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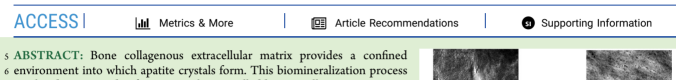
Article

¹ Collagen Suprafibrillar Confinement Drives the Activity of Acidic ² Calcium-Binding Polymers on Apatite Mineralization

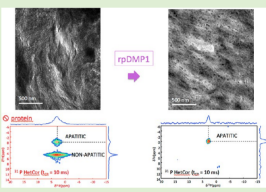
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7 is related to a cascade of events partly controlled by noncollagenous proteins. 8 Although overlooked in bone models, concentration and physical environ-9 ment influence their activities. Here, we show that collagen suprafibrillar 10 confinement in bone comprising intra- and interfibrillar spaces drives the 11 activity of biomimetic acidic calcium-binding polymers on apatite 12 mineralization. The difference in mineralization between an entrapping 13 dentin matrix protein-1 (DMP1) recombinant peptide (rpDMP1) and the 14 synthetic polyaspartate validates the specificity of the 57-KD fragment of 15 DMP1 in the regulation of mineralization, but strikingly without 16 phosphorylation. We show that all the identified functions of rpDMP1 are 17 dedicated to preclude pathological mineralization. Interestingly, transient



18 apatite phases are only found using a high nonphysiological concentration of additives. The possibility to combine biomimetic 19 concentration of both collagen and additives ensures specific chemical interactions and offers perspectives for understanding the role 20 of bone components in mineralization.

21 INTRODUCTION

22 Bone is a complex composite material, which closely associates 23 cells and an extracellular matrix (ECM). The bone ECM 24 combines various components, that is, collagen, noncollage-25 nous proteins (NCPs), carbonated hydroxyapatite (HA) 26 nanoparticles, and water. Although bone is one of the most 27 studied living materials, the exact role of its components, 28 noticeably the organic one, is still debated. Both intra- $^{1-3}$ (~40 29 nm) and interfibrillar⁴ (~1.5 nm) spaces in collagen are 30 proposed to be nucleation sites for bone mineral. The 31 interfibrillar confinement generated by the collagen matrix 32 increases as the ECM becomes mineralized, thus forming the 33 mature bone (Figure 1A) with the characteristic twisted 34 plywood pattern⁵ (Figure 1B). This suprafibrillar three-35 dimensional (3D) architecture affects the physical properties 36 of the ECM such as mechanical response^{6,7} and microenviron-37 ment (e.g., size and spatial distribution of apatite, local 38 hydrated environment of phosphate ions).⁸ In addition, *in vitro* 39 investigations on NCPs activity have indeed yielded contra-40 dictory findings when the protein is studied either in solution 41 or in gel.^{9,10} Upon working with gels, loose collagen matrices 42 imply large interfibrillar spaces due to the lack of collagen ⁴³ packing,⁸ meaning that bone is only reproduced at the fibril ⁴⁴ level.^{11,12} Consequently, aside *in vivo*-based experiments, 45 biomimetic models in terms of confinement are needed to 46 ensure that investigations of NCPs activity are conducted

under native environment. As demonstration, only models ⁴⁷ mimicking both collagen density and geometry described for ⁴⁸ mature bone (250 mg/mL)⁸ tissues led to the typical apatite/ ⁴⁹ collagen co-alignment (Figure 1C), but noticeably with higher ⁵⁰ concentrations of calcium ions (Figure 1D, left) than that ⁵¹ described in extracellular fluid (ECF).¹³ Indeed, simulated ⁵² body fluid (SBF) failed to produce intrafibrillar mineral; ⁵³ instead, apatite in the form of spherulite was found (Figure 1D, ⁵⁴ right). This result suggested that Ca-rich proteins from bone ⁵⁵ ECM concentrate ions in the gap regions.

To go further on the effect of bone ECM confinement on 57 NCPs activity, we pursue the study by entrapping dentin 58 matrix protein-1 (DMP1), an acidic calcium-binding poly- 59 mer¹⁴ within our models. DMP1 is a phosphorylated ECM 60 protein produced by osteocytes and odontoblasts that is 61 commonly accepted as mediator in bone mineralization 62 processes.^{15–17} Indeed, DMP1 knockout mice display defective 63 mineralization of dentin^{18–21} and bone^{22,23} resembling 64 phenotypes observed in human genetics. Interestingly, the 65

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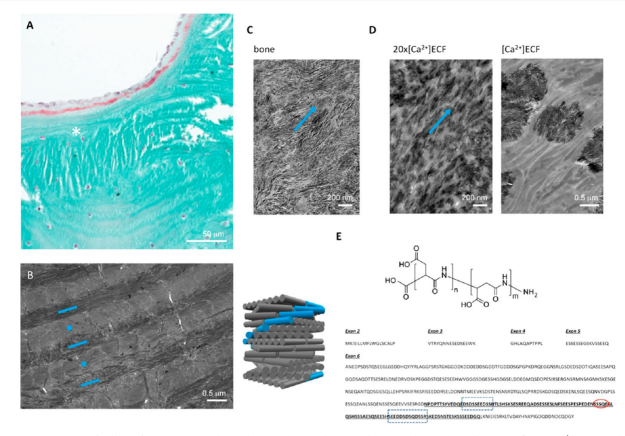


Figure 1. Comparison of collagen/apatite relationship in bone and in synthetic models reproducing the collagen confinement (density and order) found in bone. (A) Thin section of bone stained with Masson's trichrome. From top to bottom: Osteoblasts, osteoid tissue (light blue and red), and mature bone (*). Transmission electron microscopy ultrathin sections of (B) mature decalcified human bone that is characterized by a twisted plywood organization of collagen fibrils (blue dot and bars represent molecules that are perpendicular and parallel to the observation plan, respectively) schematically represented, (C) unstained sheep bone where the alignment of apatite platelets along the fibril is observed (the blue arrow shows the orientation), (D) Coll/CHA/SBF (left) and Coll/SBF/SBF (right) where $[Ca^{2+}] = 20[Ca^{2+}]ECF$ and $[Ca^{2+}] = [Ca^{2+}]ECF$, respectively. The collagen/apatite co-alignment similar to that in bone is observed in Coll/CHA/SBF (left) while spherulitic apatite crystals are obtained in Coll/SBF/SBF (right). In (E), the acidic calcium-polymer used in the biomimetic model, that is, the polyaspartate formula and the human DMP1 sequence (in bold: target peptide; blue boxed texts: collagen binding sites; red encircled text: motif found as unchanged during 220 My). The influence of organic polymers on apatite formation is studied at fixed physiological concentration of calcium, that is, $[Ca^{2+}] = [Ca^{2+}]ECF$ with two different concentrations for rpDMP1 as follows: 2.5 μ g/mL (1rpDMP1) and 25 μ g/mL (10rpDMP1).

