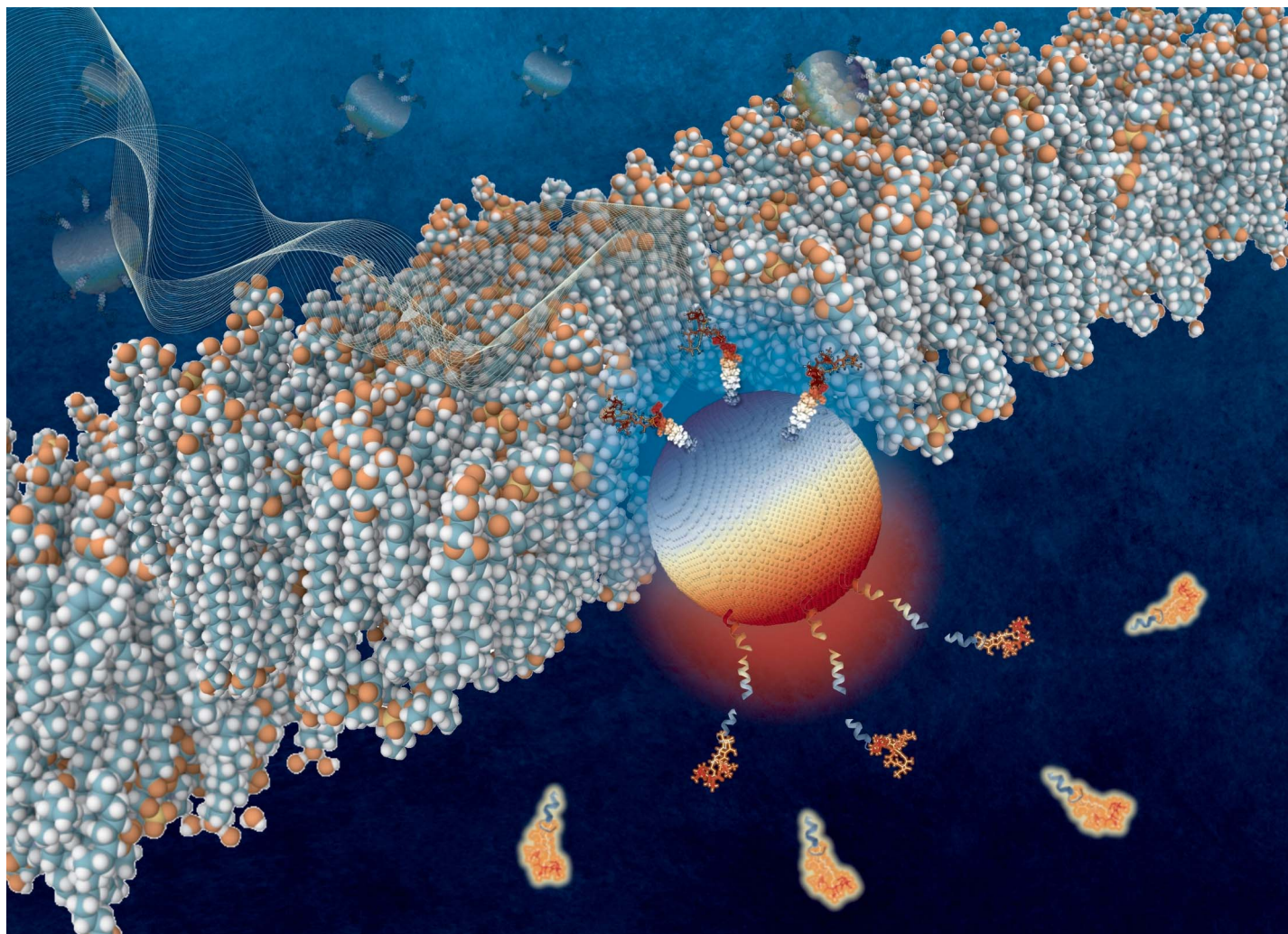




Showcasing research from the Electromagnetic Fields for Health (EMF4Health) group, Laboratory of Bioelectromagnetism “Emanuele Biondi” of CNR-IEIT – DEIB Politecnico di Milano, Milano, Italy.

Exploring magnetoelectric nanoparticles for advanced nano-electroporation and drug delivery in interventional cardiology

This work investigates the feasibility of magnetoelectric nanoparticles (MENPs) as magnetic field-mediated nanoscale electroporation and drug release agents. The research employs computational methods to study bioelectric phenomena at the interface between MENPs and biological tissues, when subjected to static and alternating magnetic fields, deriving electrical potential at the MENP surface and nanoparticle–drug bond charge displacement. This approach opens unique paths for wireless treatments in delicate medical fields such as interventional cardiology.

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Exploring magnetoelectric nanoparticles for advanced nano-electroporation and drug delivery in interventional cardiology

A. Tommasini,^{ab} G. Suarato,^{id} *^b S. Fiocchi,^b E. Chiaramello,^b A. Marrella,^b M. Lenzuni,^b M. Parazzini,^b B. Cortese^{cde} and P. Ravazzani^b

Leveraging their ability to induce an intense and highly localized electric polarization upon application of an external magnetic field, core-shell magnetoelectric nanoparticles (MENPs) constitute powerful nanotools to achieve wireless nano-electroporation and targeted drug delivery in critical applications such as cardiovascular medicine, where frequent interventions may pose general yet high risks to the patients. In this study, the magnetoelectric feature is exploited to thoroughly map *in silico* the behavior of a model MEMP under a set of either static or time-varying external magnetic fields, with the dual aim of (i) providing a series of operational parameters to elicit at the nanoscale a reversible cell membrane poration and (ii) correlating the electric potential difference developed at the activated MEMP surface with the charge displacement potentially induced in a hypothetical drug-MEMP bond. The finite element analysis framework considers a single cobalt ferrite core-barium titanate shell MEMP, either immersed in culture medium or engulfed in a blood vessel wall. Simulation results obtained under static magnetic field conditions show that maximizing magnetoelectric efficiency ($79.82 \text{ mV Oe}^{-1} \text{ cm}^{-1}$) generates electric fields in line with reversible nano-electroporation values (3.7×10^4 to $4.7 \times 10^4 \text{ V m}^{-1}$). Furthermore, a correlation between the profiles of the electric potential registered at the MEMP outer border (2.86–2.92 mV) and the charge displacement hypothetically experienced by a drug-MEMP bond is highlighted, when an alternating, low-intensity external magnetic field is used. Our computational study sets a preliminary base to gain a deeper understanding of the interface phenomena of MEMP-mediated cell poration and drug delivery, and it serves as a preliminary step towards complex future analyses.

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1. Introduction

Despite advances in medicine, particularly in the cardiovascular field, restenosis remains a common and significant problem following coronary artery revascularization. Pathologies, such as atherosclerotic plaques and cardiac ischemia, often require the use of stents to restore vessel patency and physiological blood circulation.¹ However, the incidence of in-stent restenosis (ISR), *i.e.* a progressive reduction in the vessel lumen size (reaching more than 50%) may be induced by an upregulated repair response, increases patient morbidity and may also cause

acute myocardial infarction.² Restenosis occurs when a vessel, usually a coronary artery, is damaged after percutaneous revascularization. The process that follows this event involves several interrelated mechanisms, and the trauma results in vascular injury, triggering a local inflammatory response and the production of cytokines and growth factors at the damaged site.³ The presence of these molecules causes vascular smooth muscle cells (VSMCs) to migrate into the innermost layer of the vessel, where they proliferate excessively. VSMCs begin to produce large amounts of extracellular matrix (ECM), consisting of collagen and proteoglycans, which gradually lead to the occlusion of the vessel lumen. Such ECM deposition concurs with vessel remodeling and, in parallel, thrombus formation on the damaged vessel surface, which further contributes to the restenosis process.⁴

To tackle this medical issue, in the late 1970s and mid-1980s Dotter and Judkins performed the first coronary angioplasty in a patient with a bare metal stent.⁵ Later, bare metal devices were replaced by drug-eluting stents, approved by the Food and Drug Administration in 2003,⁶ which improved physiological revascularization and lowered the mortality rate. Despite

^aDipartimento di Elettronica, Informazione e Bioingegneria, Politecnico di Milano, Piazza Leonardo da Vinci 32, Milano, 20133, Italy

^bCnr-Istituto di Elettronica e di Ingegneria Dell'Informazione e Delle Telecomunicazioni, Piazza Leonardo da Vinci 32, Milano, 20133, Italy. E-mail: giulia.suarato@cnr.it

^cInterventional Coronary Center, University Hospitals Harrington Heart & Vascular Institute, Cleveland, Ohio, USA

^dFondazione Ricerca e Innovazione Cardiovascolare, Via E. Ponti, 49, Milan, 20136, Italy

^eDCB Academy, Via E. Ponti, 49, Milan, 20136, Italy



optimization in stent design, the use of thinner struts and lightweight materials, or the research on more biocompatible coatings⁷ to further sustain the therapeutics release, persistent inflammation around the implanted stent and delayed endothelial repair remain critical in ISR.⁸ Later, a new class of devices has been developed, namely drug-coated balloons, which allow drug delivery without prosthesis implantation.^{9,10} This novel medical apparatus was designed to initially tackle ISR and later showed promising results also for the treatment of native coronary artery disease.¹¹

To this end, nanotechnology can offer a variety of curative approaches, especially when it comes to engineering platforms for targeted drug delivery. For example, micro- and nano-particles can find applications in a plethora of different medical fields (*i.e.*, tissue engineering, wound healing, and anticancer therapy, to name a few) due to their variety in terms of sizes, organic and inorganic components,^{12,13} morphologies, and fabrication and functionalization techniques.^{14,15} By tuning the surface chemistry of these nanosystems, site-specific and cellular-targeted delivery can be achieved¹⁶ and the drug loading content optimized,¹⁷ while challenges such as poor pharmacokinetics, drug instability under physiological conditions, or enzymatic degradation can be limited by properly combining cargo molecules with nanoparticle carriers.¹⁸ Leveraging their therapeutic potential, micro- and nano-polymeric particles or lipid-based micelles have been recently exploited for cardiovascular applications.^{7,19–21} To achieve a more precise spatial directionality towards a stenotic vessel region, magnetic nanoparticles, properly decorated with polymeric coatings and active drugs, have been locally and systemically administered to mice and subjected to an external static magnetic field. For example, Fellows *et al.*²² delivered heparin-coated magnetic nanoparticles for drug therapy of neointimal hyperplasia, while Polyak and co-workers²³ injected endothelial cells previously loaded with superparamagnetic nanoparticles, to allow for their blood navigation and targeted positioning near a stented area to inhibit ISR.

