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Hydroxyl radicals and oxidative stress: the dark side of Fe corrosion

This is the author's manuscript
Original Citation:
Availability:
This version is available http://hdl.handle.net/2318/1732441 since 2023-03-01T13:45:00Z
Published version:
DOI:10.1016/j.colsurfb.2019.110542
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1	Reactive Oxygen Species: the hidden face of
2	biodegradable Fe-based alloys
3	E. Scarcello ¹ , M. Tomatis ² , F. Turci ² , A. Thomas ³ , P.J. Jacques ³ , D. Lison ¹
4 5	¹ Louvain Centre for Toxicology and Applied Pharmacology (LTAP), Université catholique de Louvain, Avenue E. Mounier 53–bte B1.52.12, 1200 Brussels, Belgium.
6 7	² "G. Scansetti" Interdepartmental Center for Studies on Asbestos and Other Toxic Particulates, University of Torino, Via P. Giuria 9, 10125 Turin, Italy.
8 9	³ Institute of Mechanics, Materials and Civil Engineering (IMAP), Université catholique de Louvain Place St Barbe 2, 1348 Louvain-la-Neuve, Belgium.

10 Abstract

The present work documents, for the first time, the ability of biodegradable Fe-based
 materials to produce hydroxyl radicals (OH•) during local corrosion and its cellular impact
 on endothelial cells.

The ability of Fe-based materials to generate OH• was documented by two complementary but independent acellular tests, i.e. terephthalate (TA) hydroxylation fluorescence and electron paramagnetic resonance (EPR) spectroscopy. The cellular responses were assessed *in vitro* on HUVECs and HAOECs using colorimetry and luminescence cytotoxicity assays. Cells were exposed directly to Fe powder, or to corrosion extracts. To confirm the cellular impact of OH•, mRNA expression of oxidative stress response genes (*HO-1* and *hGCLM*) was assessed in endothelial cells directly exposed to the same particles or corrosion extracts.

21 All tested Fe-based materials showed a strong potential to generate OH•, as a result of 22 incomplete reduction of dissolved oxygen or via Fenton chemistry. The reduction of these 23 signals in presence of D-mannitol confirmed their specificity for OH•. Only direct contact 24 with Fe materials affected cell viability, indicating that released ions in corrosion extracts 25 do not contribute to the cytotoxic activity. The expression of oxidative stress genes was 26 dose-dependently increased 4 h after direct exposure to the particles, not to released ions. 27 Pre-treatment with cytochalasin-D reduced cytotoxic and oxidative stress gene expression, 28 indicating that endocytosis contributes to direct cell responses to Fe particles.

The demonstration of OH• production during corrosion and consequent oxidative stress in
 endothelial cells provides a new perspective on the biocompatibility of biodegradable Fe based alloys. These findings will influence the future design of Fe-based implants,
 especially in vascular walls.

33 Keywords: oxidative stress, hydroxyl radicals, endothelial cells, coronary stent.

34 **1. Introduction**

35 During the last decade, biodegradable metallic alloys have been developed and investigated 36 as alternatives for permanent implants, notably cardiovascular stents [1]. The most 37 favorable and suitable coronary stent should, indeed, maintain its mechanical integrity only 38 for the first 6–12 months, and be totally degraded after 12–24 months to avoid long-term 39 restenosis or late thrombosis [2]. An ideal biodegradable material for coronary stent should, 40 therefore, demonstrate a perfect compromise between degradation and mechanical 41 performances. The first biodegradable biomaterials proposed were polymers from lactic 42 acid, glycolic acid or caprolactone families but, while their biocompatibility and 43 degradation rate appear adequate, their intrinsic mechanical properties are rather poor [3]. 44 A more recent idea is to consider metals as biodegradable materials. Biodegradable 45 magnesium-based alloys were first investigated for their better mechanical properties, but 46 their specific strength remains weak compared to current permanent materials such as 47 austenitic stainless steels (316L) [4]. The suitability of Fe as a biodegradable implant 48 material has also been investigated. Fe combines a high strength, a high elastic modulus and 49 a high ductility which can be helpful for the implantation of a stent when it needs to be 50 plastically deformed. The first biodegradable Fe-based stent was made of Armco[®] iron (Fe 51 > 99.8%) and implanted in the descending aorta of New Zealand white rabbits. No 52 indication of local or systemic toxicity was detected, but the stents did not corrode 53 completely over the 18 months follow-up period [5]. The addition of Mn to Fe may improve 54 its performances, leading to excellent mechanical properties similar to 316L, combined with 55 a degradation rate higher than pure Fe and tunable by varying the Mn content [3]. These 56 degradation rates remain, however, one order of magnitude lower than those of Mg alloys. 57 Other Fe-based alloys, including the austenitic Fe-Mn-C-Pd or TWIP (twinning-induced 58 plasticity) steel alloys have been considered [6].

