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Title Page
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      Evaluation of the inflammatory responses to sol-gel coatings with distinct
 3
      biocompatibility levels
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34 Title: Evaluation of the inflammatory responses to sol-gel coatings with distinct
35 biocompatibility levels

Abstract: The immune system plays a crucial role in determining the implantation outcome, 36 and macrophages are in the frontline of the inflammatory processes. Further, cellular 37 oxidative stress resulting from the material recognition can influence how cell responses 38 develop. Considering this, the aim of this study was to study oxidative stress and 39 macrophages phenotypes in response to sol-gel materials with distinct in vivo outcomes. Four 40 materials were selected (70M30T and 35M35G30T, with high biocompatibility, and 50M50G 41 and 50V50G, with low biocompatibility). Gene expression, immunocytochemistry and 42 43 cytokine secretion profiles for M1 and M2 markers were determined. Moreover, oxidative stress markers were studied. Immunocytochemistry and ELISA showed that 50M50G and 44 50V50G lead to a higher differentiation to M1 phenotype, while 70M30T and 35M35G30T 45 promoted M2 differentiation. In oxidative stress, no differences were found. These results 46 show that the balance between M1 and M2, more than individual quantification of each 47 phenotype, determines a biomaterial outcome. 48

49 Keywords: Inflammation; macrophage plasticity; biomaterials; oxidative markers; implants

50 1. Introduction

Biocompatibility describes the appropriate biological requirements of biomaterials for medical application as well as the ability of said materials to perform with an host response in a specific application ⁽¹⁾. It is determined by the coordination of the host homeostatic mechanisms, which are disturbed upon implantation, and the consequent immune response to injury ⁽²⁾. The coordinated activation, type and action of highly specialized immune cells depends of the nature and site of the wound/damage ⁽³⁾. Macrophages represent the first line of defense on the innate immunity, being most known by their phagocytic capabilities.

Besides their major effector function of eliminating and inactivating pathogens, these cells 58 boost properties such as the clearance of apoptotic cells throughout the lifespan of an 59 organism, homeostasis and activation of tissue repair processes ⁽⁴⁾. Macrophages have the 60 capability to enter into distinct tissues, modulate and differentiate into specialized phenotypes 61 according to microenvironmental cues, stimuli from growth factors, cytokines, and 62 chemokines present in biological fluids (e.g. blood). In the case of implanted biomaterials, 63 these events are part of a whole process that could culminate in a foreign body reaction (FBR) 64 to the material ⁽⁵⁾. Once activated, macrophages can exhibit a spectrum of polarization states 65 depending on their functional nature, adopting a pro-inflammatory phenotype (M1) or an anti-66 67 inflammatory phenotype (M2), with distinct surface markers and/or different gene expression profiles. When a biomaterial is implanted into the organism, this cascade of events is 68 triggered, allowing the direct and initial migration of M1 macrophages toward the 69 implantation site, provoking the necessary inflammatory response ⁽⁶⁾, which is characterized 70 by the secretion of pro-inflammatory cytokines and chemokines, such as tumor necrosis factor 71 α (TNF- α) and interleukin 1- β (IL-1 β)⁽⁷⁾. The prolonged presence of this phenotype can lead 72 to a state of chronic inflammation, ultimately leading to implant rejection ⁽⁸⁾. The anti-73 inflammatory M2 macrophages establish themselves upon signals released by basophils, 74 including cytokines like interleukin-10 (IL-10) and interleukin-4 (IL-4) ⁽⁹⁾. This anti-75 inflammatory state is distinguishable by its role on immunoregulation, matrix deposition and 76 tissue remodeling processes ⁽⁷⁾. The increase of M2 subsets in the biomaterial surrounding 77 environment, towards a positive value of M2:M1 ratio, has been suggested as the key to a 78 positive outcome of the implanted material (10). However, the greater presence of M2 79 macrophages could increase of foreign body giant cells (FBGC) in situ, when its 80 predominance is too prolonged ⁽⁵⁾. Hence, this ratio as a marker for biocompatibility must be 81 carefully approached. 82