In addition to these more conventional systems, a novel class of multi-functional nanostructures is represented by magnetoelectric materials in which magnetic and electric fields can be strongly coupled at body temperature.²⁴ More specifically, magnetoelectric nanoparticles (MENPs) have recently emerged as a promising frontier in biomedical research due to their unique ability to act as nanoconverters.²⁵ In fact, these multi-phase structures are based on the combination of a magnetostrictive core and a piezoelectric shell, with different sizes and shapes, whose interface coupling can be optimized²⁶ to achieve desired features and performance.²⁷ Thanks to their ability to polarize and generate an intense electric field at their surface (order of magnitude of 10^5 V m⁻¹) and in their immediate nearby, MENPs have been investigated for (i) wireless brain and nervous tissue stimulation,^{28–31} (ii) cell manipulation,³² (iii) cancer treatment, through hyperthermia or targeting of cancer cells,^{33,34} (iv) tissue engineering,³⁵ and (v) nano-electroporation and drug delivery.^{36–38}

Classical electroporation methods use traditional electrodes to generate high-intensity electric fields for the

permeabilization of cell membranes. However, such electrodes are often invasive and difficult to apply in *in vivo* settings.^{39,40} Despite its efficacy, the application of a high intensity, non-localized electric field can lead to difficult and non-specific control of the process and even to loss of function or cell death (irreversible nano-electroporation). Greater spatial control of the generated electric field and high versatility in nano-electrode design can be achieved by using MENPs, which could act, when activated with a direct current magnetic field (H_{DC}), as electric field nano-generators within safe ranges (2×10^4 to 5×10^4 V m⁻¹),⁴¹ able to provide reversible and wireless cell membrane dielectric breakdown.⁴² Similarly, conventional approaches to delivering drugs must contend with poor specificity and loss of drug concentration, as well as difficulties in crossing biological barriers. MENPs can be easily and highly functionalized, as reported in extensive *in vitro* studies by Guduru *et al.*,⁴³ where nanoparticles were loaded with paclitaxel as a mitotic inhibitor for ovarian cancer treatment and labelled with a biomarker-specific antibody, and by Nair and coworkers,⁴⁴ who loaded MENPs with an anti-HIV compound. MENPs can achieve drug release following remote activation *via* alternating current, low-intensity magnetic fields (H_{AC}),⁴⁴ thus providing a field-controlled and highly specific delivery to the diseased tissue. Experimentally, the MENP-mediated drug release has been achieved with the application of an external H_{AC} of small magnitude (approximately below 50 Oe, with a frequency ranging from 10 to 100 Hz^{36,37}), which was sufficient to liberate a significant drug concentration into the cells.

By combining their ability to electroporate biological membranes with nanometric precision (when subjected to a H_{DC} field) alongside the possibility of remotely modulating their cargo molecule unloading (in the presence of a H_{AC} field), MENPs appear as a fascinating alternative to target pathological issues also in a yet unexplored field, such as interventional cardiology. In fact, the risk of restenosis and the need for further interventions could be tremendously reduced by employing MENPs, properly decorated with anti-inflammatory or anti-proliferative drugs.

With the aim of assessing the feasibility and functionality of such advanced nano-systems for cardiovascular drug release applications, the present study uses an *in silico* computational approach. A preliminary assessment of the MENP baseline behavior when subjected to an external magnetic field above its core magnetization saturation (M_s) was carried out to validate the soundness of our computational model and confirm the presence of the magnetoelectric effect (Fig. 1, step 1). Next, a series of simulations were carried out considering a single nanoparticle immersed in two different biological surroundings, such as the culture medium and the blood vessel wall tissue, exposed to magnetic fields of varying waveforms and intensities. A first parametric-sweep study under a static magnetic field (H_{DC}) (Fig. 1, step 2) was envisioned to define the operational ranges of MENP stimulation able to subsequently induce reversible membrane permeability. Afterwards, the external magnetic field assumes an alternating pattern (H_{AC}) (Fig. 1, step 3), which causes an oscillation of the generated potential difference across the nanoparticle, a phenomenon



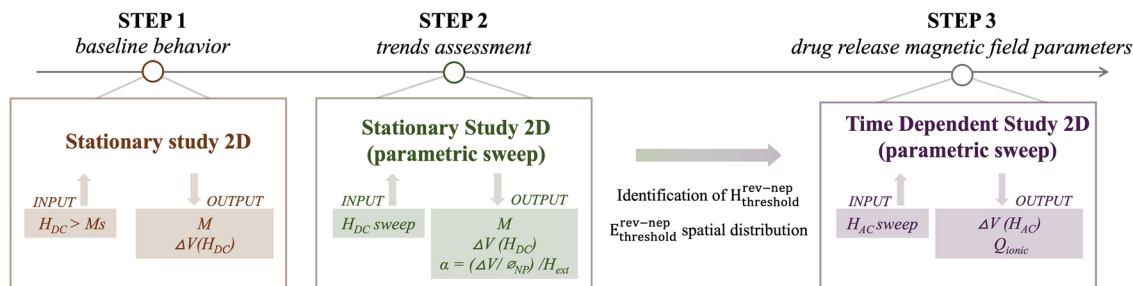


Fig. 1 MENP computational study workflow. Step 1: for the baseline behavior a static magnetic field (H_{DC}) above the material magnetization saturation (M_s) is used as input, while core magnetization (M) and the electric potential difference at the MENP surface ($\Delta V(H_{DC})$) are the output quantities. Step 2: for the sweeping stationary study, varying static magnetic fields (H_{DC}) are applied and core magnetization (M), electric potential difference $\Delta V(H_{DC})$, and magnetoelectric (ME) coefficient α are derived. This step is essential to identify $H_{threshold}^{Rev-Nep}$, which defines the minimum magnetic field required to generate an electric field ($E_{threshold}^{Rev-Nep}$) for the reversible nano-electroporation phenomenon to occur. Step 3: in the time dependent study, varying external, low-intensity alternating magnetic fields (H_{AC}) are imposed to derive the operational parameters governing the drug release process involved in a hypothetical drug-MENP ionic bond breaking. Outputs of step 3 are the electric potential $\Delta V(H_{AC})$ generated at the MENP outer shell and the term Q_{ionic} , which represents the charge displacement of a hypothetical drug-MENP ionic bond.

that has been hypothesized to be related to drug bond weakening and, consequently, drug release. The workflow reported in Fig. 1 summarizes the computational steps, highlighting the operational parameters considered in the analysis.

2. Materials and methods

2.1 Nanoparticle modeling

COMSOL Multiphysics® 5.6 (<https://www.comsol.com>), a computational tool based on the finite element approach, was adopted as the simulation environment to study the magnetoelectric response of a single, core-shell structured MENP, by means of an axisymmetric bi-dimensional (2D) model. As shown in Fig. 2, the system under study was composed of three domains: (i) a piezoelectric barium titanate shell (BaTiO_3 , in the following referred to as “BTO”), presenting a uniform thickness of 25 nm; (ii) a magnetostrictive cobalt ferrite core (CoFe_2O_4 , in the following referred to as “CFO”), with a diameter of 90 nm; and (iii) a surrounding environment

defined as a squared domain with a side length of 2000 nm. The properties of the shell and core materials, mostly derived from experimental studies and the COMSOL Multiphysics® 5.6 integrated library, are reported in Tables 1 and 2, respectively.

Two different scenarios were modeled, based on the environment in which the nanoparticle was embedded: (i) single MENP floating in the culture medium in a Petri dish; (ii) single MENP enclosed in the blood vessel wall. In both simulations, the surrounding environments were considered as purely resistive materials, with a relative permittivity (ϵ_r) of 1; the culture medium was modelled as a liquid environment, while the blood vessel was considered a solid material (properties reported in Table 3).