66 role of DMP1 has been investigated *in vitro* through different 67 models (e.g., in solution,^{24,25} adsorbed onto glass plates,^{14,26} in 68 gelatin-gel systems,^{15,27} on a transmission electron microscopy 69 (TEM) grid coated with recombinant spider silks,²⁸ with 70 preassembled collagen fibrils, 29,30 and with early calcifying 71 bone-like matrix),³¹ reaching the conclusion that the protein 72 could be involved at different levels including in apatite ⁷³ nucleation control^{27,32} and growth inhibition,³³ in mediating 74 the size of collagen fibrils,²⁹ and in stabilizing amorphous 75 calcium phosphate (ACP) phase.^{33,34} To clarify such func-76 tional diversity, a DMP1 recombinant peptide (rpDMP1) 77 containing those two collagen binding sites²⁹ and the peptide ⁴²⁷SSSQE⁴³¹ was synthesized and used here (unphosphory-78 79 lated, 121 amino acids, 13 kDa, pI = 4.14) (Figure 1E, down). 80 Indeed, four DMP1 peptides are identified in dentin and bone $_{81}$ extracts: (1) the full-length protein, (2) a N-ter fragment of 37 82 kDa, (3) a C-ter fragment of 57 kDa,³⁵ and (4) a proteoglycan 83 derived from the N-ter fragment and known as DMP1-PG 84 found in rat and mouse.³⁶ Importantly, five highly conserved 85 motifs during mammalian evolution were identified including 86 427SSSQE⁴³¹, suggesting the importance of this acidic peptide 87 for the protein structure and/or function.³⁷ Moreover, it was 88 reported that the 57 kDa fragment recapitulates the function of

full-length DMP1 in regulation of mineralization and osteocyte 89 maturation.³⁸ Finally, phosphorylation appears to be dedicated 90 to the organization of mineral when there is no collagen 91 fibrillar packing³⁹ since both phosphorylated and non-92 phosphorylated rDMP1 are proposed as apatite nucleators.⁴⁰ 93

In addition to the confinement criterion, the concentration 94 of protein closer to physiological conditions^{41,42} is of 95 importance to access to their native properties. Noticeably, 96 differences in range of concentrations (factor 10 to 100) can 97 lead to opposite activities^{14,15,24,26,29} blurring the conclusions. 98

To validate the specificity of the 57-KD fragment of DMP1, 99 experiments were also carried out with a synthetic calcium- 100 binding polymer, that is, the biomimetic poly-L-aspartic acid 101 (polyAsp, 1.2 kDa, pI = 2.98) (Figure 1E, up).^{43,44} Nowadays, 102 polyAsp is commonly used in biomineralization models for 103 mimicking acidic proteins in biological calcified tissues (nacre, 104 bone, dentin). This polymer is described to allow the 105 intrafibrillar infiltration of both DMP1⁴⁵ and apatite ion 106 precursors⁴⁶ and consequently the collagen/mineral co-align- 107 ment.⁴⁷

Here, we show that collagen suprafibrillar confinement 109 drives the activity of acidic calcium-binding polymers on apatite 110 mineralization. The difference in mineralization between 111

112 rpDMP1 and polyAsp illustrates the specificity of NCPs amino 113 acids sequence. Under confinement (i.e., intra- and interfi-114 brillar spaces), we show that all the identified functions of 115 rpDMP1 are dedicated for proper calcification to occur, 116 namely the collagen/apatite co-alignment.48,49 Indeed, we 117 show that while collagen nucleates apatite, rpDMP1 (i) 118 concentrates the apatite ion precursors locally, interfering as 119 a supporting agent for collagen (ii) to induce the formation of 120 a first highly crystalline apatite crystal. In addition, we discard 121 that rpDMP1 stabilizes the possible transient precursors of 122 bioapatite^{50,51} since they only form at high nonphysiological 123 concentrations of protein. Finally, (iii) we confirm that it 124 inhibits the apatite growth, but (iv) also show that it might 125 inhibit the homogeneous nucleation irrespective of confine-126 ment. Overall, the work demonstrates that aside from cellular 127 and biochemical processes, physicochemical parameters take 128 part in the control of bone biomineralization.

129 MATERIALS AND METHODS

Synthesis and Purification of the Recombinant DMP1 130 131 Peptide. Cloning. The coding sequence of our targeted human 132 DMP1 peptide (i.e., amino acids 367-481) was isolated from a 133 human primary osteoblast cDNA (Promocell). It was amplified in a 134 thermal cycler (Mastercycler pro, Eppendorf) by PCR using the 135 oligonucleotides 5' ATGC CATATG AACCCCGACCCCA 3' and 5' 136 GCAT CTCGAGTCA GTG GTG GTG GTG GTG GTG GTG 137 CAACTGGCCATCTTC 3' to create NdeI and XhoI restriction sites 138 at the start and stop codon, respectively. Cycling conditions were 139 initial denaturation at 94 °C for 2 min, followed by 35 cycles, each 140 cycle consisting of 30 s of denaturation at 94 °C, 45 s of annealing at 141 60 °C and 45 s of elongation at 72 °C. The final elongation lasted for 142 2 min at 72 °C. PCR products were analyzed by 1% agarose gel 143 electrophoresis and observed in an analyzer Gel Doc (BIORAD, 144 France) after ethidium bromide staining. The amplified fragment was 145 purified using the QIAquick PCR purification kit (Qiagen SA) and 146 digested with NdeI and XhoI restriction enzymes. The resulting 147 products were separated by a migration in a 1.5% agarose gel with 148 ethidium bromide and cloned into the pET22b vector (Novagen/ 149 VWR International S.A.S) which had been beforehand digested 150 similarly. Competent Novablue cells were transformed with the 151 ligation mix and selected by overnight growth on LB agarose plates 152 containing 70 μ g/mL ampicillin. Positive clones were checked for the presence of the 0.8 kb fragment by NdeI and XhoI restriction analysis. 153 Expression. BL21 (DE3) cells transformed with pET-DMP1 were 154 155 grown on a rotating table (220 rpm) overnight at 37 °C in LB 156 medium with 50 μ M ampicilin. When the optical density at 600 nm of 157 the bacterial broth reached 0.6-0.8, the induction was realized with 158 an addition of 1 mM of isopropyl β -D-1-thiogalactopyranoside. After 3 159 h, the cells were centrifuged at 5000g for 10 min, and the pellets were 160 frozen at -20 °C until purification.

161 *Purification*. Pellets were resuspended in 5 mL of buffer A (PBS 162 1×, 50 mM imidazole, complete inhibitor mix (Roche), lysozyme, 163 DNase) and lyzed by sonication three times. The bacterial extract was 164 loaded on a 5 mL bed-volume HisTrap column using an Akta Purifier 165 10 (GE-Healthcare). Buffer A was flowed through the column at 1 166 mL/min until A_{280} of the flow through reached a stable value. A 60 167 mL linear gradient of 0–100% Buffer B (PBS 1×, 500 mM imidazole, 168 complete inhibitor mix (Roche), lysozyme, DNase) in buffer A was 169 applied, and 1 mL fractions were collected. Aliquots were analyzed by 170 10% SDS-PAGE and the gels stained with Coomassie blue to 171 determine the quality of purified rpDMP1 (12 kDa, pI = 4.14) 172 (Figure S6).