Oxidative stress derives as a consequence of the surgical creation of a wound and 83 implantation, being influenced by the material properties, the degree of initial inflammation 84 and the immediate stress resulting from the procedure, occurring at all stages of the response 85 to a biomaterial. The resulting reactive oxygen species (ROS), reactive nitrogen species 86 (RNS) and lipid peroxidation subproducts (e.g. malondialdehyde - MDA) act as chemo-87 attractants and signaling molecules during healing, and are often associated with phenotypic 88 shifts of immune cells and modulation of cell response to a determined material ⁽¹¹⁾. Redox 89 interactions are responsible for stabilizing these oxidation products and glutathione (GSH), 90 synthesized from glycine, cysteine, and glutamic acid, is the most important redox-regulating 91 thiol, acting as a substrate of glutathione peroxidase (GPx) ⁽¹²⁾. The antioxidant function of 92 GSH is due to the oxidation of the sulfhydryl group (-SH), and the ratio between glutathione 93 disulfide (GSSH) and GSH is an indicator of the cellular redox potential ⁽¹²⁾. Differences in 94 ROS generation and scavenging between M1 and M2 macrophages have been studied ⁽¹³⁾. 95 Superoxide generation, namely hydrogen peroxide, is typically increased and associated to the 96 M1 macrophage phenotype, due to its phagocytic/microbiocidal activity, which depends on 97 the synthesis of ROS and RNS. Moreover, as M2 phenotypes are usually described as being 98 angiogenic, anti-oxidant and dependent on oxidative phosphorylation. A low expression of 99 100 pro-oxidants NOX2 e NOX5 and high levels of SOD, GPx and CAT have been described as required for M2 macrophage polarization ⁽¹³⁾, thus confirming the oxidative metabolic 101 differences for these immune cell subpopulations ⁽¹⁴⁾. 102

Upon implantation on a living organism, the blood is the first organic fluid in contact with the implant, leading to protein adsorption by the surface whose type, composition, quantity and conformation might impair the final outcome ⁽¹⁵⁾. This process is dependent on the physicochemical characteristics of the surface of the material and can ultimately modulate macrophage and monocyte activation and migration to the implantation site ⁽¹⁶⁾. In previous studies ⁽¹⁷⁾, we showed that a greater deposition of complement proteins onto a biomaterial is intrinsically correlated with their biocompatibility in a living host. The oxidative stress in response to the implantation process and the material itself might also directly impair the immune cellular response/differentiation and ultimately affect the implant outcome.

Following this premise, this experimental work focuses on the study of the polarization/plasticity of activated macrophages to previously described sol-gel materials with distinct biocompatibility reactions *in vivo* and the correlation of between the predominance of a determined macrophage phenotype with the oxidative stress responses.

116 **2.** Materials and methods

117 2.1. Material selection, synthesis, and preparation

Sol-gel technology was employed to synthetize four different materials using 118 methyltrimethoxysilane (MTMS), 3-glycidoxypropyl-trimethoxysilane (GPTMS), tetraethyl 119 orthosilicate (TEOS) and triethoxyvinylsilane (VTES) precursors in the proportions shown in 120 Table 1. These materials, designed in previous works, were selected due to their distinct 121 biocompatibility outcomes *in vivo* ⁽¹⁷⁾⁻⁽¹⁹⁾. For their synthesis, the corresponding alkoxysilane 122 amounts were diluted with 2-propanol (50 % vol) and hydrolyzed adding the stoichiometric 123 amount of acidified aqueous solution (0.1 M HNO₃). All the employed reagents were 124 purchased from Sigma-Aldrich (Merck KGaA, Darmstadt, Germany). The sol-gel 125 preparations were left stirring for 1 h and resting for another 1 h. The coatings were prepared 126 immediately after this resting. For that, grade- 4 Ti discs (12 mm diameter, 1 mm thick; 127 Ilerimplant-GMI S.L., Lleida, Spain) were employed as substrate for the coatings. Bare discs 128 were superficially pre-treated with a sandblasting and acid-etching treatment (SAE) 129 previously described ⁽²⁰⁾. Then, the sol-gel solutions were applied as coatings using a KSV 130 DC dip-coater (Biolin Scientific, Stockholm, Sweden). Discs were submerged into the 131

corresponding sol-gel (60 cm min⁻¹-speed) and kept immersed in it for one minute. Then, the
samples were taken out at 100 cm min⁻¹. Finally, heat treatments at 80 °C to 70M30T and
35M35G30T, and at 140°C to 50M50G and 50V50G materials were carried out for 2 h.