2.2 Stationary study

Our investigation was conducted in three main phases, as outlined in the workflow of Fig. 1. In the first phase, a stationary study (*i.e.*, the MENP was subjected to a high-intensity external static magnetic field, such as 2T, above the core magnetization saturation) was performed to evaluate the baseline magnetoelectric behavior of the MENP within the two distinct environments. Both environments were treated as purely resistive materials with the relative permittivity (ϵ_r) set to 1. For the culture medium, the electrical conductivity (σ) was set to 1.5 S m^{-1} , as found in the literature;⁵³ while for the blood vessel wall, the electrical conductivity was set to 0.232 S m^{-1} , as reported in the IT'IS Foundation's Virtual Population Tissue Properties Database.⁵¹ Additionally, for the case of the blood vessel wall, the surrounding was modeled as a solid material, characterized by mechanical properties such as Young's modulus, Poisson's ratio and density, as detailed in Table 3. Following the validated computational design previously published by our group,^{27,31,34} *in silico* MENP activation was obtained by implementing three different COMSOL Multiphysics® 5.6 modules, such as Magnetic Fields, Solid Mechanics, and Electrostatics, while the Multiphysics mode allowed for the coupling of Magnetostriction and Piezoelectric physics for a complete simulation of the

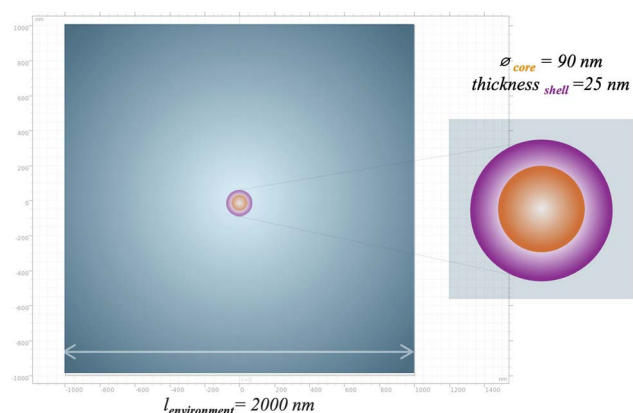


Fig. 2 Computational model of the core-shell magnetoelectric nanoparticle, immersed in the surrounding material, with sizes of the geometry.



Table 1 Barium titanate shell properties

Property	Symbol	Value	Reference
Density (kg m ⁻³)	ρ	5.2×10^3	45, COMSOL library
Relative permittivity	ϵ_r	10	COMSOL library
Electrical conductivity (S m ⁻¹)	σ	5.2×10^6	COMSOL library
Poisson's ratio	ν	0.48	46, COMSOL library
Young's modulus (Pa)	E	230×10^9	46 and 47
Saturation magnetization (A m ⁻¹)	M_s	3.69×10^5	48
Initial magnetic susceptibility	χ_0	3	49, COMSOL library
Saturation magnetostriction (ppm)	λ	-200	42 and 46

physical phenomenon. The mathematical equations governing the magnetoelectric coupled model have been thoroughly described by Flocchi *et al.*⁵⁴ and briefly summarized in the SI.

2.3 Sweep study – static external magnetic field

In the second step of the study, the magnetoelectric behavior of a single MENP subjected to a wider range of static magnetic field values was assessed. In the *Study* section of the COMSOL Multiphysics® 5.6 interface, the *Stationary Study* was added, and a parametric sweep was selected. The imposed external magnetic field was varied from 0 to 4 T, considering the following intervals: 2–50 mT, 100–800 mT, and 1–4 T. The “*entry method*” of range was set on the “*number of values*”, and fifteen values of each sub-interval were analyzed. Particular attention was given to key parameters, such as (i) the degree of magnetization of the CFO core, (ii) the electric potential generated on the BTO shell surface, and (iii) the associated ME coefficient. The latter is defined according to the equation $\alpha_{ME} = \frac{\Delta V}{\Delta H \times \phi_{MENP}}$, as the ratio between the electric potential difference on the MENP surface (ΔV), expressed in mV, and the product of the applied external magnetic field (H), expressed in Oe, and the MENP diameter (ϕ_{MENP}), given in cm.

At this step of the investigation, particular attention was paid to the electric field generated by the particle within its surroundings. To do so, three representative values of the external magnetic field (H_{ext}) were considered for the triggering of the magnetoelectric phenomenon and the following data analysis: (1) $H_{ext(1)}$ identified in the range of linear dependence between α_{ME} and the magnetic field; (2) $H_{ext(2)}$ corresponding to the maximum α_{ME} with the aim of taking advantage of the

optimal MENP performance; (3) $H_{ext(3)}$ corresponding to the minimum magnetic field yielding the highest electric potential. In particular, $H_{ext(3)}$ was identified as the value of the magnetic field at which the increment between two consecutive potential differences was less than 0.15 mV. A MATLAB code was implemented to examine the near surroundings of the nanoparticle starting from the electric field data distribution exported from COMSOL software. The regions of interest were modeled as concentric, circular semi-coronas around the nanoparticle, identifying areas within three different, increasing distances from the MENP surface. More specifically, the radii of the semi-coronas considered for the analysis were 75, 80 and 90 nm to sample electric field values located at distances of 5, 10 and 20 nm, respectively, from the shell external border ($r_{MENP} = 70$ nm). For each configuration, the electric field distribution was visualized, and its statistical metrics (in terms of median value and the 1st and the 99th percentiles) were extracted to allow for data comparison. More specifically, three regions were considered, depending on how likely the generated electric fields could cause Irreversible Nano-electroporation (Ir-Nep), Reversible Nano-electroporation (Rev-Nep) or none of the two. Such a tendency was assessed by checking the 1st and 99th percentiles of each distribution. The analysis was conducted for the two case scenarios presented in paragraph 2.1 (*i.e.*, culture medium and blood vessel wall). The outputs were processed *via* Origin-Pro 2023 (OriginLab, United States) and presented as box plots, highlighting median values and percentiles. As summarized in Fig. 1, the aim of this intermediate step was to identify the optimal magnetic field ($H_{threshold}^{Rev-Nep}$) upon activation of which an electric field comparable to those used in conventional reversible electroporation ($E_{threshold}^{Rev-Nep}$) could be

Table 2 Cobalt ferrite core properties

Property	Symbol	Culture medium		Blood vessel wall	
		Value	Reference	Value	Reference
Electric conductivity (S m ⁻¹) 76 543	σ	1.5	50	0.232	51
Relative permittivity	ϵ_r	1	—	1	—
Relative permeability	μ_r	1	—	1	—
Young's modulus (MPa)	E	—	—	1	52
Poisson's ratio	ν	—	—	0.49	52
Density (kg m ⁻³)	ρ	1007	51	1102	51



Table 3 Properties of the surrounding materials considered in the different simulation scenarios

Property	Symbol	Culture medium		Blood vessel wall	
		Value	Reference	Value	Reference
Electric conductivity (S m^{-1})	σ	1.5	50	0.232	51
Relative permittivity	ϵ_r	1	—	1	—
Relative permeability	μ_r	1	—	1	—
Young's modulus (MPa)	E	—	—	1	52
Poisson's ratio	ν	—	—	0.49	52
Density (kg m^{-3})	ρ	1007	51	1102	51

developed in the immediate vicinity of the MENP. Such H_{field} and E_{field} and their spatial distributions will be essential for the last step of the investigation.

2.4 Sweep study – time-dependent analysis

Lastly, a time dependent study was performed, with the aim of analyzing the magnetoelectric behavior of the nanoparticle exposed to a time-dependent field. This step is particularly relevant for the achievement of the final application herein envisioned, namely the use of MENPs for the combined action of nano-electroporation and controlled drug release. The analysis was divided into two phases, as sketched in Fig. 3: (1) for the reversible nano-electroporation phase, a constant H_{DC} of 300 mT (*i.e.*, the optimal value of $H_{\text{threshold}}^{\text{Rev-Nep}}$ obtained from the stationary parametric sweep study presented in paragraph 2.3) was applied and switched off after $\Delta t_{\text{Rev-Nep}}$ (time of nano-electroporation, arbitrarily chosen); (2) for the drug release phase, a sinusoidal wave was applied for Δt_{dr} (time of drug release, arbitrarily chosen) with a frequency of 100 Hz and varying amplitudes (H_{AC} ranging from 1 mT to 6.5 mT). The first phase (at constant H_{DC}) aimed at generating an electric field strong enough to induce reversible nano-electroporation on the cell membrane of a hypothetical cell located in the MENP nearby (within a distance of 5, 10 or 20 nm from the particle

outer shell), while the subsequent sinusoidal phase facilitated the generation of alternating electric potentials on the surface of the nanoparticle, which is considered prodromal to drug molecule dissociation. The analysis was conducted for the two case scenarios presented in paragraph 2.1 (*i.e.*, culture medium and blood vessel walls).

