed by the cells of the unmineralized cartilage. In the tesserae, some lacunar spaces of the central region also exhibited signals of relatively high intensity.

Fluorophore 3 had a wide distribution, in the perichondrium, the laminae of spokes in tesserae, the cartilage matrix, and in all cells except those of the central region in tesserae. Whereas the extracellular matrices of both joints and cartilage display Fluorophore 2, Fluorophore 3 was only located within the cartilage matrix.
Fluorophore 4 was the only fluorescence signature in this multiset located in the mineralized tissue of tesserae: in the border between the central and peripheral regions of the tesserae and in the outer edge surrounding the tesserae. Fluorophore 4 was also located within the cells of all regions (tesserae, joints and unmineralized cartilage). Although Fluorophores 2, 3 and 4 of this multiset were localized in cells, Fluorophore 4 showed the highest cellular AF and was absent in the extracellular matrices of the joints and unmineralized cartilage.

3.6 PMMA sample analysis

In PMMA samples, five fluorescence signatures were detected with MCR-ALS, two in 3D images (Fluorophores 1 and 2) and three in 4D images (Fluorophores 3, 4 and 5). Fluorophores 1 and 2 were excited by the 405 laser, (Table 2; Fig. 10). Fluorophore 1 was homogeneously distributed in the mineralized matrix of the tesserae, and in the extracellular matrix of the joints and unmineralized cartilage, but absent in all cells. Within the tesserae, this fluorescence signature was more intense in the laminae of spokes. The localization within the tissue of this fluorophore correlates with the distribution pattern of Coll2 observed in IHC (Supp. Figure 5). Fluorophore 2 had an intense signal in the muscles, perichondrium, center region and outer edge of the tesserae and in the cells within the joints and cartilage.

Fluorophores 3, 4 and 5 were excited with the WLL (4D images) in a broad excitation range from 470 to 670 nm (Table 2, Fig. 11). Fluorophore 3 was most intense in muscle, perichondrium, and spokes' laminae, exhibiting lower intensities within the cells and extracellular matrix of the unmineralized cartilage. In the extracellular matrix, this fluorescence signature had a homogeneous distribution, albeit forming higher intensity halos immediately surrounding cells. Fluorophore 3 was lacking in joints and the majority of tesserae cells.
Fluorophore 4 had the highest intensity in muscle. It was also present in the perichondrium, in the border between central and peripheral regions in tesserae, and in the outer edge of tesserae. In the unmineralized cartilage, it was only present within the cells, being completely absent in the extracellular matrix. Fluorophore 5 was most intense in the muscle and perichondrium, in the center, center border and outer edge of the tesserae and, to a lesser degree, in the unmineralized cartilage matrix. A weak signal could be seen within the cells of the unmineralized cartilage, but was absent in the rest of the cells.

Consolidating the information from all fluorescence signatures observed in PMMA samples, it can be noted that the cells differ in signal according to their location: cells of the unmineralized matrix present a combination of Fluorophores 3, 4 and 5, whereas cells of the joints and tesserae exhibit only Fluorophore 4, and Fluorophore 3 dominates in cells of the tesserae body zone.

3.7 Thawed samples analysis

For comparison of the AF signals of preliminary and experimental thawed samples, the different instrumental settings (e.g. bandwidth and pixel size) of the two experiments were standardized by binning to unify pixel size and by selecting only common spectral excitation wavelengths (470 and 520 nm). For both of these excitation wavelengths, the AF found in preliminary samples was nearly absent in the experimental thawed samples (Fig. 4C).

In contrast, the MCR-ALS analysis of preliminary thawed samples excited by the 405 nm laser (3D multiset) and by the 470 and 520 nm lasers (4D multiset) allowed tissue features to be distinguished. Following binning of the hyperspectral images by a factor of seven to increase spectral quality (increasing pixel size from 1.06 μm to 7.44 μm), two different fluorescence signatures were detectable, excited by the 405 nm laser (Fig. 4B, Table 2). Fluorophore 1 was generally located in the peripheral region of tesserae, whereas Fluorophore 2 was constrained largely to the central region. In this case, we excited the tissue with individual 470 and 520 nm lasers instead of a broader range, but this was still considered as a single excitation range. With these excitation wavelengths, two fluorescence signatures (Fluorophore 3 and 4) with different emission spectra were associated with each excitation wavelength (Table 2, Fig. 4B). Both fluorescence signatures were associated with the tesserae, presenting distinct spatial locations; Fluorophore 3 was generally associated with the peripheral region of the tesserae, whereas the Fluorophore 4 was associated with the central region. The detection of fluorescence signals in thawed samples indicates the presence of natural fluorophores associated with distinct spatial locations of the tissue.