59 Immediately after implantation of a metallic stent in the artery, oxygen dissolved in the 60 blood drives metal corrosion, generating free metal ions and associated degradation products. This process is complex as redox reactions generate not only ions, such as Fe²⁺ 61 62 on the anodic side, but may also contribute to the formation of reactive oxygen species 63 (ROS). While metal ions are released during corrosion, dissolved oxygen is concomitantly 64 reduced in water (4 electrons) or, when the process is incomplete, in ROS such as OH• (3 65 electrons). OH• is the most damaging ROS which highly reacts with all biological macromolecules. It is one of the most potent oxidizing agents and has significant pro-66 inflammatory properties [7]. OH• can also be formed after Fe corrosion through a Fenton 67 68 reaction in the presence of H_2O_2 present in the wounded arterial lumen [8].

Previous investigations on the biocompatibility of metallic materials often overlooked the
complexity of these biodegradation phenomena and mostly focused on soluble ionic
metallic forms released from the biocorrosion process [3, 6, 9, 10]. Most authors have so
far concluded, based mostly on indirect cytotoxicity tests according to ISO standard 109935, that Fe-based alloys could be considered as nontoxic and biocompatible materials.
Assuming that toxicity is exclusively driven by the solubilized metal ions represents,
however, an over-simplification, and testing the response to ionic constituents or leachates

76 may not provide an appropriate approach. Specifically for Fe-based materials, the possible 77 role of ROS generated by dissolved oxygen deserves a specific attention in view of the 78 capacity of this element to contribute to catalytic redox cycling such as the well-known 79 Haber-Weiss reactions. For coronary stents, the possible implication of ROS appears very 80 relevant because of the critical role of oxidative stress in the atheromatous tissue. In 81 addition, balloon angioplasty and coronary stent implantation are associated with increased 82 vascular levels of ROS in conjunction with altered endothelial cell and smooth muscle cell 83 function [11]. Because corrosion is expected to occur during the whole life time of the 84 implant, ROS would be continuously formed at the implant surface, possibly resulting into 85 prolonged inflammation and unsuccessful healing of the surrounding tissues. Consequently, 86 understanding cellular reactions to implant-induced oxidative stress and inflammatory 87 activation is important to help prevent adverse responses to metallic stents [12].

88 While it is well established, in the field of inhalation toxicology, that metal particles can 89 release ROS, see e.g. [13], it is surprising that this aspect has not been well explored in the

field of biodegradable materials used for medical devices [14]. In this work we hypothesized
that OH• accompanying biocorrosion of Fe-based material are a source of toxicity for
endothelial cells. We specifically focused on the formation of OH• which appear the most
probable and deleterious ROS during the corrosion of Fe-based materials.

94 **2. Materials & Methods**

95 **2.1 Metallic materials**

96 Carbonyl iron powder (#44890, purity \geq 99.5%) and iron chips (#267945, purity 99.98%) 97 were purchased from Sigma-Aldrich (St Louis, MO). Carbonyl iron particles were 5-9 µm 98 in spherical size (see SEM images, Supplementary data, Fig. S1). Other material sheets 99 were produced by melting iron (Alfa Aesar, 99.99%) or TWIP (FeMnC) in an arc furnace. 100 After a treatment of 10 minutes at 1000°C, the cast ingots were hot rolled to a thickness 101 lower than 2 mm. An Accutom[®] 5 automatic cut-off machine (Struer[®], Ballerup, Denmark) 102 was used to cut samples into 10x10 mm. Both sides of the samples were polished with SiC 103 water-proof paper 320 grits with water and then rinsed with 99.8 vol% absolute ethanol. 104 The specimens were finally ultrasonically washed in absolute ethanol for 10 min 105 immediately before ROS-generation tests. Micrometric crystalline silica particles were 106 Min-U-Sil[®] 5 (Berkeley Springs, West Virginia).





109	Figure \$1. Scanning Electron Microscope (SEM) images of carbonyl iron powde
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(Sigma-Aldrich®). The images were recorded using the SEM Ultra® 55 Zeiss at
 different magnitude.