To properly model the MENP behavior within a time-dependent study, the Jiles–Atherton (J–A) hysteresis model was implemented in the *Solid Mechanics* section in the software interface model builder, for the magnetostrictive core. The COMSOL entries in the core material settings for the J–A model parameters have been modified according to Table 4, considering our group's previous work.²⁷ The mathematical equations governing the J–A model are summarized in the SI. In the *Study* section of the COMSOL interface, a *Time Dependent Study* was

Table 4 Jiles–Atherton model parameters for magnetic hysteresis

Property	Symbol	Value	Reference
Magnetic saturation (A m^{-1})	M_s	3.69×10^5	48
Pinning loss (A m^{-1})	k	2×10^5	56
Domain wall density (A m^{-1})	a	1.5×10^5	56
Inter-coupling domain	α	1.5	56
Magnetic reversibility	c	0.3	56

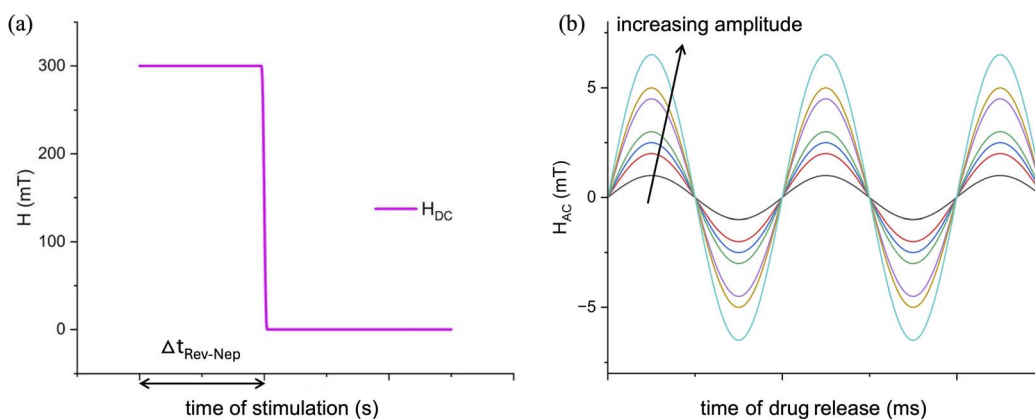


Fig. 3 Computational stimulation settings of the time-dependent sweep study: external H_{field} profiles. (a) In the first phase ($\Delta t_{\text{Rev-Nep}}$ = time of nano-electroporation), a static magnetic field with a constant amplitude phase ($H_{\text{DC}} = 300$ mT) is applied, to allow for a reversible cellular nano-electroporation, while in the following step (b) (Δt_{dr} = time of drug release) an external magnetic field presenting a sinusoidal phase with variable amplitudes (H_{AC}) is activated (keeping constant the frequency at 100 Hz), to favor the hypothetical drug–MENP bond loosening, initiating the drug delivery process.



added, and a parametric sweep was imposed. The *Parameter Value List* section was filled in with specific values (such as 1, 2, 2.5, 3, 4.5, 5, and 6.5 mT). For the nano-electroporation phase, the magnetization profile was studied, paying particular attention to the value of magnetization obtained after $\Delta t_{\text{Rev-Nep}}$ (nano-electroporation time), *i.e.* when the constant static magnetic field was switched off. For the drug release phase, the core magnetization values, the electrical potential differences (ΔV) and the α_{ME} , observed/calculated at a given time, were plotted in relation to the increasing amplitude of the sinusoidal wave. Ultimately, the concept of charge displacement was considered, derived as a function of the alternating magnetic field amplitudes and the efficiency factor (α_{ME}). Such physical quantity, referred to as Q_{ionic} , gives an indication of the stability involved in a hypothetical drug-MENP ionic bond (see the SI and Stimphil *et al.*⁵⁵). All the outputs were processed *via* OriginPro 2023 (OriginLab, United States).

3. Results

3.1 MENP baseline behavior under stationary field conditions

In the first part of the investigation (Fig. 1, step 1), a stationary study with a magnetic field above the material magnetic saturation (M_s) was carried out to evaluate the magnetoelectric baseline behavior of the MENP in two separate environments. These stationary analyses aimed at validating (i) the generation of an electric potential at the surface of the particle and (ii) the establishment of an electric field in its vicinity, starting by studying the magnetization of all CFO domains, the following strain developed, and the subsequent stress generated at the BTO piezoelectric shell.

The steps that a spherical magnetoelectric nanoparticle of our dimensions goes through, when ideally immersed in the blood vessel wall and exposed to a strong magnetic field of 2T (a field above its magnetization saturation, homogeneous and directed along the z-axis, considered the “easy axis” for the MENP magnetization) are shown in Fig. 4. Upon switching on the high intensity magnetic field, indicated by the black arrows pointing upwards, the CFO core becomes completely magnetized (Fig. 4 step 1, dark core, herein represented by the distribution of the magnetization, expressed in A m^{-1}). Through

deformation coupling of a magnetostrictive and piezoelectric material, magnetoelectricity can be exploited directly. Therefore, strain at the core level and stress at the shell level are reported (Fig. 4, step 2). The inhomogeneous stress distribution is the highest (but always within several tenths of ppm) along the sides of the nanoparticle. The magnetoelectric coupling causes a movement of charges, resulting in a maximum potential difference at a shell outer border of 6.07 mV (Fig. 4, step 3), and the consequent generation of an electric field in the MENP surrounding (Fig. 4, step 4), with values up to $7 \times 10^4 \text{ V m}^{-1}$, sharply decreasing downwards with the distance from the outer surface of the shell. A similar electric behavior, with highly comparable values, is observed when the MENP is immersed in the culture medium, and the resulting distributions are shown in Fig. S2 and Table S1 of the SI.

3.2 Trend assessment of the MENP performance under stationary field conditions

Subsequently, the magnetoelectric behavior of a single MENP in the presence of a wider range of static magnetic field values (from a low-intensity field of 2 mT to an intense field of 4 T) was investigated (Fig. 1, step 2). The magnetization profiles (magnetization, expressed in A m^{-1}) registered at the core of the MENP for either the culture medium (in the following abbreviated as “CM”) case or the blood vessel wall (in the following abbreviated as “BV”) case, as a function of the applied external magnetic fields, are shown in Fig. 5a. In both environments, for low-intensity H_{field} (2–50 mT), the magnetization profile presents a linear trend, reaching a value of $4.72 \times 10^4 \text{ A m}^{-1}$ (at 40 mT). Then, the linear tendency progressively translates into a curved profile (H_{field} of 100–800 mT and magnetization of $1.16\text{--}3.40 \times 10^5 \text{ A m}^{-1}$), until it reaches a maximum magnetization value corresponding to an applied magnetic field of about 650 mT. Beyond this point, the magnetization remains constant, defining a plateau until the maximum magnetic field value applied in this study (4 T) is reached.

The MENP behaves as an electric dipole when exposed to an external magnetic field. To evaluate the magnetoelectric coefficient (α_{ME}), according to eqn (1), the electric potential difference generated on the surface of the MENP was considered (Fig. 5b). The coefficient α_{ME} rapidly increases in the first part of

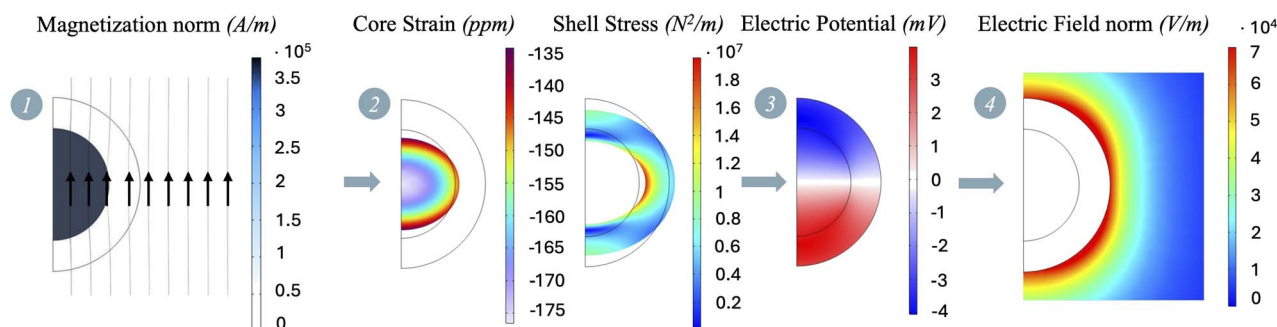


Fig. 4 MENP baseline behavior under static stimulation conditions: from the magnetization of the CFO core (1), passing through the core strain and shell stress (2), to the generation of an electric potential on the particle surface (3), and the electric field in its vicinity (4).

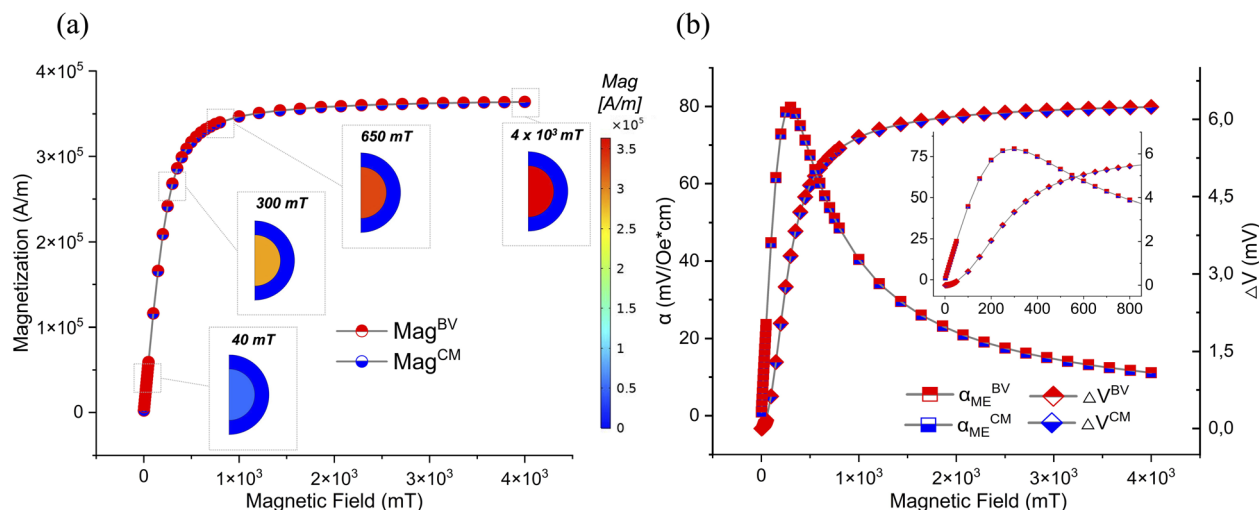


Fig. 5 Magneto-electric nanoparticle performance: (a) core magnetization trend of the MENP enclosed in the blood vessel wall (BV) or immersed in the culture medium (CM) at increasing applied magnetic fields (H_{fields}); color-coded visualization is also included for the representative cases of H_{field} of 40 mT, 300 mT, 650 mT, and 4T. (b) Profiles of the magneto-electric coefficient (α_{ME} , expressed in $\text{mV Oe}^{-1} \cdot \text{cm}^{-1}$) and the electric potential difference (ΔV , expressed in mV) developed across the MENP diameter, when the MENP is enclosed in the blood vessel wall (BV) or immersed in the culture medium (CM) at increasing applied H_{fields} .