¹⁷³ Sample Preparation. Collagen Extraction. A solution of type I ¹⁷⁴ collagen at ~3 mg/mL in 0.5 M acetic acid was prepared as previously ¹⁷⁵ described.⁵² The collagen was extracted from rat tail tendons. After a ¹⁷⁶ washing step with PBS, tendons were solubilized in 0.5 M acetic acid, ¹⁷⁷ and the solution was clarified by centrifugation (21,000 rpm, 2 h, 11 ¹⁷⁸ °C). The supernatant was selectively precipitated with 0.3 and 0.6 M of NaCl by two centrifugations (21,000 rpm, 3 h, 11 °C then 4400 179 rpm, 45 min, 11 °C), in order to remove proteins other than type I 180 collagen and collagen, respectively. The pellets were solubilized in 0.5 181 M acetic acid and dialyzed against 0.1 M acetic acid in order to 182 remove salts from the solution. A final centrifugation was performed 183 (21,000 rpm, 4 h, 11 °C), and the concentration was adjusted to a 184 final stock concentration of ~3 mg/mL. The final concentration of 185 type I collagen solution was estimated by hydroxyproline titration.⁵³

Synthesis of Collagen/Apatite Matrices. The matrices were 187 prepared according to a procedure that combines injection and 188 reverse dialysis processes (patent WO2011151587A2). The mineral-189 ization conditions for collagen matrices (concentrations of compo-190 nents) are summarized in Table 1. Collagen samples were disk-shaped 191 t1

sample designation	end collagen concentration (mg/mL)	acidic polymer (µg/ mL)	HA ion precursors in dialysis solution (acetic acid)	additional SBF bath
Coll	250	-	-	-
Coll/SBF	250	-	SBF	-
Coll/SBF/SBF	250	-	SBF	+
Coll/CHA/SBF	250	-	CHA	+
Coll/SBF(rpDMP1)	250	2.5	SBF	-
Coll/ SBF(rpDMP1)/ SBF	250	2.5	SBF	+
Coll/10rpDMP1	250	25	-	-
Coll/ SBF(10rpDMP1)	250	25	SBF	-
Coll/ SBF(10rpDMP1)/ SBF	250	25	SBF	+
Coll/SBF(polyAsp)	250	5.75	SBF	-
Coll/SBF(po- lyAsp)/SBF	250	5.75	SBF	+
CollOsteoid/SBF/ SBF	40	-	SBF	+
CollOsteoid/ SBF(10rpDMP1)/ SBF	40	25	SBF	+

^{*a*}Ionic composition of SBF and CHA solutions are detailed in Table S1.

with a thickness of ~1 mm and a diameter of ~10 mm. A soluble 192 acidic collagen solution (1 mg/mL, 0.5 M acetic acid) supplemented 193 by a $1 \times$ SBF solution mimicking the ionic compounds found in the 194 human plasma was prepared by diluting the stock solution (3 mg/mL, 195 0.5 M acetic acid) with a 1.5× SBF acidic solution (0.5 M acetic acid). 196 SBF was prepared as previously described.⁵⁴ The concentrations of 197 the salts precursors are summarized in Table S1. Two concentrations 198 of rpDMP1 were added to this solution: 2.5 μ g/mL (low) and 25 μ g/ 199 mL (high) to form the matrices referred as Coll/SBF-rpDMP1and 200 Coll/SBF-10rpDMP1, respectively. The rpDMP1 control matrix with 201 the highest concentration of rpDMP1 and without mineral (Coll/ 202 10rpDMP1) was obtained by diluting the stock solution with acetic 203 acid (0.5 M). Two matrices used as control were prepared: (i) 204 without any organo-mineral additives (Coll) and (ii) with apatite ion 205 precursors but without addition of any acidic polymer (Coll/SBF). 206 The matrices supplemented by polyaspartate (Coll/SBF-polyAsp) 207 were obtained in a similar way, by adding this acidic polymer (5.75 208 μ g/mL, Lanxess-Bayer, BaypureDS100, 1200 g/mol, pI = 2.98) to the 209 acidic collagen solution. To mimic the osteoid tissue⁵⁵ CollOsteoid/ 210 SBF/SBF, 40 mg/mL collagen matrices were also formed. In this case, 211 $25 \,\mu g/mL rpDMP1$ was added to the acidic collagen solution forming 212 the matrix referred as CollOsteoid/SBF-10rpDMP1/SBF. All these 213 solutions were continually injected in a closed dialysis chamber for 1 214 week. The bottom of the chamber contained a dialysis membrane 215 with a molecular weight cut off of 1 kDa. The reverse dialysis 216 process⁵⁶ was set against polyethylene glycol (PEG, 35 kDa, Fluka) to 217

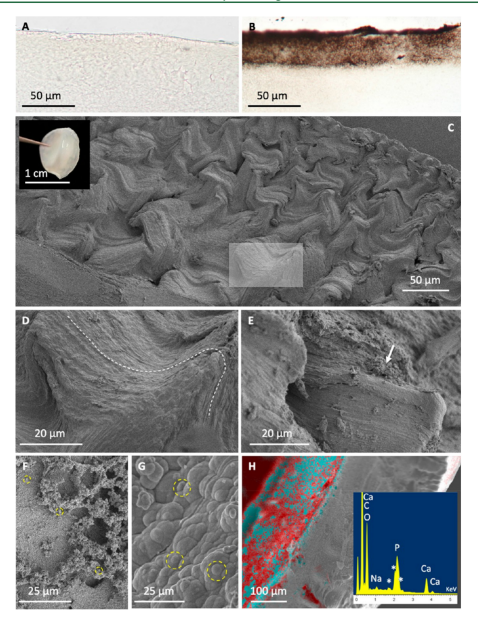


Figure 2. Structural characterizations of the hybrid collagen/apatite matrices in the presence of rpDMP1 containing the highly conserved motif. Histological sections of Coll/SBF(1rpDMP1) (A) and Coll/SBF(1rpDMP1)/SBF (B) stained with von Kossa. SEM observations of the dense collagen matrix (250 mg/mL) at low rpDMP1 content (C–F). Examination of the fractured interior of the disk-shaped matrix (C, inset) before SBF bath (Coll/SBF(1rpDMP1) (C, D; rectangle in C indicates the enlarged section that is shown in D). Examination after SBF bath (Coll/SBF(1rpDMP1)/SBF): interior (E) and surface (F). In (E), the roughness appears to increase after SBF (arrow). (G) Surface of the collagen matrix precipitated without rpDMP1 (Coll/SBF/SBF). The size and the amount of spherulitic crystals (yellow dashed circles) at the surface of the matrices decrease with rpDMP1. (H) Analysis of Coll/SBF(1rpDMP1)/SBF by EDX coupled with SEM; calcium ions are in red, and phosphorus ions in blue (SEM gold coating*).