135 2.2. In vitro assays

136 **2.2.1.** Cell culture

For the distinct experiments, mouse murine macrophage cells (RAW 264.7) were cultured on
the discs in 48-well NUNC plates (Thermo Fisher Scientific, NY, USA) at 37 °C in a
humidified (95 %) CO₂ incubator using as culture medium Dulbecco's Modified Eagle
Medium (DMEM; Gibco, Thermo Fisher Scientific) with 10 % of fetal bovine serum (FBS;
Gibco) and 1 % of penicillin/streptomycin (Gibco).

142 2.2.2. Cell fixation for SEM imaging

After 72 h of incubation, samples were washed once with PB 0.1 M and fixed with 3.5 % 143 glutaraldehyde for 45 minutes, at 37 °C, in the dark. After washing twice with PB 0.1 M, the 144 preparations were incubated with 2 % osmium for 1 h in the dark. Afterwards, samples were 145 washed with dH₂O to eliminate any osmium residues and a chain with crescent concentrations 146 of ethanol was performed for dehydration. The critical point drying was made through 147 incubation with hexamethyldisilazane (HDMS; Sigma-Aldrich). Next, samples were 148 examined in a field emission scanning electron microscope (FESEM; ULTRA 55, ZEISS 149 Oxford Instruments) at 2kV of voltage. 150

151 2.2.3. Immunocytochemistry double staining

After 24 and 72 h, samples were fixed in 4 % paraformaldehyde for 10 min (Sigma-Aldrich)
and washed five times in 1x PBS. The samples were blocked in 1x PBS containing 0.5 %
BSA and 1 % Triton X-100 (Sigma-Aldrich). They were incubated with donkey anti-mouse
CD206 primary antibody (Abcam, Cambridge, UK) diluted 1:250 in PBS containing 0.5 %

BSA and 0.5 % Tween-20 (Sigma-Aldrich), overnight at 4°C. The discs were then washed 156 five times in 1x PBS and incubated with a mixture of secondary antibodies composed of Goat 157 anti-Donkey Biotin (Jackson ImmunoResearch Europe, Ltd., Cambridgeshire, UK) diluted 158 1:500 and Streptavidin Alexa Fluor 647 (Thermo Fisher Scientific) diluted 1:500 for 1 h at 159 160 room temperature. Cells were washed five times with wash buffer (1x PBS with 0.5 % Triton X-100) and incubated with the primary antibody IL7-R (Santa Cruz Biotechnology, Dallas, 161 TX, USA) at 4 °C overnight. After five washes with wash buffer, the discs were incubated 162 with the secondary antibody Goat anti-Rabbit Alexa Fluor 488 (Thermo Fisher Scientific) for 163 1 h at room temperature. After the next five washes with wash buffer, the discs were 164 incubated with DAPI (Roche, Basel, Switzerland) for another hour to stain the cell nuclei. 165

The discs were then removed from the wells, mounted on coverslipped slides with mounting medium to prevent the sample from drying out (4.8 % poly(vinyl alcohol-co-vinyl acetate), 12 % glycerol, 0.2 M Tris-HCl, 0.02 % sodium azide) and stored at 4°C until the fluorescence microscopy analysis (Keyence International, Mechelen, Belgium).

170 2.2.4. RNA extraction, cDNA synthesis and quantitative real-time PCR measurements

After 24 and 72 h, total RNA was extracted using TRIzol (1 M guanidine thiocyanate, 1 M 171 ammonium thiocyanate, 3 M sodium acetate, 5 % glycerol, 38 % aquaphenol). To each 172 sample 300 µL of TRIzol were added followed by an incubation at room temperature. After 173 centrifugation (5 min, 13000 rpm, 4 °C), 200 µL of chloroform were added to the supernatant, 174 and the samples were centrifuged (5 min, 13000 rpm, 4 °C). The aqueous layer was mixed 175 with 550 µL of isopropanol and kept at room temperature for 10 min. Samples were 176 centrifuged (15 min, 13000 rpm, 4 °C), and washed twice with 0.5 mL of 70 % ethanol. The 177 resulting pellet was dissolved in 30 µL of RNAse free water. RNA concentration, integrity, 178 and quality were measured using NanoVue® Plus Spectrophotometer (GE Healthcare Life 179 Sciences, Little Chalfont, UK). Approximately 1 µg of total RNA was converted into cDNA 180

using PrimeScript RT Reagent Kit (Perfect Real Time; TAKARA Bio Inc., Shiga, Japan) and
the reaction was conducted with the following conditions: 37 °C for 15 min, 85 °C for 5 secs
and a final hold at 4°C. The resulting cDNA quality and quantity was measured using a
NanoVue® Plus Spectrophotometer (GE Healthcare Life Sciences), then diluted in DNAsefree water to a concentration suitable for reliable qRT-PCR analysis and stored at -20 °C.