the curve (H_{field} in the range of 2–50 mT), until it reaches a peak of about $79.82 \text{ mV Oe}^{-1} \cdot \text{cm}^{-1}$, corresponding to a magnetic field of 300 mT. Beyond this value, the curve gradually decreases asymptotically, dropping down to $11.15 \text{ mV Oe}^{-1} \cdot \text{cm}^{-1}$ at high-intensity H_{field} values. Fig. 5b also reports on the evolution of the electric potential (ΔV , calculated as the difference between the maximum and minimum values of the two opposite poles generated on the surface of the MENP, and expressed in mV), which shows a similar profile to that previously described for the magnetization: an initial linear trend at low-intensity H_{field} (reaching a value of 0.11 mV at 40 mT), followed by the attainment of a plateau (around 6.22 mV). Furthermore, the trend of this parameter in the two different environments under study appears to perfectly overlap.

3.3 Reversible nano-electroporation activation area

Next, the electric field generated by the particle in the different surroundings was analyzed. The external magnetic field (H_{ext}) values considered are: $H_{\text{ext}(1)}$, identified as 150 mT, corresponding to a value within the linear dependency between α_{ME} and H ; $H_{\text{ext}(2)}$, *i.e.* 300 mT, corresponding to the maximum α_{ME} (Fig. 5b), with the aim of taking advantage of the optimal performance of the MENP; and $H_{\text{ext}(3)}$, namely 650 mT, corresponding to the magnetic field value at which the increase of two consecutive potential differences was below 0.15 mV (Fig. 5b). According to Fig. 1 (proceeding from step 2 towards step 3), the aim was to identify the optimal magnetic field ($H_{\text{threshold}}^{\text{Rev-Nep}}$) that could generate an electric field such as $E_{\text{threshold}}^{\text{Rev-Nep}}$ in the particle nearby.

The left side of Fig. 6 shows the electric field distributions at distances of 5, 10 and 20 nm from the outer border of the MENP shell ($r_{\text{MENP}} = 70 \text{ nm}$), as extrapolated from COMSOL software and processed with MATLAB (see paragraph 2.2 in the Materials

and methods section). On the right side of Fig. 6, the statistical metrics of the electric field distributions (in terms of median values and the 1st and the 99th percentiles) are reported, for each of the three progressively larger areas considered. Furthermore, the statistical metrics of the electric field distributions generated by the external magnetic fields under study are presented in Table 5, to ease the data comparison. While the data presented in Fig. 6 and Table 5 refer to the case of a MENP engulfed in the blood vessel wall, a similar analysis was also carried out for the case of the culture medium as the surrounding environment (Fig. S3 and Table S1). According to the data, stimulation with a magnetic field of 150 mT appears to be below the threshold that triggers a biological membrane response (above $2 \times 10^4 \text{ V m}^{-1}$), while at 650 mT, E_{fields} distributions falling in the range of irreversible nano-electroporation are obtained (above $5 \times 10^4 \text{ V m}^{-1}$). Therefore, maintaining conservative conditions, the following analyses have been carried out considering an external H_{DC} of 300 mT, whose generated E_{fields} distributions completely remain in the range of reversible nano-electroporation (2×10^4 to $5 \times 10^4 \text{ V m}^{-1}$).

3.4 Drug release operational ranges: time-dependent study

To complete our investigation (Fig. 1, step 3) and analyze the magneto-electric behavior of the nanoparticle exposed to a time-dependent field, a time-dependent study was carried out. This step is crucial for achieving the final application herein envisaged, such as the exploitation of MENP-based nanotechnology for combined nano-electroporation and controlled drug release in cardiovascular applications.

The analysis comprises two phases: (i) induction of reversible nano-electroporation on the cell membrane of a hypothetical cell in the vicinity of the MENP, triggered by a static magnetic field (H_{DC}); (ii) promotion of the dissociation of drug



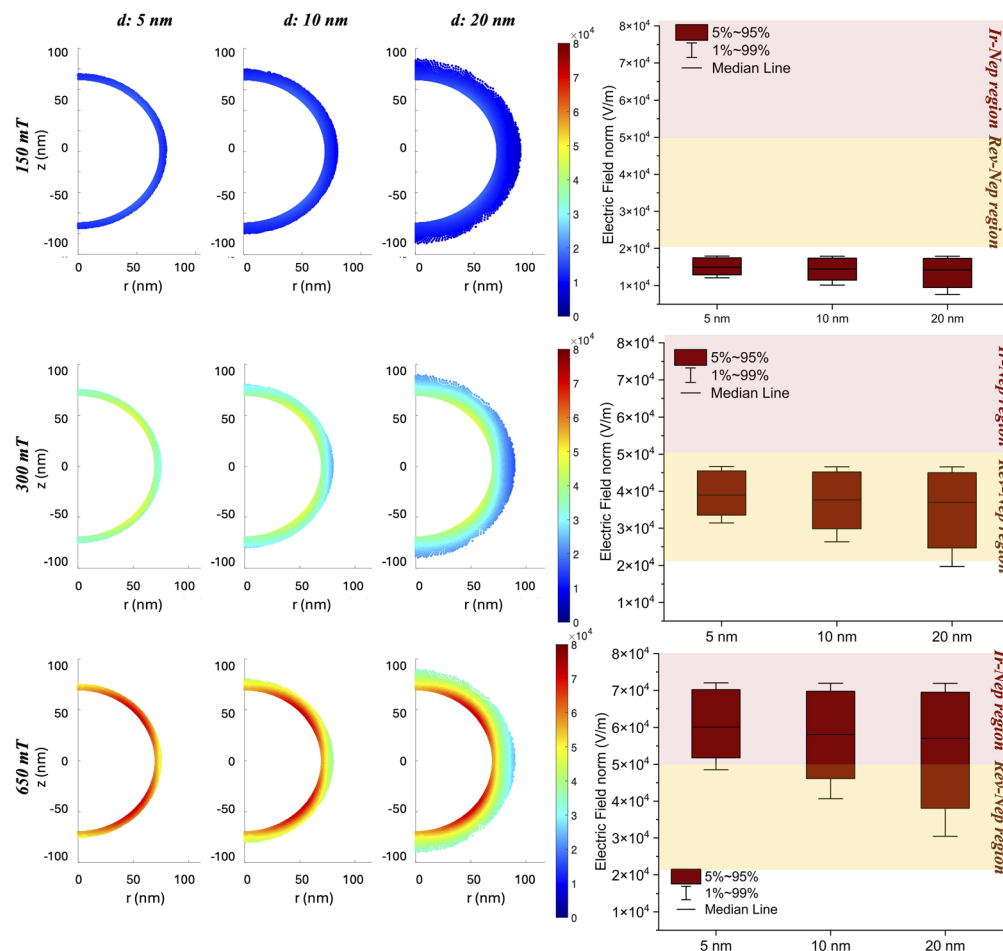


Fig. 6 Nano-electroporation activation area. Left: distribution of E_{field} obtained after H_{DC} stimulation of a MENP engulfed in the blood vessel wall, at gradually increasing areas from the nanoparticle ($d = 5$ nm, 10 nm and 20 nm, distance from the outer shell); right: box plots of statistical metrics of E_{field} distributions in terms of median values, and the 1st and the 99th percentiles. Different nano-electroporation regions are highlighted: irreversible nano-electroporation (Ir-Nep region, in red) and reversible nano-electroporation (Rev-Nep region, in yellow).

molecules hypothetically bonded at the nanoparticle surface, by applying a H_{AC} stimulation, thus leveraging the generation of an alternating electric potential at the MENP outer shell. To achieve the reversible nano-electroporation, a H_{DC} of 300 mT (as obtained from the analysis reported in Fig. 6 and Table 1) is initially applied for $\Delta t = t_{\text{Rev-Nep}}$ (Fig. 7a, pink line) and then switched off. As this study is not intended to propose the definition of a precise protocol for nano-electroporation but is primarily interested in assessing the MENP-mediated process feasibility, specific time intervals are not considered. The magnetization profiles of the MENP core over time generated by

such static magnetic field stimulation are reported in Fig. 7a for the case of both environments (blood vessel – red line, and culture medium – dashed blue line), with the aim of understanding how the nanoparticle core behaves during and after the removal of the stimulation. According to Fig. 7a, the core magnetization obtained at 300 mT is roughly $5.65 \times 10^4 \text{ A m}^{-1}$ and remains constant over $\Delta t_{\text{Rev-Nep}}$. When the H_{DC} is switched off, the core retains a residual magnetization of approximately $5.2 \times 10^4 \text{ A m}^{-1}$ ($\sim 10 \text{ emu g}^{-1}$) for both simulation environments, except for a negligible delta. Such residual