112 **2.2 Electron paramagnetic resonance / spin trapping**

The ability of the different samples to generate OH• was monitored by electron paramagnetic resonance (EPR) spectroscopy with a Miniscope MS 100 (Magnettech, Berlin, Germany) EPR spectrometer using DMPO as spin-trapping agent. The instrument settings were as follows: modulation 1000 mG, scan range 120 G, center of field approximately 3355 G.

118 2.2.1 Carbon-centred free radicals detection during metal corrosion

25 mg of the powder or a square massive sample (ARMCO[®] / Pure iron / TWIP) were 119 120 suspended in 1 ml of PBS (0.5 M, pH 7.4, Sigma-Aldrich, Milan, Italy) of sodium formate 121 (1.0 M, Sigma-Aldrich, Milan, Italy) and DMPO (0.04 M, Cayman Chemical Company, 122 Ann Arbor, Michigan). After 10, 30 and 60 min of incubation under continuous stirring, 123 aliquots of 50 µl were withdrawn, filtered (pore diameter 0.22 µm, Merck Chemicals SA, 124 Tullagreen, Carrigtwohill, Co. Cork, Ireland) and EPR spectra were recorded at room 125 temperature. Blanks were run in parallel in the absence of sample. Iron(II, III) oxide was 126 used a positive control [15]. All the measurements were performed in triplicate.

127 2.2.2 Surface-driven Fenton reactivity (target molecule H₂O₂)

128 25 mg of the powder or a square sample (ARMCO[®] / Pure iron / TWIP) were suspended in 129 1 ml of a phosphate-buffered saline (PBS 0.5 M, pH 7.4) containing DMPO (0.04 M) and 130 H_2O_2 (0.1 M). After 10, 30 and 60 min of incubation under continuous stirring, aliquots of 131 50 µl were withdrawn, filtered (0.22 µm) and EPR spectra were recorded at room 132 temperature. Blanks were run in parallel in the absence of sample. All the measurements 133 were performed in triplicate.

134 **2.3 Fluorimetric determination of hydroxyl radicals**

135 Disodium terephthalate (99%, Alfa Aesar, Tewksbury, MA) was dissolved in 0.01 M PBS 136 (pH 7.4) at a final concentration of 10 mM [16]. Iron powder or square samples were 137 suspended in TA solution (30 mg dust or a metal square piece / ml) and incubated for 30 138 min at 25°C under continuous stirring. In some assays, the OH• scavenger D-mannitol (75 139 mM, Sigma-Aldrich, St Louis, MO) was added to the reaction mixture. After incubation, the suspensions were filtered on cellulose acetate membrane (0.22 μ m) and the fluorescence 140 141 of the filtrate was measured on a SpectraMax i3x Multi-Mode microplate reader with an 142 excitation light $\lambda_{ex} = 324$ nm and reading the maximum emission intensity at $\lambda_{em} =$ 143 425 nm.

144 **2.4 Controls in an oxygen-free environment**

145The absence of OH• generation in an oxygen-free environment was verified for both146procedures described above (EPR and TA). In these assays, the aqueous buffered solution147containing DMPO or TA were deoxygenated by energetic bubbling of N2 for 30 min before148adding samples and throughout the duration of the test.

149**2.5 Cell culture**

Cell studies were performed with HUVEC and HAOEC endothelial cell lines (Cell Application, Sigma-Aldrich, San Diego, CA) because endothelial cells are in direct contact with the stent surface immediately after implantation [6]. HUVECs have been chosen because they are arguably the most well-characterized primary human EC type [17] and HAOECs as a stand-in for the arterial tunica intima [18]. The cells were cultured on 0.2% gelatin-coated 75-cm² culture flasks (Corning Incorporated, Corning, NY) in Endothelial
Cell Growth Medium (Cell Application, Sigma-Aldrich, San Diego, CA) and maintained in
a humidified incubator (New Brunswick Galaxy[®] 170S) containing 5% CO₂ at 37°C. Cells
were grown to confluency and harvested by trypsinization. A maximum of 7 and 5 passages
was used for HUVECs and HAOECs, respectively, to maintain phenotypic characteristics
of endothelial cells. Cells were also routinely tested for the absence of Mycoplasma
infection (PCR Mycoplasma Test Kit I/RT, PromoKine, Huissen, the Netherlands).