To evaluate the effects of the materials on the inflammatory responses, genes corresponding 186 to pro and anti-inflammatory phenotypes were selected (Table 2). GADPH was used as a 187 housekeeping gene. Primers were designed using DNA sequences for these genes available 188 from NCBI (https://www.ncbi.nlm.nih.gov/nuccore), employing PRIMER3plus software tool 189 (http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi) and purchased to Thermo 190 Fischer Scientific. Quantitative real-time PCR (qRT-PCR) were carried out in 96-well plates 191 (Applied Biosystems[®], Thermo Fisher Scientific) and individual reactions contained 1 µL of 192 cDNA, 0.2 µL of specific primers (forward and reverse at 10 µM L⁻¹) and 5 µL of SYBR 193 Premix Ex Taq (Tli RNase H Plus; TAKARA, Bio Inc., Shiga, Japan) in a final volume of 10 194 195 µL, and were carried out in a StepOne Plus[™] Real-Time PCR System (Applied Biosystems[®]). The cycling parameters were an initial denaturation step (95°C, 30 s) followed 196 by 95 °C for 5 s and 60 °C for 34 s, for 40 cycles. The final melt curve stage comprised a 197 cycle at 95 °C for 15 s and at 60 °C, for 60 s. Fold changes were calculated using the $2^{-\Delta\Delta Ct}$ 198 method and the data was normalized in relation to the blank wells (without any material). 199

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2.2.5. Cytokine quantification by ELISA

To measure secreted cytokines (TNF- α , IL-1 β , TGF- β and IL-10), the cell culture supernatants used for immunocytochemistry were collected and frozen until further analysis. The concentration of these cytokines was determined using an ELISA (Invitrogen, Thermo Fisher Scientific) kit and according to the manufacturer's instructions.

After 24 and 72 h, cells were washed three times with PBS and incubated at 4 °C for 10 min 206 in lysis buffer (0.2 % Triton X-100, 10 mM Tris-HCl, pH 7.2). Glutamic acid, glutathione 207 208 (GSH) and glutathione disulfide (GSSG) concentrations were quantified chromatographically using the method proposed by Reed ⁽²¹⁾. Shortly, this method is based in the reaction of the 209 Sanger Reactant (1-fluoro-2,4-dinitrobencene) with amino groups and iodoacetic acid to 210 block free thiol groups. Samples were measured after derivatization using a high-performance 211 liquid chromatographic system equipped with a diode array detector. Glutathione peroxidase 212 activity (GPx) was determined by the desaparition of NADPH monitored at 340 nm as 213 proposed Lawrence et al.⁽²²⁾. Briefly, a solution containing 50 µL of samples, 550 µL of 214 potassium phosphate buffer 0.1 M pH 7.0, EDTA 1 mM and NaN₃ 1 mM was mixed with 100 215 μ L GSH disulfide reductase (0.24 U mL⁻¹), 100 μ L glutatione reduced 1 mM and 100 μ L 216 NADPH 0.15 mM. The resulting solution was incubated for 3 min at 37 °C. Then, 100 µL of 217 hydrogen peroxide 1.5 mM were added to start the reaction. Glutathione reductase activity 218 was determined using the method proposed by Smith and et al.⁽²³⁾. The method consists in 219 220 monitoring spectroscopically the 2-nitrobenzoic acid formation. This is formed as subproduct of the GR catalyzed reduction of GSSG to GSH in presence of 5,5'-dithiobis(2-nitrobenzoic 221 acid) (DTNB). The GSSG reduction was started by adding 25 µL of sample to a solution 222 containing 450 µL 0.2 M phosphate buffer pH7.5 and 250 µL of DTNB 3 mM prepared in 10 223 mM phosphate buffer, 50 µL of 2 mM NADPH and 50 µL of 10 mM EDTA. Total volume 224 was adjusted to 1 mL using ultrapure water and the wavelength set at 412 nm. MDA 225 concentration was determined chromatographically using an HPLC system using Richard et 226 al. proposed method ⁽²⁴⁾ with modifications introduced by Romero et al. ⁽²⁵⁾. Sample 227 preparation consisted in mixing samples (100 µL) with 0.75 mL of thiobarbituric acid with 228 0.37 % and perchloric acid 6.4 % (2:1, v/v) and heated to 95 °C for an hour. Then, pH was 229