3.8 Tissue AF imaging at larger scales

In thick samples imaged by SPIM, muscle, cells, tesserae and perichondrium all emitted distinct AF signals, which differed in their excitation and emission wavelengths. The strongest excitation/emission pairings for each tissue are listed in Table 2. The perichondrium was the tissue with the strongest AF (Fluorophore 2; Fig. 8A), and needed to be saturated in image post-processing to allow other features to be visualized.

Tesserae were clearly visible beneath the perichondrium (Fig. 8A–D), exhibiting three spatially distinct regions, similar to those described above for PMMA samples, with distinctive AF wavelengths and intensities. The peripheral region of tesserae showed AF in the 482/25 nm filter window when excited with the 405 nm laser (Fluorophore 1, red in Fig. 8C). This region was encapsulated by a thin outer border delineating the edge of tesserae (~15–30 μm), which emitted a very localized AF signal in the 520/50 window (Fluorophore 2, green in Fig. 8C), when illuminated with a 488 nm excitation laser. The center region of tesserae emitted a signal with the same intensity and wavelength as the outer edge, forming a small, hazy autofluorescent region approximately the same cross-sectional shape as the tessera (in vertical and planar sections).

The unmineralized matrix emitted AF in the 593LP window when excited with the 561 nm laser (Fluorophore 3—dark blue in Fig. 8C), with low signal intensity in comparison with the cells and tesserae. However, even with low signal strength, local intensity differences could be appreciated. In particular, individual cells and cell groups were haloed by a zone of AF of lower intensity than that of the bulk cartilaginous matrix. Cells within the cartilaginous matrix and joints exhibited a strong AF (Fluorophore 2), whereas those within tesserae exhibited no visible signal. Although the resolution was not high enough to distinguish cellular form or structures (e.g. nucleus) to any degree, it was possible to visualize the large-scale, 3D arrangements of cells within the unmineralized matrix. For example, cells located directly beneath the tesserae tended to be isolated, yet with increasing distance into the cartilaginous core, tended to be organized into groups of two and then four.
4 Discussion

All major structural components of tessellated cartilage are autofluorescent, sensitive to a broad range of excitation wavelengths (405–600 nm). In particular, these components strongly reacted when excited with UV (405 nm), emitting a diversity of tissue-specific green-yellow AF (i.e. within the 480–580 nm range). We show that it is possible to image both soft and hard tissue elements of tessellated cartilage by exciting tissue AF with confocal and SPIM microscopes, resolving tissue components at a broad range of size scales, from millimeter- down to subcellular spatial resolution (1 μm).

1. Effect of preparation methods (PMMA, paraffin and thawed) on AF

The specific fluorescence signatures we observed—although all within the excitation/emission ranges mentioned in the previous paragraph—were not consistent among the different sample preparations studied, differing in the number of fluorescence signatures detected, their spectral features, and where they were expressed. The reasons for these differences are unclear, but variation in autofluorescence is known to be linked to multiple aspects of sample preparation, as well as tissue composition. For example, the loss of AF in thawed samples could be related to the destructive influence of freezing and thawing on cartilage, which causes disruption of chondrocytes [37, 38] and proteolytic degradation of fibers (e.g. collagens, actin) in the extracellular matrix [39–41]. The AF signatures evident in preliminary thawed samples of tesserae, however, are promising for future research, indicating that neither embedding media nor fluorescent markers are required for the study of this tissue. Cryosectioning of samples should facilitate AF imaging by ensuring more uniform slices resulting in more features in-plane.