162 **2.6 Direct cell viability assays**

163 The day of cell exposure, Fe particles were heated at 200°C (WTB, Binder[®] drving oven) for 2 h for sterilization and destruction of any possible trace of endotoxin, and suspended in 164 165 cell culture medium at a stock concentration of 5 mg/ml. Immediately before cell exposure, 166 each particle suspension was diluted to final concentrations and vortexed. Direct cell 167 viability was assessed with the colorimetric WST-1 assay from Roche Diagnostics GmbH (Mannheim, Germany) and the luminescent CellTiter-Glo[®] 2.0 assay from Promega Corp. 168 169 (Madison, WI). In the WST-1 assay, the amount of formazan dye formed is directly related 170 to the number of metabolically active cells. The assay was carried out as described in the 171 manufacturer's instruction. In brief, cells (2.10⁴ HUVECs or HAOECs) were seeded in 96-172 well transparent plates and exposed the day after to different concentrations of iron powder 173 for 24 h. Cells were washed and incubated in fresh medium with 10% WST-1 reagent for 174 2h. Absorbance was measured at 450 nm, with 690 nm as reference, in a multiplate reader 175 (Infinite F200, Tecan[®]). Results are reported as relative WST-1 activity, where 1.0 176 corresponds to the absorbance measured in untreated control cultures. The CellTiter-Glo® 177 2.0 assay determines the number of viable cells in culture by quantitating the amount of 178 ATP which indicates the presence of metabolically active cells. In brief, cells (2.10^4) 179 HUVECs or HAOECs) were seeded in 96-well white plates and exposed as above. After 180 24h, the CellTiter-Glo reagent was added. Luminescence was read on a luminometer 181 (Victor[™] X4, PerkinElmer[®]). Results are reported as for WST-1.

For endocytosis inhibition studies, endothelial cells were treated with cytochalasin-D (2.5 μM, Sigma–Aldrich, St Louis, MO) 30 min before particle exposure. Min-U-Sil[®] 5 or Co (II, III) oxide (Aldrich, St Luis, MO) were used as positive controls.

185 **2.7 Indirect cell viability assays**

186 HUVECs or HAOECs were also used to assess the in vitro cytotoxicity of corrosion extracts 187 generated after the immersion of increasing concentrations of iron powder in cell culture 188 medium for 24 h. The concentration of released Fe ions was quantified by inductively 189 coupled plasma - mass spectroscopy (ICP-MS) after filtration of the suspension (Amicon® 190 5000 NMWL, Millipore Corporation, Bedford). Endothelial cells were seeded in a 96-well 191 plate with a cell density of 2.10⁴ per well. After 24 h of incubation, the cell medium was 192 replaced by corrosion extracts and cells were further incubated for 24 h. The mitochondrial 193 activity or the ATP amount in viable cells were measured as described for the direct contact 194 test.

Indirect contact testing was also performed using a 96-well Transwell[®] chamber with 0.4
µm pore polycarbonate membrane insert (Corning[®] HTS, Sigma-Aldrich, St Luis, MO).
Cells were seeded in the lower chamber and allowed to adhere. After 24 h, increasing
concentrations of iron powder were added in the upper chamber. In this condition, only
degradation products released from the dust could reach cell culture at the bottom of the
well.

201 **2.8 Gene expression experiments**

202 Transcripts of oxidative stress response genes HO-1 and hGCLM were measured in 203 HUVECs or HAOECs exposed to increasing concentrations of iron powder or corrosion 204 extracts for 4 h. Cells were washed twice and RNA was extracted with the TriPure Isolation 205 Reagent (Roche, Mannheim, Germany) according to the manufacturer's protocol, followed 206 by DNase treatment (Invitrogen Inc., Camarillo, CA). A quantity between 10 ng and 5 µg 207 of RNA was reverse transcribed by M-MLV-Reverse Transcriptase (Invitrogen) with 700 208 pmol/µl random hexamers (Eurogentec, Seraing, Belgium) in a final volume of 25 µl. The resulting complementary DNA was then diluted 10-fold in sterile UltraPure® water 209 210 (Invitrogen) and used as template in subsequent real-time polymerase chain reactions 211 (PCR). Five microliters of diluted cDNA or standards were amplified using SYBR Green technology in a total volume of 20 µl on a StepOnePlus[™] Real-Time PCR System Thermal 212 213 Cycling Block (Applied Biosystems, Foster City, CA) according to the following program: 10 min 95°C and 40 cycles of (15 s 95°C + 1 min 60°C). After amplification, a melting 214 215 curve was generated and data analysis was performed with the StepOne[™] Software v2.3 216 (Applied Biosystems). Primers for HO-1, hGCLM and β -actin were purchased from 217 Invitrogen Inc.