218 control the final concentration of collagen. The PEG was dissolved in 219 0.5 M acetic acid up to ~300 mg/mL for all Coll matrices or ~50 220 mg/mL for all CollOsteoid matrices. To form the mineralized 221 matrices, the ionic precursors of SBF or CHA were dissolved in the 222 PEG/acetic acid solution. After injection of the total amount of 223 collagen, dialysis was continued for 1 week in order to obtain a 224 homogeneous concentration in the samples and a good maturation of 225 the peptides. The pH was then increased to a range of 9–10 by 226 ammonia gas diffusion for 3 days to induce collagen fibrillogenesis, 227 stabilize the liquid crystalline organization into dense fibrillar 228 matrices, and precipitate the mineral phase. These matrices were 229 then removed from the dialysis chamber and washed several times in 230 sterile double-distilled water until complete neutralization. The final 231 concentration of type I collagen in the different collagen matrices was 232 assessed by hydroxyproline titration and found to be ~250 mg/mL for all Coll matrices and ~40 mg/mL for all CollOsteoid matrices. 233 The different matrices were split into halves, and the mineralization 234 degree of one-half was increased by a bath in 45 mL of 1.5× SBF 235 solution at pH = 7.4 under mild rotary stirring (220 rpm) at 37 °C for 236 1 week.⁸ 237

Mineral Characterization. *Histology*. Bone samples were fixed in 238 4% paraformaldehyde and embedded in paraffin for sectioning. Ten 239 μ m-thick serial sections perpendicular to the cell layer were dewaxed, 240 rehydrated, and stained specifically by von Kossa, thus identifying 241 divalent ions. The slides were rinsed, dehydrated, and mounted for 242 observation with an optical microscope (Nikon E600 POL) or an 243 epifluorescence microscope (AXIO 100 Zeiss). 244

Scanning Electron Microscopy and Energy-Dispersive X-ray 245 Spectroscopy. Each sample was fixed in 3.6% glutaraldehyde in a 246 cacodylate/saccharose buffer solution (0.05 M/0.6 M -pH 7.4). The 247

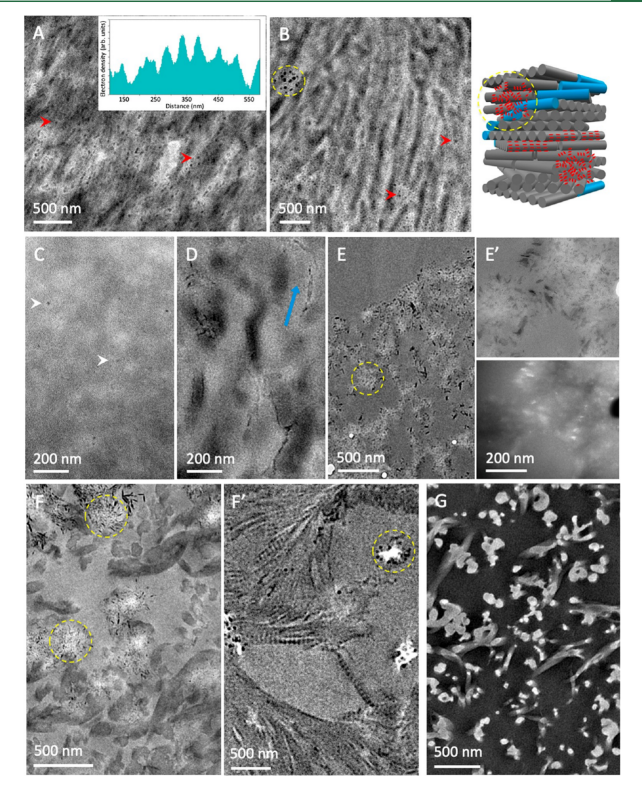


Figure 3. Investigations by TEM of calcium phosphate precipitation in rpDMP1 containing a mature bone-like matrix. Observations of unstained ultrathin sections of (A) Coll/SBF(1rpDMP1) and (B) Coll/SBF(10rpDMP1) before the SBF bath. Both matrices present nanometric electrondense precipitates (red arrows) along and/or within the collagen fibrils with periodicity highlighted by the intensity profile. In addition, aggregates of the "dot-like" precipitates are seen in Coll/SBF(10rpDMP1); the scheme illustrates the corresponding distribution of dot-like precipitates (red) related to the organization of collagen fibrils. They are found along the fibrils or as aggregates (yellow dashed circle). (C) A control matrix prepared with the high concentration of rpDMP1 (25 μ g/mL) and without ionic apatite precursors (Coll/10rpDMP1) shows that collagen fibrils are hardly observed in contrast to the precipitated proteins (yellow arrows). After immersion in SBF, (D) axial alignment of platelets are observed inside the matrix. (E) Observations within the most superficial layer of the matrix show the presence of spherulitic crystals (E') according to bright-/dark-field images. Observations at the surface of Coll/SBF(10rpDMP1)/SBF (F) without and (F') with staining where the dashed yellow circles highlight the presence of spherulite. Observations at the surface of (G) Coll without staining. 248 samples were dehydrated through successive ethanol bath (50%, 70%, 249 80%, 90%, and 100%), and a supercritical CO_2 drying process was 250 performed on a BAL-TEC 030. Then samples were sputter-coated 251 with a gold layer of 10 nm and observed in a Hitachi S-3400N 252 operating at 3 kV.

Energy-dispersive X-ray (EDX) microanalysis was used for the mapping of mineral deposits inside the matrices. The EDX instrument X-Max (Oxford Instruments) was coupled to a scanning electron microscope Hitachi S-3400N operating at 12 kV, and the Oxford Microanalysis Group XAN.70 software was used for this analysis.

Transmission electron microscopy. This protocol is similar to the protocol for the scanning electron microscopy (SEM). Then, samples were rinsed, dehydrated, and embedded in Epon 812. For a few amples (always mentioned in the text), an additional postfixation was carried in 2% osmium tetroxide in a cacodylate/saccharose buffer solution (0.4 M/0.6 M-pH 7.4) during 1 h at 4 °C, otherwise neither osmium nor uranyl acetate (staining) were added to avoid artifacts on the ultrathin sections. Sections (~80 nm) were observed with a Tecnai protection and the section of the spirit G2 operating at 120 kV.

267 Wide-Angle X-ray Diffraction (Transmission Mode). Matrices 268 were inserted in X-ray cylindrical borosilicate capillary tubes. The 269 tubes were wax-sealed to keep the samples hydrated and placed 270 directly in the vacuum chamber beam. X-ray diffraction experiments 271 were performed with a S-MAX 3000 RIGAKU using a mono-272 chromatic CuK α radiation. The diameter of the cylindrical beam 273 dimension of the specimen was 400 μ m. The data were collected in 274 the 3–60° range (2 θ). The sample-to-detection distance was 0.059 m 275 with a diameter of capillary tubes at 1 mm or 0.058 m with a diameter 276 of 2 mm. The two-dimensional (2D) wide-angle X-ray diffraction 277 (WAXD) patterns were collected with imaging plates then scanned. 278 The data were analyzed using Image (LPS, U-psud) software.

279 ζ -Potential Measurements. The matrices were washed before 280 characterization precluding the involvement of free ions in the 281 resulting global charge. Samples were crushed in liquid nitrogen, and 282 the resulting powders were dispersed in PBS solution. Measurements 283 were carried out using a Malvern Zetasizer Nano ZS90.

Thermogravimetric Analysis. Samples were dried under a laminar hood overnight to minimize the mass loss of loosely bounded water. The analysis was performed on a thermo-microbalance instrument (NETZSCH STA 409PC). The measurement was performed from temperature to 1000 °C in an oxidizing atmosphere (air) with a heating rate of 5 °C/min.

290 Nuclear Magnetic Resonance. Solid-state nuclear magnetic 291 resonance (ssNMR) experiments were realized on hydrated samples. 292 Magic angle spinning (MAS) spectra were acquired at a frequency of 293 8 kHz, with samples packed into 4 mm zirconia rotors. ${}^{1}\text{H}-{}^{31}\text{P}$ cross-294 polarization (CP) experiments were performed on an Avance 300 295 Bruker spectrometer operating at frequencies of 300.13 MHz (${}^{1}\text{H}$) 296 and 121.50 MHz (${}^{31}\text{P}$). The contact times (CT) were set at 10 and 1 297 ms, and the recycle delay (RD) at 2 s. A 2D ${}^{1}\text{H}-{}^{31}\text{P}$ heteronuclear 298 correlation (HETCOR) was performed with the following parame-299 ters: RD = 2 s, CT = 10 and 1 ms, 1280 transients for each 128 t1 300 increments. ${}^{1}\text{H}$ and ${}^{31}\text{P}$ chemical shifts were referenced ($\delta = 0$ ppm) 301 to adamantane and to 855% aqueous H₃PO₄, respectively.