With regard to embedded samples, the spectral differences between paraffin and PMMA multisets were not due to fixative effects, as both were fixed following the same protocol. Additionally, although tissue fixation with aldehydes (PFA, glutaraldehyde) can induce tissue AF due to the formation of chemical ring structures, the AF generated by PFA is generally low and hardly detectable with spectroscopy [42, 43]. Paraffin samples presenting fewer fluorophores could be related to the demineralization process. Decalcification using EDTA has been demonstrated to increase AF or produce changes in fluorophores location and/or spectra during mineral removal [44–46]. On the other hand, the capacity for AF from the mineral itself—carbonated apatite in tesserae, as well as in mammalian calcified cartilage and bone [40]—is still under debate, with several studies arguing that it is not autofluorescent [44–49]. It should be noted though that apatite AF has previously only been investigated at lower excitation-emission wavelengths than those used in this study and should be explored. Additionally, several studies have demonstrated that high temperatures can quench tissue AF and, in our study, paraffin embedding was carried out at higher temperatures (60ºC) than PMMA (4ºC). Lastly, we are confident that AF differences among preparation types were not due to the embedding media themselves: we verified that both paraffin and PMMA exhibited negligible AF (the latter supported by other studies; e.g. [50, 51]) and moreover, the embedding media were removed from sections prior to imaging.

2. Tissue biology insights from AF imaging

Our demonstration of tissue-specific AF in tessellated cartilage opens new opportunities for high-resolution optical imaging and characterization of this tissue. This is particularly true for PMMA samples, which resulted in the highest quality images of tessellated cartilage, with well-defined mineralized and non-mineralized structures (e.g. mineralization fronts, Liesegang lines, spokes, cell nuclei), many not seen in other preparation methods.
4.1 Tesserae

The mineralized tissue (tesserae) in stingray cartilage exhibited consistent and strong AF, similar to mammalian calcified cartilage, which exhibits a strong two photon fluorescence signal [52]. Tesserae AF was typically partitioned into three discrete regions: a center region, peripheral region and a narrow border at the outer margin of the tesserae. Curiously, this regional partitioning differs from that typically used to describe tesserae structure, especially in histology, where tesserae are divided into an upper “cap” and a lower “body” zone (sensu [10]).

In contrast, the three-zone AF we observed in tesserae shows no correspondence with any published histological work that we know of, although some images suggest the center can in fact stain differently than both cap and body (e.g. Figure 9b in [8]). AF images of tesserae from another stingray species (Urobatis; [11]), however, suggest a similar three-zone pattern. In studies of shark species where calcium-binding dyes had been injected to study skeletal growth [53, 54], fluorescence was also observed in central and outer border regions of tesserae, but attributed to areas of active mineralization and/or material turnover, with AF never considered [11]. For such skeletal age and growth work, our study underlines the importance of characterizing tissue AF to avoid erroneous conclusions, particularly in tissues where growth mechanisms are poorly known. The observations supporting the bipartite (histology-based) and tripartite (AF-based)
schemes of tesserae organization argue that both represent real—but different—aspects of tissue composition and architecture.

Whereas differences in collagen and proteoglycan type could, for example, explain cap and body histological staining variation, differences in AF signal must be due to other factors (e.g. density variation of an autofluorescent component), perhaps linked to tesserae development and mineralization (e.g. the growth of tesserae at their margins). The center region exhibits several distinctive characteristics, having comparatively high mineral density [19], association with acid-based stainings [8], and larger and more spherical cells than the remainder of the tessera [18]. We propose that these aspects and the center’s AF are also linked to a specific early developmental stage of the
skeleton, with the center’s AF reflecting the size and shape of tesserae from that ontogenetic stage. Tesserae first form in the unmineralized cartilage, some distance from the perichondrium and not yet in contact with each other [8, 55], gradually increasing in size during ontogeny by accreting mineral on their margins until they come into contact and approach the perichondrium. The ontogenetic transition from isolated tesserae to abutting, perichondrally-associated tesserae therefore correlates with new tissue associations and perhaps the incorporation of different autofluorescent components, and so could be signified by the shift in fluorescence characteristics between the central and peripheral regions.

In contrast, the distinctness of outer border AF relative to that of the periphery could be related to the tesseral edge being an active site of mineralization [9, 25, 26], since mineralization fronts in osteonal bone also show defined autofluorescent bands [47]. In this regard, however, the multiple fluorophores shared between the center and outer border are more difficult to reconcile, as the center is believed to be a mineralization hub only earlier in skeletal development (see above). In the case of the center region, perhaps the AF signal is related to tissue age and degree of mineralization, as in bone [56]. The tissue properties and components at the root of the AF differences among the center, peripheral and outer border regions require characterization; however, their autofluorescent characteristics and the accretionary growth of tesserae argue these differences are linked to the regions having specific developmental roles and/or representing important turning points in the mineralization of the skeleton.