218	β -actin:	(sense) 5' CCCGTGCTGCTGACCG G 3'
219		(antisense) 5' CGTCACCGGAGTCCATCAC 3';
220	<i>HO-1</i> :	(sense) 5' GCAACAAAGTGCAAGATTCTGC 3'
221		(antisense) 5' GCTGTAGGGCTTTATGCCATGT 3';
222	hGCLM:	(sense) 5' CAGCCTTACTGGGAGGAATTAGAA 3'
223		(antisense) 5' TTACTATTTGGTTTTACCTGTGCCC 3'.

224 Results were calculated as a ratio of *HO-1* or *hGCLM* expression to the expression of the 225 reference gene, β -actin.

226 **2.9** Antioxidants

Endothelial cells were treated with D-mannitol (125 mM), D-sorbitol (1.5 mM), sodium formate (10 mM), N-acetyl-L-cysteine (NAC, 5 μ M), uric acid (1 mM), ascorbic acid (250 μ g/ml), 2,6-di-tert-butyl-4-methylphenol (BHT, 0.01 mM), trolox (5 μ M) or catalase (250 units/ml) immediately prior to iron powder exposure as described in section <u>2.6</u>. All antioxidants were purchased from Sigma-Aldrich (St Louis, MO).

232 2.10 Statistical analysis

Data are presented as means ± standard error of the mean (S.E.M.) of independent experiments (N) conducted in replicates (n). The data were analyzed with GraphPad Prism (GraphPad software, La Jolla, CA) and OriginPro9.0 software (OriginLab Corp., Northampton, MA). Differences between groups were analyzed by one-way analysis of variance (ANOVA) followed by a post-hoc Dunnett's pairwise comparison test. Differences with p value < 0.05 compared to control group were considered statistically significant.

3. Results

241 **3.1 Fe-based materials generate hydroxyl radicals**

242 Our first goal was to determine the ability of Fe-based materials to release OH• in aqueous 243 conditions. The formation of OH• during the corrosion of Fe-based materials was assessed 244 by monitoring the formation of carbon-centered free radicals in the presence of formate 245 [19]. The cleavage of a C–H bond was detected by spin trapping and quantified by EPR spectroscopy. Representative EPR spectra of the [DMPO-CO₂]-adducts indicated that iron 246 powder (Fig. 1b), pure iron piece (Fig. 1c), pure iron chips (Fig. 1d), ARMCO[®] piece (Fig. 247 248 1e) and TWIP piece (Fig. 1f) strongly generated carboxyl radicals (•CO2–) in solution. No 249 signal was detected in the absence of Fe material (blank) (Fig. 1a). The strong reactivity of 250 Fe-based materials was maintained over time, up to 60 min.



251 252	Figure 1. Surface driven release of hydroxyl radicals from the corrosion of Fe-based materials. EPR spectra recorded on suspensions of iron powder (75 mg, b), pure iron
253 254 255 256 257 258 259 260 _	piece ($10x10x1.5$ mm, c), pure iron chips (670.6 mg, d), ARMCO [®] piece ($10x10x1.5$ mm, e) and TWIP piece ($10x10x1.5$ mm, f) compared to blank (a) in a sodium formate- buffered solution in the presence of DMPO as spin-trapping agent. Aliquots of 50 µl of suspension were withdrawn after 10, 30 and 60 min of incubation under continuous stirring at room temperature, filtered and analysed for EPR spectra. The panel shows a representative spectrum out of at least three experiments. x 1,5 means one and a half EPR signal intensity.

261 In order to support the EPR results, we applied a complementary test using a different 262 readout, i.e. the capacity of OH• to hydroxylate TA. Compared to the blank (Fig. 2), all Fe-263 based samples were able to hydroxylate TA, regardless of weight, form or Fe content. The 264 specificity of the TA measurement was verified by adding D-mannitol, an OH• scavenger 265 frequently used in biological systems [20]. D-mannitol significantly reduced the signal, 266 confirming the generation of OH• during Fe-based materials corrosion.





determination of hydroxyl radicals released from Fe-based materials using the terephthalate (TA) assay. Samples were immersed in a buffered (PBS) solution of disodium terephtalate (10 mM) for 30 min at RT under continuous stirring in the absence/presence of D-mannitol, an hydroxyl radical scavenger (75 mM). Supernatant was recovered, filtered and the fluorescence was measured (excitation $\lambda_{ex} = 324 nm$, emission $\lambda_{em} = 425 nm$). Data are means ± SEM (N=4, n=3).