Table 5 Statistical metrics in terms of median values and the 1st and the 99th percentiles of E_{field} distributions generated by $H_{\text{ext}(1)}$, $H_{\text{ext}(2)}$ and $H_{\text{ext}(3)}$

	$H_{\text{ext}(1)} = 150 \text{ mT}$			$H_{\text{ext}(3)} = 300 \text{ mT}$			$H_{\text{ext}(3)} = 650 \text{ mT}$		
(V m^{-1})	5 nm	10 nm	20 nm	5 nm	10 nm	20 nm	5 nm	10 nm	20 nm
Median	1.50×10^4	1.45×10^4	1.43×10^4	3.90×10^4	3.77×10^4	3.70×10^4	6.01×10^4	5.81×10^4	5.71×10^4
99th percentile	1.80×10^4	1.80×10^4	1.79×10^4	4.67×10^4	4.66×10^4	4.66×10^4	7.21×10^4	7.19×10^4	7.19×10^4
1st percentile	1.21×10^4	1.02×10^4	0.76×10^4	3.14×10^4	2.63×10^4	1.97×10^4	4.85×10^4	4.06×10^4	3.04×10^4

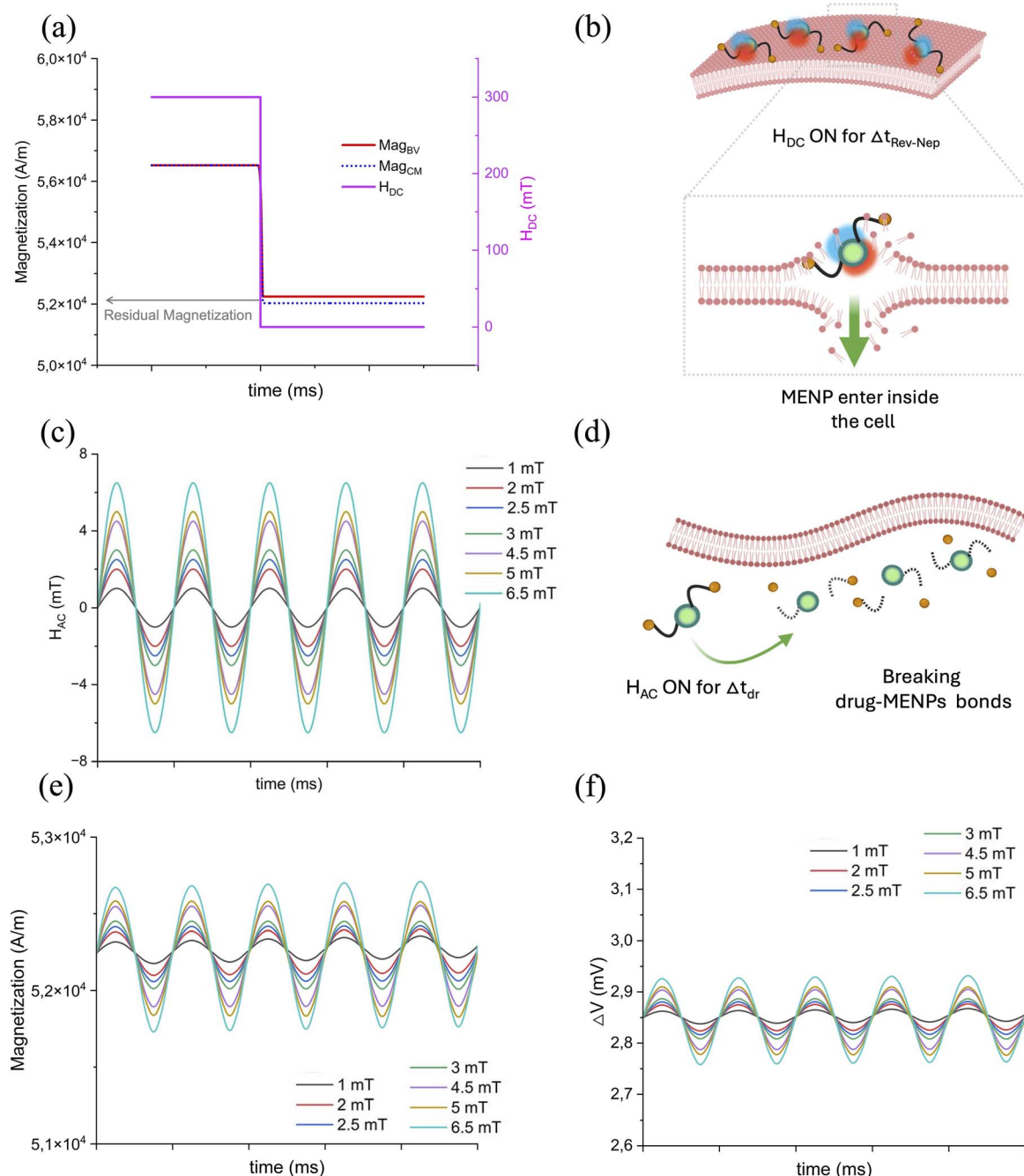


Fig. 7 Time dependent study for drug release: (a) static magnetic field profile (H_{DC} , expressed in mT), switched off after $\Delta t_{Rev-Nep}$, and core magnetization trend over time (magnetization, expressed in $A\ m^{-1}$) in the blood vessel wall (red line) or in the culture medium (dotted blue line), to allow for a reversible cellular nano-electroporation; (b) schematic illustration of the hypothesized MENP journey to cross a cell membrane after application of a H_{DC} for the time required for the reversible nano-electroporation phenomenon to occur; (c) alternating magnetic field (H_{AC} , expressed in mT) with variable amplitude profiles for Δt_{dr} prodromal to the drug delivery process; (d) graphical representation of the hypothetical drug-MENP bond dissolution and drug release under H_{AC} stimulation; (e) MENP core magnetization behavior; and (f) electric potential difference profile over time for different H_{AC} .

magnetization is retained for $t > t_{Rev-Nep}$, thus leaving the particle in a "magnetized" state.

Next, the drug release step is considered to investigate how the alternating magnetic field may affect the MENP specific behavior. In Fig. 7c, sinusoidal waves with the different amplitude values considered in the study are graphed. The core

magnetization and the electrical potential difference profiles are plotted in Fig. 7e and in Fig. 7f, respectively, for a single MENP engulfed in the blood vessel well. For the case of the CM scenario, the corresponding plots and values are reported in SI, Fig. S4 and Table S2. As noticeable from Fig. 7e and f and Table 6, higher values of magnetization and ΔV are obtained when the

Table 6 MENP core magnetization, electric potential difference and magnetoelectric coefficient values in relation to the increase in the amplitude of the applied H_{AC} for the BV case. Each parameter is calculated at the instant when the peak of the H_{AC} sinusoidal wave is reached

H_{AC} (mT)	Magnetization ($A\ m^{-1}$)	ΔV (mV)	α_{ME} ($mV\ cm^{-1}\ Oe^{-1}$)
1	5.23×10^4	2.86	87.36
2	5.24×10^4	2.87	84.00
2.5	5.24×10^4	2.88	83.43
3	5.24×10^4	2.88	83.88
4.5	5.25×10^4	2.90	81.70
5	5.26×10^4	2.91	82.07
6.5	5.27×10^4	2.92	79.85

amplitude of the sinusoidal stimulation is greater (H_{AC} ranging from 4.5 mT to 6 mT), with more pronounced oscillations, and values reaching $5.27 \times 10^4\ A\ m^{-1}$ and 2.92 mV, respectively. Similarly to the previous analysis carried out for the trend assessment of MENP performance under steady state conditions (paragraph 3.2), the values of core magnetization,

electrical potential difference and magnetoelectric coefficient related to the increase of the alternating field amplitude at a given time point (arbitrarily chosen) are reported in Table 6.