The potential for AF imaging to inform our understanding of mineralization processes is further supported by the observation of mineralized spheroids in several regions of the tesserae in PMMA samples. These “calcospherites” have been observed by multiple authors at mineralization fronts in elasmobranch skeletons [10, 26, 57–59], but also in mammalian calcified cartilage and bone [60–63], where they are believed to act as accretive building blocks for mineralized tissue. In tessellated cartilage, such calcospherites have been demonstrated predominantly at interfaces between unmineralized cartilage and mineralized tissue, particularly those associated with cells: forming enclosures around chondrocytes in incipient tesserae in young animals or along the chondral border of established tesserae [9, 10, 25, 64]. In our study, AF images highlighted calcospherites in similar locations, but also lining the walls of lacunae in tesserae and additionally at the border between the center and peripheral regions, deeply embedded within the mineralized matrix of tesserae. This visible record of a past mineralization front supports our hypothesis of the autofluorescent center region as a snapshot of a specific developmental period in tesserae, while underlining AF as a useful tool to locate both active and former globular mineralization fronts.

### 4.2 Extracellular matrices

Cartilages are composed of a combination of several organic components (e.g. different collagen types, elastin, glycosaminoglycans), where their variation determines the mechanical properties and function of the tissue [2, 65]. In contrast to mammalian cartilage, the composition of elasmobranch cartilage is not well characterized, with only a few major components identified, such as the primary structural collagens (e.g. Coll1, Coll2). This lack of information is mainly related

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Table 1: Summary of image multisets analyzed, MCR-ALS model parameters, and primary results (number of unmixed fluorescence signatures resolved)

| Type of image | Embedding medium | Nr. of images | Nr. of unmixed fluorescence signatures | LOF (%) | Explained variance (%) |
|---------------|------------------|--------------|--------------------------------------|----------|------------------------|
| 3D Paraffin   | 10 (5/2/3)       | 1            | 13.85                                 | 98.08    |
| 3D PMMA      | 9 (3/3/3)        | 2            | 13.62                                 | 98.14    |
| 3D Thawed    | 1 (1/0/0)        | 2            | 8.68                                  | 99.25    |
| 4D Paraffin   | 6 (0/3/3)        | 3            | 26.31                                 | 93.08    |
| 4D PMMA      | 11 (3/3/5)       | 3            | 25.39                                 | 93.55    |
| 4D Thawed    | 3 (1/0/0)        | 2            | 24.25                                 | 94.12    |

Multisets are organized by image type (3D = 405 nm, 4D = WLL) and sample preparation methodology (paraffin, PMMA or thawed samples), with the number of images comprising each multiset listed and the number of images per ROI in each multiset detailed in parentheses immediately below (in the order: Tesserae/ Joint/ UC). The number of unmixed fluorescence signatures identified by SVD for each multiset and estimates of model quality parameters (LOF and explained variance) are also listed. No images could be analyzed for the tesserae ROI in the paraffin 4D multiset, since the image settings were not comparable with the other multisets. Tesserae information, however, was available and analyzed from joint ROI images in the paraffin 4D multiset (see Fig. 9). The most relevant ROIs analyzed can be found in Figs. 9, 10, 11; Supp. Figure 3—4
to the difficulty in identifying tissue components unequivocally, as their identification requires protocols that are either
time-consuming/costly (e.g. development of elasmobranch-specific antibodies) or involve steps that can even hamper
cOMPONENT identification or ultrastructural investigation (e.g. degradation of surrounding matrices). In comparing our
AF data to existing information on tessellated cartilage structure and composition and our own data from other imaging
modalities (histology, BSEM), we propose likely sources for some observed fluorophores, particularly those associated
with tissues of known collagen composition.