302 **RESULTS AND DISCUSSION**

Structural Characterizations of rpDMP1-Loaded Min-304 **eralized Collagen Matrices.** Experimental conditions and 305 matrices composition (see Materials and Methods) are 306 summarized in Table 1. It includes collagen matrices without 307 additives used as control, namely Coll, Coll/SBF (Figure S1), 308 Coll/CHA/SBF (Figure 1D, left), and Coll/SBF/SBF (Figure 309 1D, right).⁸ This means that after coprecipitation of collagen 310 with either CHA (~20 times more concentrated in apatite ion 311 precursors see Table S1) or SBF, these two last matrices were 312 then immersed in SBF to mimic further steps of mineral 313 growth. With additives (rpDMP1, polyAsp), fibril precipitation 314 was only performed in SBF ($[Ca^{2+}] = [Ca^{2+}]ECF)^{57}$ to set physiological-like conditions. Two concentrations of rpDMP1 315 were used here, for which we will refer to low (2.5 μ g/mL, 316 1rpDMP1) and high (25 μ g/mL, 10rpDMP1). For a better 317 reading, data for 10rpDMP1 are displayed in the main text 318 only when differences in mineral formation are observed 319 between the two concentrations. Additional data for 320 10rpDMP1 are shown in Figure S2.

We first investigated the spreading of mineral within 322 collagen matrices before and after immersion in SBF for 323 both rpDMP1 concentrations. For this purpose, we used von 324 Kossa staining on histological sections as it allows observations 325 at large scale. Interestingly, staining is hardly observed without 326 SBF extra-bath (Coll/SBF(1rpDMP1), Figure 2A), whereas a 327 f2 strong staining is observed for both matrices at the superficial 328 layer (~50 μ m thick) after immersion (Coll/SBF(1rpDMP1)/ 329 SBF, Figure 2B). Because von Kossa staining may interfere 330 with any divalent ions,⁵⁸ further characterizations were 331 performed to confirm the presence of mineral. 332

To identify the distribution of the minerals at a lower scale, 333 investigations were performed at the surface and inside 334 fractured disk-shaped matrices (Figure 2C-G) using SEM 335 coupled with EDX microanalysis (Figure 2H). Observations 336 inside the matrix reveal that the entrapment of rpDMP1 does 337 not disturb the self-assembly of collagen fibrils since twisted 338 plywood structures form over large distances (Figure 2C,D 339 versus Figure S1, Coll/SBF). After immersion in SBF (Coll/ 340 SBF(1rpDMP1)/SBF), the surface roughness is more 341 pronounced by the deposition of particles on the closely 342 packed oriented fibrils (Figure 2D versus 2E, arrow). On the 343 surface of the matrices, spherulites usually consisting of 344 aggregates crystals⁵⁹ are observed, and, interestingly, they 345 appear to be smaller in the presence of the protein (~0.5 μ m 346 versus ~4 μ m in Coll/SBF/SBF, yellow dashed circles in 347 Figure 2F,G). In addition, they appear less abundant since they 348 do not cover the whole surface. EDX analysis shows that the 349 spherulites are mainly composed of calcium and phosphorus 350 atoms with an average Ca/P ratio of about 1.66-1.71, 351 suggesting the formation of apatite in the presence of 352 rpDMP1 (Figure 2H).⁶⁰ 353

Confinement Effect on Apatite/rpDMP1 Distribution. ³⁵⁴ TEM investigations were performed on thin sections of Coll/ ³⁵⁵ SBF(1rpDMP1) and Coll/SBF(10rpDMP1) to access higher ³⁵⁶ magnifications. TEM sections were not stained to avoid the ³⁵⁷ presence of staining deposits that are difficult to distinguish ³⁵⁸ from CaP crystals.⁸ Observations evidence the presence of ³⁵⁹ nanometric electron-dense ("dot-like") precipitates (~30 nm) ³⁶⁰ with low and high rpDMP1 concentrations (red arrows in ³⁶¹ Figure 3A,B, respectively) within the dense collagen network at ³⁶² f3 this scale. The striated pattern (67 nm) is observed locally ³⁶³ (Figure 3A, inset), indicating that some minerals localize inside ³⁶⁴ the gap regions.^{8,61} Interestingly, additional aggregates of ³⁶⁵ nanoprecipitates are observed for high rpDMP1 content ³⁶⁶ (Figure 3B, yellow dashed circle). ³⁶⁷

To help identify the nature of the "dot-like" precipitates, a 368 control matrix was prepared without apatite ion precursors 369 (Coll/10rpDMP1) (Figure 3C). Likewise, "dot-like" particles 370 are observed (white arrows), confirming that the nano- 371 precipitates are also composed of the protein, in agreement 372 with previous observations.¹⁴ Although it is difficult to 373 conclude due to the low contrast of organic components in 374 TEM, the precipitates appear less abundant and not distributed 375 along the fibrils (Figure 3C versus Figure 3A,B). Nevertheless, 376 spherulites are observed without rpDMP1 in SBF (Coll/SBF/ 377

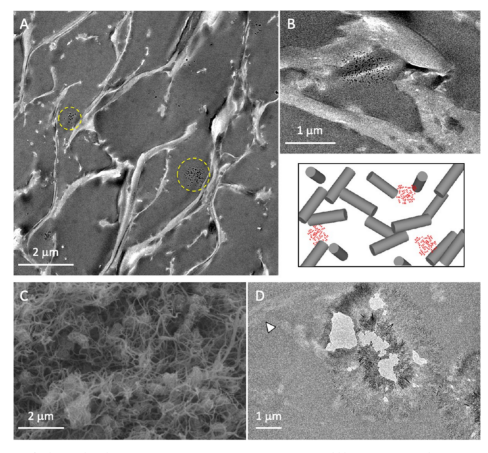


Figure 4. Investigations of calcium phosphate precipitation in rpDMP1 containing osteoid-like matrix. TEM observations of unstained ultrathin sections of CollOsteoid/SBF(10rpDMP1)/SBF (A) at low magnification where "dot-like" precipitates aggregate locally (yellow dashed circle). They are clearly seen (B) at higher magnification. Below, the corresponding relationship between mineral and collagen is represented in the scheme. (C) SEM and (D) TEM micrographs of CollOsteoid/SBF/SBF. The absence of staining makes the observation of cross-striated fibrils usually seen in a longitudinal cut (white arrow) difficult.

378 SBF in Figure 1D and Coll/SBF⁸), indicating that rpDMP1 379 may interfere with some ions to localize specifically inside the 380 gap zone.