275 To confirm that dissolved oxygen drives the generation of OH•, spin trapping and 276 fluorimetric measurements were performed in an oxygen-free environment. The absence of 277 oxygen completely suppressed the corrosion mechanism and the generation of OH• 278 (Supplementary data, Fig. S2).



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Figure S2. Generation of hydroxyl radicals by Fe-based materials is oxygendependent. EPR spectra recorded on suspensions of iron powder (75 mg, **a**), pure iron chips (632 mg, **b**), ARMCO[®] piece (10x10x1.5 mm, **c**) and TWIP piece (10x10x1.5 mm, **d**) in a aqueous buffered solution in the presence of DMPO as spin-trapping agent. The solution was previously deoxygenated by vigorously bubbling N₂ for 30 min; the oxygen-free environment was maintained for the duration of the test. Aliquots of 50 µl of suspension were withdrawn after 10, 30 and 60 min of incubation under continuous stirring at room temperature, filtered and analysed for EPR spectra.

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When Fe cations are generated in the presence of hydrogen peroxide, additional reactions known as the Fenton reaction (Eq.¹) can take place, also leading to the generation of OH• :

$$293 \qquad Fe^{2+} + H_2O_2 \rightarrow Fe^{3+} + OH \cdot + OH^-$$

294Representative EPR spectra of [DMPO-HO]-adduct recorded in the presence of H_2O_2 are295reported (Fig. **3b-d**). No signal was detected in the absence of Fe material (Fig. **3a**). A time-296dependent increase in the intensity of the EPR signal was observed with all tested samples.



298 Figure 3. Fenton-mediated generation of hydroxyl radicals during corrosion of Fe-299 based materials. EPR spectra recorded on suspensions of iron powder (75 mg, b), pure iron piece (10x10x1.5 mm, c), pure iron chips (2.19 g, d), ARMCO[®] piece 300 (10x10x1.5 mm, e) and TWIP piece (10x10x1.5 mm, f) compared to blank (a) in a 301 302 aqueous buffered solution in the presence of H_2O_2 (0,1M) and DMPO as spin-trapping 303 agent. Aliquots of 50 µl of suspension were withdrawn after 10, 30 and 60 min of 304 incubation under continuous stirring at room temperature, filtered and analysed for 305 EPR spectra.

306 The panel shows a representative spectrum out of at least three experiments.

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Alternative reactions could also lead to radical adduct artifacts such as nucleophilic addition of water at the nitrone carbon (or C-2 position) of DMPO in the presence of Fe(III) ions [21]. Hence, we recorded the [DMPO–HO]–adduct without H₂O₂ and no signal was detected, demonstrating that, under our experimental conditions, DMPO exclusively forms
 the radical adduct by trapping the OH• (data not shown). We, therefore, concluded that the
 corrosion of Fe-based materials generates OH•.

314 **3.2 Direct cellular contact with Fe powder induces endothelial responses**

315 We next used Fe powder to assess the responses of endothelial cells. We reasoned that if 316 OH• are generated, they need, due to their short half-life, an intimate contact with cells to 317 exert a cytotoxic activity. We, therefore, compared the response to Fe powder in direct and 318 indirect cytotoxicity assays, hypothesizing that only the direct assay could capture the 319 activity of OH•. Experiments were carried out in two cell lines, HUVECs and HAOECs, 320 using two cytotoxicity tests based on different principles, i.e. colorimetry and luminescence. 321 Both assays showed that only direct contact with Fe powder significantly and dose-322 dependently affected the viability of endothelial cells after 24 h exposure. In contrast, 323 indirect assays (corrosion extracts or Transwell exposure) did not reveal a cytotoxic activity 324 of the Fe powder (Fig. 4a, b, c).