Coll1 and Coll2 are well-known fluorophores in skeletal tissue [2]. In the elasmobranch skeleton, Coll1 is known to
be present in both muscles and perichondrium [8]. Among the diverse fluorophores we observed associated with both
the perichondrium and muscle (Table 2), the one with the highest intensity is likely associated with Coll1, having also a
similar spectral profile of Coll1 in other taxa [4, 63–65] (Fig. 12A). Surprisingly, this fluorophore was also associated with
chondrocytes of the unmineralized matrix, which are not known to express Coll1 [66, 67]

Additionally, based on correspondence with immunostained samples, that the two fluorescent signals
associated with the extracellular matrix (ECM) of joints and uncalcified cartilage represent or are linked to Coll2 (Fig. 12B).
This collagen, in addition to being autofluorescent [41, 68], is indeed a primary component of the ECM of joints, unmin-
eralized cartilage and tesserae [8, 66, 69] (see also Supp. Figure 5). The specific distribution of these putative “Coll2 fluo-
rophores” raises several interesting points relevant to growth and patterning of tessellated cartilage. First, the relatively
homogeneous signal of the two fluorophores, seen throughout the ECM of joints, uncalcified cartilage and tesserae,
supports the hypothesis that tesserae arise in and are predominantly composed of Coll2, despite some local variation of
other materials (e.g. the mineral in tesserae or the Coll1 Sharpey’s fibers that insert into the cap from the perichondrium).
Additionally, the localization of these two fluorophores in the tesserae indicates a potential advantage of AF imaging
of tesserae over immunohistochemistry, where Coll2 antibody reactivity in tesserae was only possible after stringent
digestion techniques [8]. Lastly, one of the “Coll2 fluorophores” presented an extremely high signal in the laminae of

| Sample preparation | Image type | Fluorescence signals | $\lambda_{ex}$ (nm) | $\lambda_{em}$ (nm) | Distribution |
|--------------------|------------|----------------------|-------------------|------------------|-------------|
|                    |            |                      | range             | (peak)           | Perichondrium |
| 3D                 | Fluo. 1    | 405                  | -                 | 425 - 700        | Tesserae: central region + outer edge |
|                    |            |                      |                   | 480 - 500        | Joints: cells |
|                    |            |                      |                   |                  | CM: cells    |
| Paraffin           | Fluo. 2    | 470 - 570            | (470)             | 475 - 650        | Joints: matrix + cells |
|                    |            |                      |                   | 525 – 575        | CM: matrix + cells |
|                    | Fluo. 3    | 480 - 570            | (530 – 560)       | 500 - 620        | Tesserae: spokes |
|                    |            |                      |                   | 540 – 575        | CM: matrix    |
|                    | Fluo. 4    | 470 - 670            | (470 – 500)       | 490 - 700        | Tesserae: peripheral region + central border |
|                    |            |                      |                   | 500 – 590        | CM: cells     |
Table 2 (continued)

| Fluo. 1 | 405 | - | 421 - 680 (480 - 500) |
|---------|-----|---|-----------------------|
| **4D** |     |   |                       |
| Fluo. 4 | 470 - 600 (470 - 540) | 490 - 670 (540 - 580) |
|         |     |   |                       |
| Fluo. 5 | 470 - 570 (470 - 480) | 490 - 640 (510 - 560) |

**PMMA**

**3D**

| Fluo. 2 | 405 | - | 421 - 700 (495 - 500) |
|---------|-----|---|-----------------------|

**Perichondrium**

**Tesserae:** center region + outer edge

**Joint:** cells

**CM:** cells

**Muscle**

**Perichondrium**

**Tesserae:** matrix + spokes

**Joint:** matrix

**CM:** matrix

**Muscle**

**Perichondrium**

**Tesserae:** spokes + cells

**CM:** matrix + cells + PCM

**Muscle**

**Perichondrium**

**Tesserae:** center border + outer edge

**Joint:** cells

**CM:** cells

**Muscle**

**Perichondrium**

**Tesserae:** center + center border + outer edge

**CM:** matrix
spokes, suggesting the composition of these distinct features, which have otherwise proved challenging to characterize \[8, 19\]. The factors driving the formation and periodicity of spokes are unknown, but their strong AF signal suggests spoke laminae are formed by packing Coll2 fibrils in particularly high density.

Given the limited data on tessellated cartilage tissue composition, the remaining fluorescence signatures are more difficult to assign to specific tissue components (Fig. 12C–F). In vertebrates, skeletal tissues contain a variety of fluorescent structural proteins and so, additional collagens or proteins like elastin may also be present in tessellated cartilage. Our results, therefore, offer guides for future targeted characterization of tessellated cartilage structures, whose tissue compositions have yet to be determined.
4.3 Cells