Fig. 8 reports simulation data comparison involving the concept of average charge displacement (referred to as Q_{ionic} ; the detailed explanation is available in the SI) to which hypothetical drug-MENP ionic bonds may be subjected when the system is immersed in a homogenous, low-intensity H_{AC} field at 100 Hz. The Q_{ionic} profiles at different H_{AC} amplitudes over the studied period reveal a trend comparable to that of the core magnetization and potential difference (Fig. 8a): an increase in the stimulation sine wave amplitude is associated with more visible oscillations. The susceptibility to rupture of hypothetical drug-MENP bonds is related to the alternating magnetic field values, which, in turn, cause oscillation of the electric potential difference. Therefore, the trend of Q_{ionic} versus ΔV at a given time is also shown (Fig. 8b), highlighting a linear relationship, with charge shift values ranging from 1.29 to 1.32×10^{-17} coulomb (C), which are in line with Nair *et al.*⁴⁴ Lastly, Fig. 8c compares the evolution over time of the electric potential difference and the average charge displacement of the hypothetical ionic drug-MENP bonds from their equilibrium

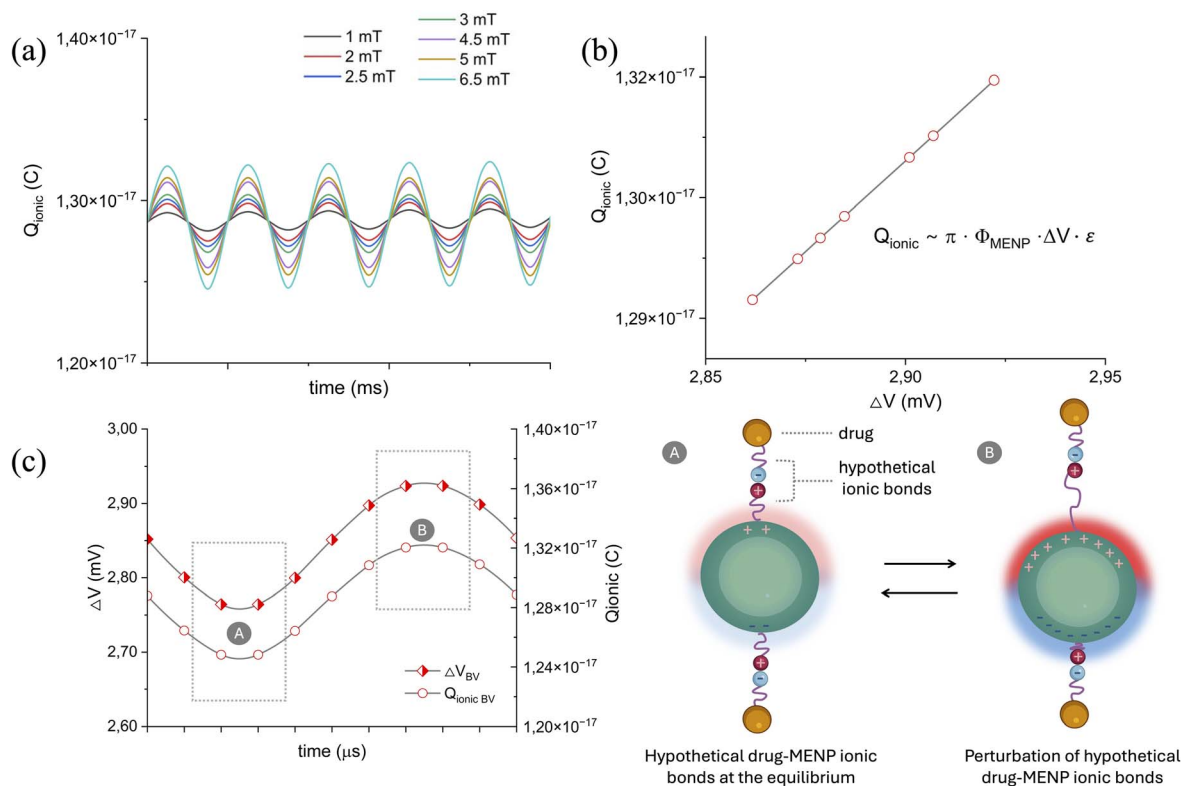


Fig. 8 Ionic bond charge displacement parameter: (a) charge displacement in the hypothetical drug-MENP ionic bond profile over time for different H_{AC} amplitudes; (b) charge displacement in the hypothetical drug-MENP ionic bond values in relation to the various electric potential differences generated at the MENP surface at varying H_{AC} stimulations, considered at a specific time point; (c) comparison between the trends of the charge displacement in the hypothetical drug-MENP ionic bond and the electric potential difference over a certain period of time. In (c), as a representative example, the $H_{AC} = 6.5$ mT case was considered for the data presentation. Drug-MENP sketch: case A illustrates two hypothetical drug-MENP ionic bonds at equilibrium, where the electric potential difference is at its minimum and so the average displacement of charges is negligible; case B represents a perturbed state of the ionic bonds, characterized by the movement of the positive and negative charges from their equilibrium positions, which causes bond length stretching and bond length contraction. In A, less intense red and blue colorations indicate lower ΔV at the MENP surface, while in B more intense blue and red shadows refer to higher ΔV .

positions. As a representative example, the $H_{AC} = 6.5$ mT stimulation case was considered for the BV scenario: as expected, Q_{ionic} and ΔV oscillate in a reciprocal manner. For the CM case, equivalent results are shown in the SI (Fig. S5).

4. Discussion

Nanotechnology advancements have recently brought to light new nanomaterials, which greatly facilitate the development of cutting-edge biomedical strategies to interact with biological systems and tissues in a wireless and on-demand way. Nanomaterials appear to be an intriguing option to face pathological problems in fields such as interventional cardiology and coronary artery surgery, due to their capacity to pass through biological barriers and allow a stimuli-responsive modulation of the anti-inflammatory or anti-proliferative drug release. Notably, magnetoelectric (ME) nanomaterials, able to induce electrical polarization on their surface when exposed to a magnetic field, constitute a one-of-a-kind tool, whose properties and stimuli-responses can be remotely activated and tuned by acting on their material composition, size, geometrical anisotropy, and combination with organic compounds.^{57,58} Among the most widely studied ME material configurations are core-shell magnetoelectric nanoparticles (MENPs), whose structure allows the maximization of interfacial coupling between the magnetostrictive core phase and the piezoelectric shell phase, thus optimizing the performance.^{27,54} The most widely used and promising material combination is based on a cobalt ferrite (CFO) core and a barium titanate (BTO) shell, presenting good biocompatibility and a stable magnetoelectric effect.⁵⁹ A quantification of the MNP performance is given by the magnetoelectric coefficient (α_{ME}), which considers the magnetoelectric coupling in terms of the generated electric field (output) and the external magnetic field (input). There are a limited number of studies in the literature reporting or mapping α_{ME} , due to difficulties in measuring it experimentally and the lack of standardized protocols (e.g., single particle *vs.* bulk materials, in-house measurement apparatus). Such a limitation is related to the intrinsic material parameters of the components, which, in turn, are influenced by complex structural features such as magneto-crystalline anisotropy, domain distribution, domain wall motion of the phases, crystal lattice mismatch and the presence of point defects at the interface.²⁸ Computational analyses come to the rescue by deriving significant parameters and predicting the overall behavior of the system. In this regard, recent simulation studies have reported values of α_{ME} ranging from $0.01 \text{ V cm}^{-1} \cdot \text{Oe}^{-1}$ to approximately $2.5 \text{ V cm}^{-1} \cdot \text{Oe}^{-1}$, to define the MNP conversion efficiency under various combinations of static and alternating magnetic fields and by examining various shapes and geometries.^{27,28}

The *in silico* work starts with a validation of the ME effect of a MNP engulfed in the blood vessel wall or immersed in the culture medium, as modeling environments, to observe the generation of an electric potential on the MNP surface and an electric field in the surroundings, in response to a 2T static magnetic field (above the core magnetization saturation). The MNP baseline behavior is analyzed in Fig. 4: the magnetization

involves a realignment of the magnetic domains within the core, a compression along the direction of the applied magnetic field (*i.e.*, the *z*-axis), and an expansion in the other two dimensions, leading to a core dimensional change (lattice deformation) on the order of a hundred ppm. This deformation is imparted to the piezoelectric shell, transforming the stress exerted by the core into a change in the charge position within the atoms of the piezoelectric lattice, thus generating a surface potential and an electric field distribution on the MNP and in its vicinity, which are consistent with those found in previous studies conducted by our group,³⁰ where MNPs have been proposed for peripheral nerve stimulation^{29,60} and sensing,⁶¹ and nerve regeneration,⁶² highlighting the versatility of the adopted computational framework. After having confirmed the basic response of the MNP, a wider range of DC magnetic field values is assessed, and the CFO-BTO nanoparticle parameters are thoroughly characterized. The results of the calculated magnetization recorded at the core of the MNP (Fig. 5a) describe a curve whose trend agrees with the one of Ma *et al.*⁶³ In fact, when the applied magnetic field gradually increases, the domains become progressively aligned with its direction and the magnetization values rise. At a certain point, the magnetization profile reaches a plateau, because of the complete orientation of the magnetic domains in the direction of the applied field, and any further increase in the external stimulus does not translate into a stronger magnetization. The trend of the magnetoelectric coefficient as a function of the applied field (Fig. 5b) is in accordance with the work of Kumari *et al.*²⁸ Considering that MNPs of different sizes and material properties are modelled, our maximum value of α_{ME} is about $0.1 \text{ V Oe}^{-1} \cdot \text{cm}^{-1}$, registered at 300 mT, while they reported a peak of $0.4 \text{ V Oe}^{-1} \cdot \text{cm}^{-1}$ corresponding to an external H_{DC} of 600 mT. It is therefore possible to conclude that the MNP performance does not necessarily improve as the potential difference increases, but it rather diminishes at the expense of a progressively higher magnetic field.