325 Figure 4. Direct exposure to iron powder affects endothelial cell viability. Cells (2.104 326 HUVECs or HAOECs) were seeded in 96-well transparent or white plates and exposed 327 the day after to different concentrations of iron powder or corrosion extracts for 24 h. 328 Extracts were obtained from culture medium incubated during 24 h with increasing 329 concentrations of iron powder. Supernatants were collected and centrifuged. The 330 chemical concentration of released iron ions was quantified by inductively coupled 331 plasma - mass spectroscopy (ICP-MS) after filtration of the suspension (d). In some experiments a 96-well Transwell[®] chamber with 0.4 µm pore polycarbonate 332 333 membrane insert was used, where cells were seeded in the lower chamber allowing 334 to adhere for 24 h and increasing concentrations of iron powder were added in the 335 upper chamber (c). The cells were washed and incubated in fresh medium with 10% 336 WST-1 reagent for 2 h. Absorbance was measured at 450 nm, with 690 nm as 337 reference, in a multiplate reader (a.1, b.1, c). The white plate was replenished with fresh medium with the CellTiter-Glo[®] reagent and luminescence was read on a 338 339 luminometer (a.2, b.2). Results are reported as relative WST-1 activity or luminescence

To further support the response of endothelial cells to OH• when in contact with Fe powder, we documented the oxidative stress response by monitoring the expression of heme oxygenase-1 (HO-1). HO-1 was increased dose-dependently 4 h after direct exposure, whereas corrosion extract exposure did not induce such a response (Fig. 5). Similar results were obtained for another oxidative stress marker, human glutamate cysteine ligase modifier subunit (*hGCLM*, Supplementary data, Fig. S3). These results supported the concept that OH• induce oxidative stress responses in endothelial cells.





Figure 5. Corrosion of iron powder induces oxidative stress in EC. HUVECs (a) or HAOECs (b) were exposed to different concentrations of iron powder or corrosion extracts for 4 h. RNA was extracted and reverse transcribed for real-time polymerase chain reactions (PCR). Values of heme oxygenase-1 (HO-1) mRNA were normalized to β -actin amplified from the same samples and are presented as fold increase compared to untreated cells. Data are means ± SEM, * p < 0.05 (N=2, n=4).

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We tried to protect endothelial cells from oxidative stress with antioxidants as described in section <u>2.9</u>. No evidence of HUVECs or HAOECs protection from OH• was recorded, as assessed by cytotoxicity or oxidative stress genes induction.

To clarify the mechanism of their response to Fe particles, endothelial cells were pre-treated with cytochalasin-D, an inhibitor of actin polymerization that blocks >90% of endocytosis [22]. When cells where pre-treated with cytochalasin-D no cytotoxic or oxidative stress gene expression was observed (Fig. **6a, b**).



Figure S3. Corrosion of iron powder induces oxidative stress in EC. HUVECs (a) or HAOECs (b) were exposed to different concentrations of iron powder or corrosion extracts for 4 h. RNA was extracted and reverse transcribed for real-time polymerase chain reactions (PCR). Values of glutamate cysteine ligase modifier subunit (hGCLM) mRNA were normalized to β -actin amplified from the same samples and are presented as fold increase compared to untreated cells. Data are means ± SEM, * p < 0.05 (N=1, n=4).











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381 382 383 384 385 386 387	Figure 6. Only iron powder endocytosis induced endothelial cytotoxicity. Cells (2.10^4 HUVECs or HAOECs) were seeded in 96-well transparent or white plates and exposed the day after to different concentrations of iron powder or corrosion extracts with or without pre-exposure with cytochalasin-D (2.5μ M, 30 min before particle exposure). After 24 h, the cells were washed and incubated in fresh medium with 10% WST-1 reagent for 2 h. Absorbance was measured at 450 nm, with 690 nm as reference, in a multiplate reader (a.1, 3). The white plate was replenished with fresh medium with
388 389	the CellTiter-Glo [®] reagent and luminescence was read on a luminometer (a.2, 4). Results are reported as relative WST-1 activity or luminescence (R.L.U.), where 1.0
390 391 392 393	corresponds to the value measured in untreated control cultures. Min-U-Sil [®] 5 and Co(II, III) oxide were used as positive controls. Data are means \pm SEM for at least three samples (N=4), * p < 0.05.

395 **4. Discussion**

396 We documented, for the first time, the ability of biodegradable Fe-based materials to 397 produce OH• during corrosion. Metallic materials that undergo corrosion, release 398 degradation products at the implanted site thus constantly causing the formation of ROS 399 [14], but no previous study had been carried out in order to evaluate the OH•-producing 400 activity of biodegradable Fe-based alloys. This finding is important because it sheds a new 401 light on the biocompatibility of Fe-based alloys investigated as implant. These have been 402 generally considered as having good biocompatibility properties mostly based on indirect 403 contact cellular tests [3, 6, 9, 23, 24]. These assays can, however, not capture the cytotoxic 404 activity of the short-lived oxygen species.