After a SBF bath, few platelets with axial alignment (blue 381 382 arrow in Figure 3D) are observed for both rpDMP1 383 concentrations together with spherulitic particles at the surface 384 and within the most superficial layer of the matrix (dashed 385 yellow circles in Figure 3E,F). Noticeably, they appear smaller 386 in size (~300 nm versus ~2 μ m) than those found without 387 rpDMP1 (Figure 1D, right), but remain crystalline according 388 to the contrast observed in the dark-field TEM image (Figure 389 3E'). Further observations at the surface of Coll/SBF-390 (10rpDMP1)/SBF allow the visualization of both the resin 391 and individual fibril due to a lower density of collagen locally. 392 A comparison between mineralized collagen fibrils without or 393 with staining (Figure 3F,F', respectively) confirms the 394 precipitation of fibrils with the cross-striated pattern. 395 Observations of an unmineralized collagen matrix without 396 staining (Coll, Figure 3H), where the resin is darker than 397 collagen, confirm that the contrast in mineralized samples comes from the spreading of mineral (ions or precipitates) 398 over the collagen fibrils. Note that we cannot conclude on the 400 involvement of DMP1 in mediating the size of fibrils,²⁹ since 401 the average diameter of collagen fibrils appears unmodified 402 (~100 nm) regardless the rpDMP1 concentration.

403 Going further, thermogravimetric analysis (TGA) inves-404 tigations were performed to better characterize the effect of rpDMP1 concentration on apatite formation (Figure S3). 405 Considering the standard deviation recorded for the matrices 406 $(\pm 5 \,^{\circ}C)$, the difference in mineralization degree should be 407 considered as a trend here. After SBF bath, the mineral content 408 increases for low rpDM1 concentrations (from ~3.5% to 409 ~18%), whereas it does not change significantly at a high 410 content of rpDMP1 (from ~12.5% to ~15%). Since both 411 matrices exhibit spherulites on the surface after SBF bath 412 (Figure 2F and Figure S2C), it indicates that the main mineral 413 content is localized inside the matrix in the form of 414 nanoprecipitates. 415

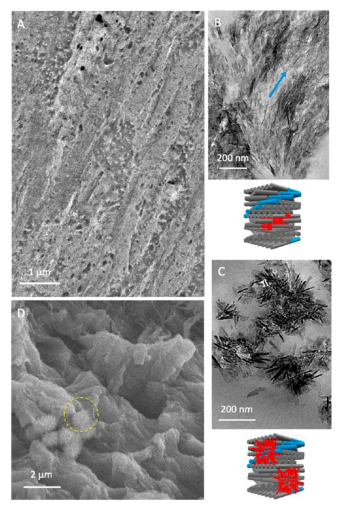
In addition, the ζ -potential was investigated to help 416 understand the interactions between collagen, rpDMP1, and 417 the mineral. Before the SBF bath, the global surface charge 418 tends to be neutral in the presence of rpDMP1 whatever the 419 concentration of the protein (-0.79 (± 0.66) mV and 0.22 420 (± 0.78) mV) for low and high rpDMP1 concentrations, 421 respectively). Indeed, the Coll/SBF matrix exhibits a negative 422 ζ -potential (-4.3 (±2.5) mV.⁸ This result agrees with a partial 423 coating of collagen by the selected DMP1 domain, which 424 includes the two collagen binding sites²⁹ and the highly 425 conserved acidic peptide.³⁷ Indeed, it is reported that DMP1 426 binds to the N-telopeptide region in collagen.^{29,45} After 427 immersion in SBF, Coll/SBF(1rpDMP1)/SBF exhibits a more 428 negative ζ -potential (-9.77 (±0.48) mV)), while it is slightly 429 unchanged for Coll/SBF(10rpDMP1)/SBF (-1.79 (±0.33) 430 mV). The ζ -potential of HA being negative between pH 5 and 431

 $_{432}$ 8 (-5 to -37 mV respectively)⁶² is in agreement with the $_{433}$ TGA analysis.

To go deeper in the understanding of how the suprafibrillar 435 confinement affects the spreading of rpDMP1 over collagen, 436 the peptide was entrapped in a scaffold mimicking the fibrillar 437 arrangement of osteoid tissue (Figure 4A,B). As mentioned 438 above, this means that (i) the fibrils network is not as dense as 439 the previous mature bone model with larger interfibrillar spaces 440 which are above the micron size (macroporous gel) and (ii) 441 the collagen fibrils are not organized (no plywood geometry). 442 Note that spherulitic crystals form in the control matrix 443 without peptide (CollOsteoid/SBF/SBF) as shown by SEM 444 (Figure 4C) and TEM (Figure 4D), which is in agreement 445 with previous observations on a matrix with a collagen gradient 446 concentration.⁸

Interestingly, "dot-like" precipitates are also observed, but 447 448 they do not localize inside the fibrils (Figure 4A). Although 449 most of the investigations have focused on electrostatic 450 interactions to explain the intrafibrillar infiltration of both 451 mineral and proteins in collagen, this difference in apatite 452 crystals distribution in osteoid- and mature bone-like matrices 453 evidence that the suprafibrillar confinement provided by the 454 collagen assembly in bone tissue (i.e., cholesteric geometry) 455 plays a key role on this phenomenon. Thus, it may explain the 456 need of other proteins to infiltrate collagen in models lacking 457 biomimetic interfibrillar spaces (<1.7 nm).⁴⁵ The fact that 458 collagen here is continuously in contact with ions (even before 459 mineralization) as occurring in bone strengthens the need of a 460 balance between osmotic equilibrium and electroneutrality for 461 intrafibrillar mineralization.⁶³ In addition, the use of biomi-462 metic collagen interfibrillar spaces (in both osteoid- and 463 mature-like matrices) contradicts that rpDMP1 favors the 464 templating of crystal growth.¹⁴ Conversely, according to (i) 465 TEM and SEM investigations where it is observed that the size 466 of spherulites consist of either "dot-like" aggregate or "mature" 467 apatite platelets with (Figures 2F, 3B, and 4A,B) or without 468 rpDMP1 (Figures 2G and 4C,D) respectively, (ii) the mineral 469 amount found by TGA, and (iii) the resulting surface charge 470 probed by ζ -potential, we confirm its role as a growth 471 inhibitor.³³ This effect may be related to the confinement $_{472}$ which promotes the protein folding by destabilizing the $_{473}$ unfolded state $_{64,65}^{64,65}$ and thus further specific interactions 474 between rpDMP1 and apatite nuclei; the structural character-475 istics of growing apatite being driven by the involvement of 476 ionic substitutions (specifically from carbonate ions).⁶⁶

Toward Evidence of the Specific Activity of rpDMP1 477 478 under Confinement. When polyAsp is supplemented to 479 collagen, it is difficult to observe a difference in mineral 480 spreading over the collagen matrix before immersion in SBF 481 (Coll/SBF(polyAsp), Figure 5A) as compared to rpDMP1. 482 Here also, a lower concentration of polyAsp (5.75 μ g/mL) was 483 used than that in the literature (usually between 10 and 100 $_{484} \mu g/mL$) (i) to reach at least 2 magnitude order lower amounts 485 of negative charge and (ii) to reproduce a more "realistic" 486 collagen to NCPs ratio since such a high polyAsp 487 concentration can be used for one single fibril.³ Note that 488 our samples were not lyophilized before characterization, as it 489 is commonly performed with this polymer in the literature.⁶⁷ 490 This is of importance since the resulting phase and degree of 491 mineralization are here related to a hydrated biomimetic 492 process. The difference between polyAsp and rpDMP1 in 493 apatite mineralization becomes clearer after the SBF bath 494 (Coll/SBF(polyAsp)/SBF, Figure 5B,C). TEM observations