By tuning the magnetoelectric coupling, a wide range of diagnostic and therapeutic applications based on MNPs can be explored. In this context, ME properties can be exploited to wirelessly and selectively manipulate cells and even reversibly permeabilize their membrane to accomplish targeted drug delivery. Efficient permeabilization of the membrane is conventionally achieved by following protocols based on the use of electric fields (*i.e.* electroporation), which generally rely on bulk electrodes placed in the proximity of cells and/or tissues. Plate electrodes for non-invasive stimulation or needle electrodes inserted into the tissue for invasive stimulation are two types of traditional devices.^{39,40} Invasive stimulation is usually the most effective strategy for electroporation, but it suffers from drawbacks common to all invasive procedures (complications can include bleeding, inflammatory responses and infections or systemic issues). These latter hinder the use of such a technique for broad-spectrum treatments or when the localization of the diseased area might be difficult to reach (e.g., cardiovascular applications). Recently, some studies have focused on the use of inorganic nanoparticles, such as gold nanoparticles or MNPs, as a completely contactless method to



achieve electroporation with enhanced spatial specificity at the nanoscale.^{42,64} In particular, in the very recent work of Bryant *et al.*,⁶⁵ MENPs have been explored as nanotheranostic agents for specific targeting of cancer cells and damaging *via* irreversible electroporation (IRE) by means of a high amplitude DC magnetic field (on the order of magnitude of 7T). From the simulations carried out in our study (Fig. 6 and Table 5), the electric field values considered suitable for reversible nano-electroporation (*i.e.*, ranging from 2×10^4 to 5×10^4 V m⁻¹, according to Lv *et al.*⁴¹ which we have taken as reference despite referring to bulky electrode-mediated pulsed stimulation) are reached when a H_{DC} of 300 mT is used to excite the MENP. This corresponds to the optimal performance scenario where the maximum efficiency (α_{ME}) is observed. In contrast, when the induced electric field exceeds 5×10^4 V m⁻¹ (corresponding to a H_{DC} of 650 mT), cell death caused by irreversible nano-electroporation has been ascribed to several reasons,⁶⁶ such as permanent membrane lysis, loss of cell homeostasis, flush in/out of molecules, and even cytotoxic effects as a consequence of electrolysis and by-products generated around the electrodes.⁶⁶

Having verified the occurrence of reversible nano-electroporation in the presence of an external 300 mT H_{DC} , we proceeded with a time-dependent static simulation. The MENP, which exhibits its non-superparamagnetic properties at room temperature,²⁴ presents a residual magnetic flux density⁴⁸ when a H_{DC} of 300 mT is applied for $\Delta t = t_{Rev-Nep}$ and subsequently removed (Fig. 7a). To further investigate the potential of MENPs not only as wireless permeabilizing nanodevices of biological membranes but also as emerging tools for the controlled release of therapeutic agents into specific tissues, we focused our investigation on the alternating magnetic field simulation. Ideally, it has been previously postulated that the application of a H_{AC} onto a magnetized nanoparticle decorated with drug molecules would create conditions for an efficient drug-MENP bond breaking process, thus enhancing the drug release efficacy.⁴⁴ Such phenomenon is influenced by several factors, either related to the physico-chemical features of the system (*i.e.*, organic coating on the MENPs, nature of the drug-MENP bond, and type and amount of drug loaded), or the conditions of magnetic stimulation (duration, amplitude and frequency of H_{AC} and possible temperature increase during the stimulation).^{43,67} In this preliminary computational study, the main parameter considered was the variation of the H_{AC} amplitudes (values in the range of the experimental published literature^{43,67}), from which the core magnetization and the subsequent electric potential difference at the MENP surface are dependent. In line with the work of Kozielski *et al.*,⁶⁸ where it is affirmed that applying a sinusoidal magnetic field to magnetoelectric materials produces a sinusoidal electric field with a frequency that matches the input, Fig. 7e and f display comparable trends for the core magnetization and the electric potential difference profiles. As a consequence of the ME effect, the electric dipole generated at the MENP outer border induces a charge surface density (defined as σ_{MENP}) of the order of magnitude of $\pm\alpha H$ (following the theoretical dissertation of Stimphil *et al.*,⁵⁵ eqn (18) of the SI). Assuming a hypothetical

drug molecule attached at the MENP outer shell *via* an ionic bond, its release would occur if such σ_{MENP} equals and/or exceeds the charge density involved in the drug-MENP ionic bond. Moreover, fundamental to the dissolution of such a hypothetical ionic bond is the effective alternation of stretching and contracting of the ionic bond length, which ultimately results from the movement of positive and negative charges that constitute the ionic bond itself from their equilibrium positions. Considering that multiple drug molecules may be bound to a single MENP surface, an average positive/negative charge displacement can be reasonably defined (labelled as Q_{ionic} , according to the theoretical modelling of nanoparticle-bound drug release by Stimphil *et al.*,⁵⁵ eqn (19)–(22) of the SI). Our simulations conducted under varying H_{AC} conditions (Fig. 8b) highlight that such an average charge displacement in the hypothetical ionic bond is indeed governed by a linear dependence on the electrical potential difference (ΔV) developed at the MENP surface. ΔV oscillations over time would promote Q_{ionic} variations, thus perturbing the equilibrium positions of the positive and negative charges. We speculate that the length of the hypothetical drug-MENP bonds would be far from their equilibrium when such oscillations become more readily detectable (*i.e.*, H_{AC} equal to or greater of 4.5 mT, Fig. 8c). Higher alternating magnetic field values would result in larger oscillation amplitudes, thereby spanning over a wider range of electrical potential differences at the MENP outer shell, which, in turn, would be responsible for a stronger variation in the average bond charge displacement. In other terms, such stimulation conditions would cause a strong ‘shaking effect’ on the system, favoring the delivery of the therapeutic agent.⁴³

4.1 Study limitations

To the best of our knowledge this work marks the first *in silico* assessment of the operational parameters (*i.e.*, H_{DC} and H_{AC} external stimulations, with varying field strengths and amplitudes) involved in MENP-mediated nano-electroporation and drug release. However, several factors that are known to contribute to both phenomena are not included in the herein computational model and data analysis. While our study focused on the MENP general electrical outputs (*i.e.*, E_{fields} in the MENP nearby and ΔV at its surface after remote activation *via* external magnetic fields) and how they set the conditions for MENP-biological structure interface events to occur, other aspects are neglected. In fact, our model is an approximate simplification of the MENP structure, that does not consider additional outer layers, such as organic coating linkers and drug molecules, which are expected to slightly modify the overall electrical behavior. Regarding the nano-electroporation process, membrane pore dynamics formation has not been considered at this stage of the investigation and any possible effect of the thermal contribution related to MENP stimulation is herein overlooked. In providing a high-level physical perspective on the feasibility of using DC electric fields generated by MENPs, the concepts of free moving charges in conductive media and Debye length have not been considered



in the present work. More detailed investigations will be addressed in future studies to investigate the electric phenomena at the MENP–cell membrane interface, taking into account the screening effects of free charge carriers on the expected cross membrane electric potential generated by the MENPs located within the Debye length. Such a limitation is also related to the computational challenge of modeling systems with highly dissimilar dimensions and spatial occupation, ranging from the nanometer sized features of the MENPs to hundreds of micrometers of extension of cell membranes. Lastly, future studies more specifically centered on the modeling of the MENP–therapeutic agent bond are needed, to correlate both the DC and AC magnetic stimulation with the drug delivery process in terms of bond nature, the amount of drug loaded, and the inorganic particle/organic molecule assembly stability.

5. Conclusion

Today, MENPs are at the forefront in the development of innovative, wirelessly powered biomedical devices to actively interact with cells and tissues. Based on a spherical MENP model previously validated by our group, the current work aims to explore the use of MENPs for reversible nano-electroporation and drug release. The first part of our step-by-step analysis showed how to best exploit the operating principle – maximizing α_{ME} – so that the induced electric field in the MENP nearby, following static external magnetic field activation, is within the range of a reversible nano-electroporation, not harmful for the cells. The second part focused on the analysis of the drug release phenomenon, shedding some light on the reciprocal dependence of the magnetoelectric effect and the subsequent changes in the electric potential on the charge displacement. The preliminary concept herein presented constitutes a precursor to detailed analyses of gradually more complex systems, which will take into account the effects of organic coatings and multiple drug molecules on the overall MENP electrical behavior and on the charge displacement at the MENP–molecule interface. This will provide further knowledge to support the synthesis of drug-loaded MENPs and to define protocols for specific delivery systems adapted to cardiovascular surgery applications, in view of a progressively more personalized and less invasive medicine.

Author contributions

AT: investigation, visualization, data curation, writing – original draft; GS: conceptualization, visualization, and writing – original draft; SF, EC, and MP: data curation and methodology. SF, EC, MP, ML, AM, and BC: writing – review and editing; PR and EC: funding acquisition. MP, BC and PR: supervision. All authors contributed to manuscript revision and read and approved the submitted version.

Conflicts of interest

The authors declare no conflict of interest.

Data availability

The data supporting this article have been included as part of the SI. Data for this article, including material properties are taken from the COMSOL Multiphysics Library, available at <https://www.comsol.com>, while the tissue properties are taken from the IT'IS Foundation Tissue Database of the Sim4Life database, available at <https://itis.swiss/virtual-population/tissue-properties/database/dielectric-properties/>.

Supplementary information include the mathematical expressions for (a) the COMSOL multiphysics model of the magnetoelectric phenomenon; (b) the COMSOL multiphysics model of the magnetic hysteresis phenomenon; (c) the theoretical model of the drug delivery phenomenon; (d) magnetization loop of the cobalt ferrite core; (e) static study results to validate the MENP baseline behavior in the culture medium environment; (f) distributions of the electric field produced by various external H_{DC} fields within culture medium surroundings; (g) time dependent study for the drug release step in the CM environment; (h) MENP core magnetization, electric potential difference and magnetoelectric coefficient values in relation to the increase in amplitude of the applied H_{AC} for the CM case scenario. See DOI: <https://doi.org/10.1039/d5na00438a>.

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