The AF signal provided information on cellular morphology, for example, demonstrating a clear pericellular matrix (PCM) region, a narrow tissue zone haloing chondrocytes in the unmineralized cartilage. In mammalian cartilage, the PCM acts as an interface between the cells and their environment and is crucial in determining how chondrocytes interact both mechanically and physicochemically with the surrounding matrix [70–72]. A factor in mediating these behaviors is the distinct composition of the PCM which, for example, has a higher proteoglycan content than the surrounding cartilage and is composed uniquely of collagen type VI [73, 74]. Indeed, the proteoglycan aggrecan is considered a potential source of pericellular AF in two-photon fluorescence microscopy (TPF) [75, 76]. Despite its importance in mammalian cartilage, a defined PCM has never been described in elasmobranchs, neither with histology nor electron microscopy. Although we cannot yet identify the source of PCM AF, its defined signal in elasmobranch cartilage suggests it may have distinct compositional, architectural and/or mechanical characteristics, as in mammalian cartilage.
Despite fluorophores having different tissue associations, nearly all fluorophores observed were associated with chondrocytes. Cell AF is typically due to NADH, FAD+ (flavins) and lipofuscins (lipopigments), cytoplasmic components with distinct spectral profiles [22, 77–79]. Among these components, flavins are the only ones with a spectral signature located within the spectral ranges in this study ($\lambda_{ex} = 440–470$ nm / $\lambda_{em} = 480–540$ nm; [22]), whereas the other autofluorescent cellular components are associated with lower excitation wavelengths (< 405 nm). Although the source(s) of cellular AF cannot be identified in this study, we note that chondrocytes emitted different signals depending on their location in the tissue. This may be indicative of cells’ production of different components and, ultimately, variation in cell function. This hypothesis is supported by the cells in the upper part of the joint emitting a brighter AF signal than other cells, reinforcing the recent findings of Marconi et al. [67], where joint cells exhibited higher production of Coll2.

5 Conclusions

Tessellated cartilage contains a diversity of fluorophores that emit a native AF signal which, particularly after fixation, embedding and sectioning, is strong enough to image the tissue at a level of detail rivaling backscatter electron microscopy and histology. The AF signal was affected by sample preparation, which should be more deeply explored to clarify how tissue architectures and composition are distorted through common investigation techniques. Images of samples prepared for PMMA showed exceptional preservation of tissue morphology, especially mineralized parts, even allowing identification of active and former mineralization fronts through the localization of calcospherites. AF in tessellated cartilage is mainly emitted in the green–yellow region of the spectrum (480–580 nm), excited with a broad range of wavelengths (405–600 nm), but reacting most strongly to 405 nm excitation (UV). Several fluorophores could be associated with Coll1 and Coll2, the main structural collagens of tessellated cartilage; the diverse and specific distributions of other unidentified fluorophores, however, suggest the presence of additional collagens and/or other autofluorescent molecules yet to be characterized in elasmobranch cartilage. The localization of the varied fluorescent signatures allowed description of several tissue morphologies, including a novel organizational scheme of three concentric autofluorescent regions in tesserae and a pericellular matrix surrounding chondrocytes. Tissue AF is therefore demonstrated to be a powerful diagnostic tool to image tessellated cartilage that, with further studies and standardized methods, will allow us to generate composition maps without the need of more demanding preparation techniques.

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Authors’ contributions JC was responsible for experiment design, sample preparation, data collection and analysis, made the majority of manuscript figures, and wrote the manuscript. MM was involved in experiment design, sample preparation and data collection with the confocal microscope and wrote for the corresponding sections of the manuscript. AGS and AJ were involved in autofluorescence signal analysis with MCR-ALS and wrote and generated figures for the corresponding manuscript sections. MB was involved in experiment design, was responsible for the preparation of paraffin samples and histological stainings and wrote the corresponding manuscript sections. EJG collected the light-sheet microscopy (SPIM) data and wrote for the corresponding sections of the manuscript. PL-A was involved in experiment design and provided access to the Super Resolution Light Microscopy facilities at ICFO. MND supervised the project, was involved in experiment design, participated in data collection and analysis, and wrote the manuscript. All authors edited the manuscript and approved its submission. All authors read and approved the final manuscript.

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Data availability and code availability https://doi.org/10.17617/3.4h
Declarations

Ethics approval and consent to participate This study did not involve any experiment on living animals. All experiments were obtained from fisheries by-catch and therefore did not require the approval of an ethical committee.

Consent to publish This study did not involve any experiment with patients and, therefore, did not require any consent to publish.

Competing interests The authors declare no competing interests.

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