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Figure 5. Calcium phosphate precipitation investigated by EM in the presence of polyAsp containing mature bone-like matrix. Observations by TEM of unstained ultrathin sections of matrices before and after SBF immersion: (A) Coll/SBF(polyAsp) and (B,C) Coll/SBF(polyAsp)/SBF, respectively. The corresponding schematic representation of the apatite crystal (red)/collagen fibril (gray) relationship in a 3D perspective is shown. The lack of staining in (A) indicates that there is an homogeneous spreading of SBF ions over the collagen. After immersion in SBF (B, C), HA platelets are observed in the matrix; they either (B) co-align with the long axis of the fibril (blue arrow) or (C) form spherulites as schematically presented, respectively, below the (B) and (C) TEM micrographs. (D) SEM micrograph of Coll/SBF(polyAsp)/SBF showing that spherulites (yellow dashed circle) are also observed inside the matrix.

reveal large domains where the mineral platelets co-align with 495 the collagen fibrils (Figure 5B), as seen in mature bone (Figure 496 1C) strengthening the affinity of polyAsp for gap regions in 497 line with previous conclusions in the literature.^{44,45,68} 498 However, in contrast to rpDMP1 results, nonbiomimetic 499 spherulitic crystals are also observed in the matrix (Figure 500 5C,D), strengthening that only small molecules such as 501 osteocalcin can penetrate alone the intrafibrillar space.⁶⁹ 502

These observations evidence the specificity of biological 503 rpDMP1 versus synthetic polyAsp amino-acids sequences in 504 mediating apatite mineralization and also suggest that, in 505 addition to its role as growth inhibitor, it may prevent 506 homogeneous nucleation.³³ Finally, because the rpDMP1 507 sequence is not phosphorylated, this result tends to strengthen 508

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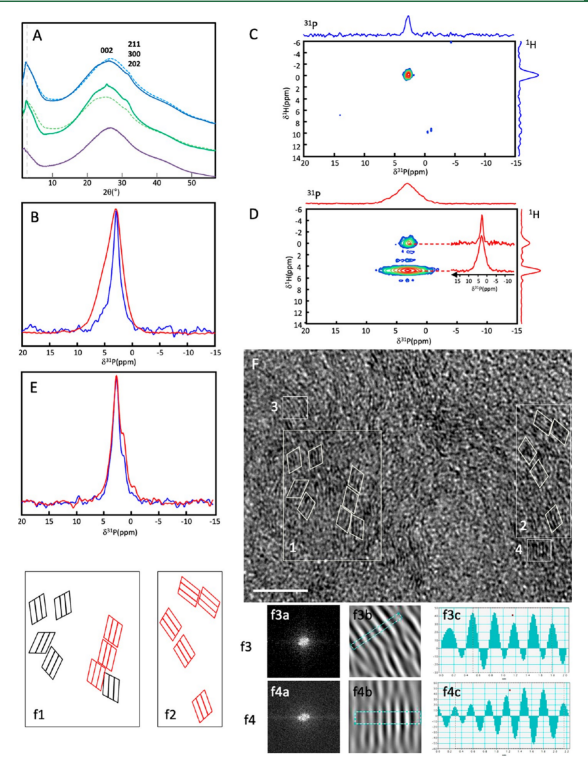


Figure 6. Characterization of the mineral phase. (A) 1D radial average of the WAXD patterns of matrices at 2.5 μ g/mL, without (Coll/SBF(1rpDMP1), dashed green) and with (Coll/SBF(1rpDMP1)/SBF, plain green) immersion in SBF; at 25 μ g/mL, without (Coll/SBF(10rpDMP1), dashed blue) and with (Coll/SBF(10rpDMP1)/SBF, full blue) immersion in SBF, and a collagen matrix as control (purple). The vertical gray dashed line points out the shift of the reflection which corresponds to the lateral distance between collagen molecules. (B) 1D ³¹P CP MAS spectra (CT = 10 ms) of the hybrid matrices Coll/SBF(1rpDMP1) (blue) and Coll/SBF(1rpDMP1)/SBF (red). (C) 2D ¹H-³¹P HETCOR spectrum (CT = 10 ms) of Coll/SBF(1rpDMP1) with extracted ³¹P slices at δ (¹H) = 0 and 4.9 ppm. (D) 2D ¹H-³¹P HETCOR spectrum (CT = 10 ms) of Coll/SBF(1rpDMP1)/SBF. (E) 1D ³¹P CP MAS spectra (CT = 10 ms) of the hybrid matrices Coll/SBF(1rpDMP1) with extracted ³¹P slices at δ (¹H) = 0 and 4.9 ppm. (D) 2D ¹H-³¹P HETCOR spectrum (CT = 10 ms) of Coll/SBF(1rpDMP1)/SBF. (E) 1D ³¹P CP MAS spectra (CT = 10 ms) of the hybrid matrices Coll/SBF(10rpDMP1) with extracted ³¹P Slices at δ (¹H) = 0 and 4.9 ppm. (D) 2D ¹H-³¹P HETCOR spectrum (CT = 10 ms) of Coll/SBF(1rpDMP1)/SBF. (E) 1D ³¹P CP MAS spectra (CT = 10 ms) of the hybrid matrices Coll/SBF(10rpDMP1) (blue) and Coll/SBF(10rpDMP1)/SBF (red). (F) TEM micrograph of Coll/SBF(1rpDMP1)/SBF (scale bar = 5 nm) showing crystalline HA nanodomains. (f1) and (f2) are schematic representations of the selected areas 1 and 2 showing the orientation of the HA (210) planes; some crystalline nanodomains appear to align along the same direction (red). (f3) and (f4) FFT (f3a and f4a) and inverse FFT (f3b and f4b) performed on the selected areas 3 and 4, confirming the presence of crystalline nanodomains. The *d* spacings are measured using the inverse FFT profile along one direction (f3c and f4c).

509 that the organic phosphate concern may be attributed to 510 polyadenosine diphosphate ribose.⁷⁰

Structural Features of rpDMP1-Mediated Mineral 511 512 under Confinement. WAXD studies were performed to 513 better characterize the mineral phase and collagen network. 514 The 1D radial average of the WAXD patterns is shown in 515 Figure 6A. Without the SBF extra bath, the signal from 516 collagen fibrils is strong, revealing a low degree of 517 mineralization (dashed green and blue lines). The character-518 istic HA diffraction peaks, that is, (002) and the merged (211), 519 (300), (202) reflections, are observed more clearly after the 520 SBF bath (full green and blue lines), especially for Coll/ 521 SBF(1rpDMP1)/SBF, confirming an increase in mineral 522 content. In addition, the lateral distance between collagen 523 molecules decreases from 1.5 to 1.1 nm in the matrices loaded 524 with rpDMP1 (with or without SBF bath) in comparison to 525 the pure collagen matrix (full purple line) as observed by the 526 peak shift toward high angles (vertical gray dashed line). This 527 distance is shorter than that reported for a mineralized dense 528 collagen scaffold without protein,⁷ but agrees with densely 529 packed microfibrils⁷¹ and fibrillar collagen in a dry state.⁷ 530 Since the sample is studied in its native hydrated form, this 531 decrease in the intermolecular distance in addition to the 532 contrast observed in fibrils by TEM confirm that rpDMP1 533 localizes inside the gap zones. The peptide may colocalize with 534 the mineral as shown for polyelectrolytes.⁶³ Besides confine-535 ment, SBF ions are needed for the protein to localize in 536 intrafibrillar spaces, especially calcium ions, considering the s37 model of a periodic assembly of DMP1 into a β -sheet template 538 with high calcium-binding capacity.¹⁴ Then the Ca-binding 539 protein can interact with a significant fraction of the π , that is, 540 possibly covalently bond to the collagen.⁷³ At a critical 541 concentration of Ca^{2+} , the apatite nucleation which strongly 542 depends on the degree of confinement⁷⁴ that occurs inside 543 collagen. However, since apatite forms without organic 544 additives under collagen confinement,⁸ it shows that 545 rpDMP1 increases the local ions concentration but does not 546 initiate the nucleation here.