adjusted to 6 and precipitates removed by centrifugation (10000 rpm, 1 min). Separation was 230 carried out in a HPLC system equipped with a C18 250x4.6 mm 5 µm chromatographic 231 column using an isocratic separation. Flow was set at 1 mL min⁻¹ and fluorescence detector 232 was set to 527 nm for excitation and 532 nm for emission. Mobile phase consisted in 50 mM 233 phosphate buffer (pH 6.0): methanol (58:42, v/v) and 1,1,3,3-tetramethoxypropane was used 234 as standard solution. All standards and mobile phases were prepared daily. Protein levels were 235 determined from cell culture lysates using a Pierce[™] BCA Protein Assay Kit (Thermo Fisher 236 Scientific) and used to normalize oxidative stress values. 237

238 **2.3.** Statistical analysis

Based on the normal distribution and equal variance assumption test, the data were analyzed via one-way analysis of variance (ANOVA) with Newman-Keuls post hoc test and expressed as mean \pm standard deviation (SD). Statistical analysis was performed using GraphPad Prism 5.04_software (GraphPad Software Inc., La Jolla, CA, USA). The asterisk (*) indicates statistically significant ($p \le 0.05$) differences between the four materials.

244 **3.** Results

245 **3.1. Morphological analysis**

To evaluate cellular morphology, macrophages seeded on the distinct materials were studied with SEM. The obtained images of cell spreading revealed that macrophages seeded for 72 h on 70M30T and 35M35G30T treatment acquired an elongated morphology (Fig. 1a', 1b'). When seeded on 50M50G and 50V50G, macrophages adhered and spread to a typical rounded shape (Fig. 1c', 1d').

251 **3.2.** Immunocytochemistry double staining

To evaluate the expression of markers associated with M1 and M2 phenotypes, immunocytochemistry was performed. IL7-R, an M1-phenotype marker, showed significant increased fluorescence of the macrophage cultures on the 50V50G and 50M50G when
compared to the other two materials (Fig. 2). No differences were observed on the CD206
M2-marker fluorescence intensity.

257 **3.3. Gene expression analysis**

The expression of pro and anti-inflammatory markers by the RAW264.7 cells cultured onto 258 the distinct materials is shown in Fig. 3. At 24h, the expression of TNF- α was significantly 259 higher on 35M35G30T, generally decreasing at 72 h on all materials (Fig. 3a). On the other 260 hand, IL1- β expression peaked at 24 h and then decreased on all materials at 72h (Fig. 3b). No 261 statistical differences were found for iNOS expression. Regarding anti-inflammatory markers, 262 a significant increase of TGF-B was observed for 50M50G at 24 h, but after 72 h no 263 differences between materials were observed (Fig. 3d). The expression of IL-10 showed 264 differences at 72 h with a significantly higher expression on 50V50G (Fig. 3e). The 265 expression of EGR2 was significantly lower on 70M30T at 24 h compared to the other 266 materials and decreased at 72 h (Fig. 3f). 267

268 **3.4.** Cytokine quantification by ELISA

To obtain data about inflammatory induction by these materials, secretion profiles of pro- and 269 270 anti-inflammatory cytokines of RAW264.7 macrophages were assessed by ELISA (Fig. 4). RAW264.7 macrophages cultured on both 50M50G and 50V50G treatments showed a clear 271 increased secretion of TNF-a at 24 h compared to those cultured on the 70M30T and 272 35M35G30T materials. At 72 h, a marked high secretion of TNF- α for 50V50G was observed 273 (Fig. 4a). Further, an increasing IL-10 release was observed on this material, with significance 274 regarding the other materials (Fig. 4d). IL-1 β was not detected until 72 h of culture, revealing 275 no differences between materials. 276

278 **3.4.** Oxidative stress

Fig. 5 shows the macrophage oxidative stress markers (GSH, GSSG, GR, GPx and MDA)
when cultured on sol-gel materials. No significant differences were found between materials
at any time measured.