- To cover a wide range of implanted materials used in medicine and their different surface reactivity, we tested several samples of Fe-based materials differing in shape, size, surface area, roughness and composition. All samples examined here showed a strong potential to generate OH• in acellular systems, via EPR and TA hydroxylation assays, two complementary techniques [25]. Any quantitative comparison of OH• yields must, however, be considered carefully because the samples were not tested under similar conditions of dose, surface area, geometry and Fe content.
- 412 Implant corrosion releases degradation products that can be found in various forms, 413 including free metallic ions, colloidal complexes, inorganic metal salts or oxides and wear 414 particles [14]. Particle size and shape change with the passage of time and corrosion 415 particles can further contribute to corrosion, as the surface in contact with the surrounding 416 fluids becomes larger [26]. Incomplete reduction of dissolved oxygen could produce ROS, 417 such as OH•, on material surface that will immediately react with almost every molecule in 418 its environment. It can also be expected that local corrosion of Fe implants releases metal ions which undergo the Fenton reaction in presence of H₂O₂, also present in the wounded 419 420 area. As biocorrosion persists through the life of the implant, OH• are likely to be 421 continuously formed at the implant site, worsening the local oxidative stress levels of the 422 diseased tissue. This could influence the success of the device implantation and the healing 423 of the surrounding tissues [27]. Small debris as result of metal degradation migrate in the 424 tissue surrounding the material and are immediately phagocytosed, whereas metal 425 nanoparticles can pass through the cell plasma membrane mainly by diffusion or 426 endocytosis [28]. Hence, ROS derived from the material may cause intracellular oxidative 427 damage, including to the nucleus proteins and lipids, resulting in inhibition of DNA repair 428 pathways, impair of nuclear signal transduction and defective gene expression [29]. We 429 found that endocytosed Fe particles directly affected endothelial cells viability probably via 430 intra-cellular corrosion and OH• production. This toxicity mechanism could explain that 431 antioxidants, able to scavenge OH• in a cell free environment were unable to protect 432 endothelial cells. Indeed, OH• would be mostly produced in a microenvironment poorly 433 accessible to water-soluble antioxidants, possibly after endocytosis. Moreover, during the 434 24 h cell exposure, antioxidant molecules may have reacted with cell culture medium 435 components and lead to different forms of antioxidant, or simply degrade, whereas iron 436 particles are immediately cell-internalized. It is possible that using a massive sample (e.g. 437 cube or disk [30]) instead of our model particles, may lead to different results, because the 438 degradation rate of the bulk material during a cell culture test is expected to cause less 439 particle formation and consequent endocytosis.
- 440 Overall, acellular results showed that dissolved oxygen drives the corrosion of Fe-based 441 materials and generates OH• by two different ways, directly on the metal surface or through 442 Fe ions released that undergo a Fenton reaction, as shown in Fig. 7. Direct OH• release 443 emerges as the most significant to induce cell toxicity or oxidative stress as exposure to 444 extracts, that include iron ions able to react with the H₂O₂, did not affect cell responses.



Figure 7 Summary of mechanism of OH• from the corrosion of Fe-based materials.
OH• are directly generated from the incomplete reduction of dissolved oxygen
(surface driven release, in red) or mediated by Fe ions that react with H₂O₂ (Fentonmediated generation, in yellow).

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452 The mechanisms highlighted here can act as a first approximation of the biodegradation of 453 a Fe-based implant and its impact on surrounding tissues. These results represent a 454 particularly interesting aspect because, until now, little was known about biomaterials and 455 oxidative stress, and the mechanism underlying metal toxicity is still not fully understood 456 [14]. The present data suggest that some of the classic experiments used for evaluating *in* 457 vitro biomaterials toxicity, such as indirect contact test [3, 6, 9, 23, 24, 31], are not 458 appropriate to capture the impact of implanted biodegradable Fe-based materials on 459 surrounding tissue. In the light of our results, assuming that toxicity is exclusively driven 460 by the solubilized metal ions represents an over-simplification, missing the contribution of 461 the short-lived ROS. In addition, our data indicate that oxidative stress might contribute to 462 the toxicity of biodegradable Fe-based materials and this aspect appears particularly 463 relevant to take into account for an atheromatous tissue already submitted to oxidative stress 464 [32].

This work was supported by an ARC grant (principal promoter Pascal J. JACQUES).

465 **5. Acknowledgments**

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468 **References**

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