To investigate the presence of calcium-phosphate minerals 547 548 described as transient phases in bone formation^{51,75,76} and 549 better understand the local phosphate environment in 550 collagen-containing rpDMP1, ³¹P ssNMR experiments were 551 performed (Figure 6B-E). The ³¹P CP MAS spectrum of 552 Coll/SBF(1rpDMP1) displays a single ³¹P resonance centered 553 at 2.8 ppm, typical of apatite (Figure 6B, blue). Interestingly, 554 the 2D¹H-³¹P HETCOR spectrum of Coll/SBF(10rpDMP1) 555 does not show the correlation resonance characteristic from 556 HPO₄²⁻ (Figure 6C), excluding the presence of a hydrated 557 disordered surface layer.^{66,77} In addition, the line width of the 31 P resonance (LW = 1 ppm) is thinner than matrices prepared 558 559 without rpDMP1 (Coll/SBF, LW = 2 ppm) (Figure S4). Both $_{560}$ the thinner resonance and the absence/low amount of HPO₄²⁻ 561 surface species demonstrate that the mineral "dot-like" 562 precipitates of the matrices are highly crystalline and that 563 rpDMP1 drives the higher crystallinity of apatite. This result is 564 in agreement with previous works proposing that DMP1 565 impacts on the initial stages of apatite formation, providing a 566 structural local order.¹⁴ Interestingly, DMP1 is found more 567 abundant in the boundary between the dentin and enamel 568 where apatite is well crystallized.³⁰

⁵⁶⁹ Finally, it is interesting to note that such "dot-like crystals" ⁵⁷⁰ were found at the very early stages of normal *in vivo* ⁵⁷¹ calcification.¹ After the SBF bath, the ³¹P LW in CP MAS spectrum increases (Figure 6B, red), reaching the value found 572 for bone apatite (4.1 ppm).⁷⁸ The fact that the 2D ${}^{1}H{-}^{31}P$ 573 HETCOR spectrum of Coll/SBF(1rpDMP1)/SBF (CT = 10 574 ms) (Figure 6D) displays the two typical resonances of bone 575 minerals, that is, $\delta({}^{31}P) = 2.8$ and 3.2 ppm, shows that the 576 apatite spherulites observed at the surface of the matrices by 577 microscopy (Figure 2F) dominate the spectroscopic signature 578 here.

At a higher rpDMP1 content, 1D ³¹P CP-MAS spectra 580 (Figure 6E) display two shoulders (centered at 1.5 and -1 581 ppm) in addition to the apatite resonance at 2.8 ppm before 582 and after SBF. These resonances correspond to the HPO₄²⁻ 583 ions from the OCP phase. Noticeably, the presence of a 584 relatively low content of rpDMP1 leads to the physiological 586 concentration of protein stabilizes an additional CaP phase 587 like OCP. Such high concentrations are classically used in the 588 literature, questioning the role of DMP1 in stabilizing the ACP 589 phase. ^{33,34}

HRTEM observations performed on ultrathin sections of 591 Coll/SBF(1rpDMP1)/SBF (Figure 6F and Figure S5) confirm 592 the highly crystalline nature of the "dot-like" precipitates. 593 Spaced thin layers that are mostly concentric, that is, onion-like 594 morphology that appears to be composed of crystalline 595 nanodomains (~1 nm) (some are depicted in areas 1 and 596 2), are observed. Schematic representations of two selected 597 areas (1 and 2) show the orientation of possible crystal planes 598 (f1 and f2). Some crystalline nanodomains appear to align 599 along the same direction (in red). Their morphology is difficult 600 to identify; they are presented here as parallelogram. In (f3) 601 and (f4), fast-fourrier tranform (FFT) (f3a and f4a) and 602 inverse FFT (f3b and f4b) performed on the (3 and 4) selected 603 areas confirm the presence of crystalline nanodomains. The 604 crystal planes were indexed by comparing the measured d_{605} spacings using the FFT profile along one direction with 606 calculated values of HA (f3c and f4c). Two lattice spacings 607 were measured as 0.31 and 0.34 nm, corresponding to the 608 known (210) and (002) crystal planes of apatite (Figure 6F 609 and Figure S5, respectively). Among the 20 analyzed planes, 610 80% correspond to the (210) and the remaining 20% to the 611 (002).

The possible consequence of a crystalline apatite nano- 613 domain versus an ACP phase can prevent the formation of 614 pathological calcifications. Indeed, ACP as the first solid phase 615 of calcium phosphate formed in bone would delay apatite 616 formation, allowing the precipitation of other transient phases. 617 This hypothesis is in agreement with the fact that the 618 mineralization process of bone is described to occur rapidly 619 as soon as collagen molecules self-assemble into collagen fibrils 620 in the extracellular space.⁷⁹ In fact, according to our 621 knowledge, there is no report on bone pathologies related to 622 the presence of one of the transient apatite phases, although it 623 is extensively described in synthetic models lacking interfi-624 brillar confinement *in vitro* as discussed above. 625

CONCLUSION

The bone ECM is a dynamic (in terms of structure, 627 remodelling, and ECF) and complex environment where the 628 confinement effect occurs that is not only critical for the cells 629 behavior⁸⁰ but also for the proteins and even the solvent 630 (water).⁶⁴ Indeed, confinement provided by suprafibrillar 631 organization of collagen strongly impacts the activity of 632 mineralizing polymers. The results suggest the occurrence of 633

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634 sequential cooperative effects during the early stages of bone 635 apatite precipitation through the formation of Ca-DMP1 636 complex, which then binds to collagen allowing the 637 concentration of apatite ion precursors in gap regions. 638 Formation of highly apatite nuclei and inhibition of both 639 homogeneous nucleation and crystal growth are consistent 640 with a common role of DMP1 controlling the physiological 641 (versus pathological) bone formation. Strikingly, these effects 642 are reached with the nonphosphorylated 57-KD amino acid 643 sequence under confinement. This versatile bone-like model 644 will be useful to provide insights into the role of other bone 645 components (e.g., citrate, proteoglycans, or even different 646 sequences related to NCPs kinetic of maturation (i.e., post-647 translation modification)) during the successive events that 648 orchestrate mineralization.

ASSOCIATED CONTENT 649

650 Supporting Information

651 The Supporting Information is available free of charge at 652 https://pubs.acs.org/doi/10.1021/acs.biomac.1c00206.

Composition of the solutions used to mineralize 653 collagen matrices, SEM and TEM images, EDX spectra, 654 and TGA curves (PDF) 655

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Notes

The authors declare no competing financial interest. 700

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