282 4. Discussion

Implanting a biomaterial foreign body into a living host leads to immediate tissue damage and cell disruption resulting from the surgical procedure. The blood protein adsorption onto the surface of the material causes platelet degranulation, forming a provisional matrix that kickstarts tissue healing responses, inducing immune cell activation and migration ⁽²⁶⁾.

The composition, conformation and amount of the bound proteins is regulated their specific 287 288 affinity and the biomaterial characteristics. Distinct biological responses can result by changing the surface and consequent protein adsorption; more specifically, emerging data 289 suggest that the modulation of immune cells is directly driven by complement protein 290 adsorption, affecting the *in vivo* biocompatibility of a material ⁽²⁷⁾. Immune cells interact 291 closely with complement proteins inducing an initial inflammatory response that propagates 292 depending on multiple factors and at implantation site activate and promote additional cellular 293 294 events.

Macrophages present a high plasticity and can adopt a wide battery of phenotypes. The M1 phenotype is characterized a pro-inflammatory response, the M2 phenotype presents antiinflammatory characteristics. At initial stages of inflammatory responses, the M1 is the most prevalent but, with time, macrophages undergo a transition to the M2 phenotype. However, the extent of the diversity of the M2 phenotype is not completely understood, and several M2 subtypes have been described (M2a, M2b, M2c, and M2d) ⁽²⁸⁾. These phenotypes attenuate acute and chronic inflammation through different mechanisms and signals ⁽²⁹⁾ even though 12

this classification still fails to cover the wide range of signals and functions related to M2 302 macrophages ⁽³⁰⁾. With a prolonged presence of a M1 phenotype on the local 303 microenvironment surrounding the material, fibrous structures can be observed ⁽⁵⁾. Thus, the 304 hypothesis that a biomaterial leading to the formation of connective tissue structures possibly 305 induces the differentiation of macrophages to a M1 phenotype arises. Previous work has 306 shown that the materials with low biocompatibility (50M50G and 50V50G) lead to the 307 formation of a fibrous capsule, while the materials with good biocompatibility (70M30T and 308 30M35G30T) did not present inflammatory structures. To understand these distinct in vivo 309 responses, protein adsorption of these two groups was compared. Results revealed higher 310 311 adsorption of inflammatory-related proteins onto the surfaces related to biocompatibility problems ⁽¹⁷⁾. The morphology acquired by macrophages when in contact with good 312 biocompatible materials cells displayed an elongated form, with cytoplasmic projections on 313 the apical edges, typical of M2-phenotype; on the other hand, on the materials with low 314 biocompatibility, the cells adopted an round shape, with very frail extensions of the 315 cytoplasm, characteristic of a M1 phenotype $^{(31),(32)}$. Furthermore, higher quantities of TNF- α 316 and IL-10 were secreted by the cells on the materials with low biocompatibility. This 317 increased release of TNF- α , a M1 marker ⁽³³⁾, is observed for cells cultured on both 50M50G 318 319 and 50V50G after short times of incubation (24 h). In addition, 50V50G showed this greater cytokine liberation even after 72 h, revealing a strong inflammatory potential with respect the 320 other treatments. The upregulated secretion of IL-10 on 50V50G, often considered a key M2 321 marker ⁽³³⁾, is dependent on the cell line ⁽³⁴⁾. In RAW264.7 cells exposed to LPS, IL-10 322 secretion is increased (35). As described in Araújo-Gomes et al. (27), GPTMS presents an epoxy 323 ring in its structure that might mimic LPS. However, IL-10 secretion was not significantly 324 higher on 50V50G. This might be due to the vinyl group of this formulation, as it was 325 described to induce inflammation in hepatic murine cells ⁽³⁶⁾. These results point out that IL-326

10 biomarker could lead to incorrect conclusions in murine cells as it is dependent on the 327 material chemistry. Interestingly, an overexpression of EGR2 was observed at 24 h on the 328 materials with low biocompatibility. The EGR2 is described to have a specific role on RAW 329 264.7 macrophage plasticity. Specifically, EGR2 is described to be expressed by non-330 activated and M2 macrophages, whereas it is downregulated in M1 macrophages ⁽³⁷⁾, being 331 modulated by the transcription factor CEBPB. Moreover, this gene is described as being a 332 "master controller" of inflammation by regulating B and T cell function to achieve immune 333 homeostasis ⁽³⁸⁾. We hypothesize that the greater expression of this gene during the first 24 h 334 on the GPTMS-based materials is due to the greater inflammatory induction, to regulate and 335 336 attenuate the inflammation caused by those specific materials. The immunocytochemistry supports the data obtained on by ELISA, disclosing higher tendency for the materials with 337 low biocompatibility to induce the RAW 264.7 to differentiate toward a pro-inflammatory M1 338 phenotype. This distinct polarization points out to the increased inflammatory potential of the 339 50M50G and 50V50G coatings, which is coherent with the data obtained in a previous study 340 and could explain the dissimilar biocompatibility associated with each of these materials ⁽¹⁷⁾. 341 However, it appears that 35M35G30T is also inducing an M1 phenotype compared to the 342 70M30T coating. This fact can be associated with the 35 % of GPTMS incorporated in the 343 344 coating network. GPTMS-derived sol-gel materials showed an increased inflammatory potential, which in turn was directly correlated with a higher affinity of complement proteins 345 to the material surface ⁽²⁷⁾. However, when comparing to 50M50G and 50V50G, we can 346 conclude that this may be due to the lower percentage of the compound, therefore not 347 compromising biocompatibility. 348

Although this data seems to identify clear and distinct cellular behavior when exposed to the materials, these differences were not be translated into the oxidative stress induction. Data obtained from oxidative stress measurements showed no differences between materials, suggesting once more that the inflammation is driven by the complement protein attachment,
consequent cytokine liberation and immune cell activation, and the materials do not represent
immediate harm for the cell and/or induce oxidative stress.

355 **5.** Conclusion

The aim of this study was to evaluate how sol-gel coatings with distinct *in vivo* outcomes modulate oxidative stress and inflammatory responses. Although there was no differences in oxidative stress, coatings with low biocompatibility (50M50G and 50V50G) had proinflammatory profiles with higher secretion of TNF- α . Moreover, these materials showed a higher expression of M1 receptors (IL7-R); however, the expression of M2 receptors (CD206) was not significantly different, indicating that M1 and M2 balance is key to define inflammatory responses to a biomaterial.

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493 Figures



Figure 1. Cell morphological analysis by SEM. Sample microphotographs of RAW 264.7
cultured on (a-a') 70M30T, (b-b') 35M35G30T, (c-c') 50M50G, (d-d') and 50V50G sol-gel
hybrid coatings after 72h. The experiment was carried out with two replicates. Scale bar: 10
µm and 4 µm.



Figure 2. Immunostaining of RAW264.7 cells cultured on (a-a'') 70M30T, (b-b'') 35M35G30T, (c-c'') 50M50G, and (d-d'') 50V50G sol-gel hybrid coatings, after 72h. IL7-R (a'-d') was used as a M1 marker and CD206 (a''-d'') was used as a M2 marker. The relative corrected total cell fluorescence (CTCF) of these markers (e and f) was quantified using ImageJ. The experiment was carried out with three replicates. Data are presented as mean \pm SD. The asterisk (*) indicates differences between materials (p < 0.05). Scale bar: 100µm.

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Figure 3. Gene expression of RAW264.7 cells cultured on 70M30T, 35M35G30T, 50M50G and 50V50G on the sol-gel hybrid coatings after 24 and 72h: (a) TNF- α (a), (b) IL-1 β , (c) iNOS, (d) TGF- β , (e) IL10, and (f) EGR2. The experiment was carried out with six replicates. Data were normalized to blank wells (without material) and are presented as mean ± SD. The asterisk (*) indicates differences between materials (p < 0.05).



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Figure 4. Cytokine secretion of RAW264.7 cells cultured on 70M30T, 35M35G30T, 50M50G and 50V50G on the sol-gel hybrid coatings after 24 and 72h: (a) TNF- α , (b) IL1- β , (c) TGF- β , and (d) IL-10. The experiment was carried out with four replicates. Data are presented as mean \pm SD. The asterisk (*) indicates differences between materials (p < 0.05).



518 Figure 5. Oxidative stress markers of RAW264.7 cells cultured on 70M30T, 35M35G30T,

519 50M50G and 50V50G on the sol-gel hybrid coatings after 24, 72 and 168h: (a) GSH, (b)

520 GSSG, (c) GR, (d) GPx, (e) MDA, (f) GSH/GSSG. The experiment was carried out with four

521 replicates. Results are shown as mean \pm SD.