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Irradiation-based design of mechanically resistant microstructures tuned via multiscale phase-field modeling

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We present a multi-scale phase field modeling of stationary microstructures produced under 1 MeV krypton ion irradiation in a phase separating concentrated solid solution of silver and copper. We show that the mixture reaches ultimately a stationary micro-structural state made of phase domains with composition and size distribution mapped to the values of the incident flux of particles and of the temperature, variables that help defining a non equilibrium phase-diagram for the irradiated alloy. The modeling predicts the formation of diverse microstructures likely connected to spinodal hardening, thus opening the perspective of the on-purpose tuning of mechanically resistant microstructures and the preparation of metastable alloys with mechanical properties improved by comparison to counterparts obtained via classical thermo-mechanical treatments.

I. INTRODUCTION

Patterned microstructures forming in driven materials have been for long a hot subject of applicative research since it is foreseen that thereby new preparation routes can emerge for industrial materials with desired physical properties, surpassing these obtained via classical preparation routes [1, 2]. Stationary irradiation microstructures are made of phase domains with composition and size distribution controlled by the temperature, the flux and the nominal composition of the alloy. Under irradiation at finite temperatures, these have been shown to stem from competing mass transfer processes and from the large difference existing between the characteristic length scales of the associated non local forces [3, 4]. Examples of irradiation microstructures are periodic defect walls [5], bubble or void lattices [6, 7] and periodic composition modulations in concentrated alloys [8].

Early suggestions for micro-structural engineering targeting the preparation of mechanically resistant materials were based on the known benefit of homogenising an initially heterogeneous microstructure via ion-beam mixing [9–12]. A challenging step further is to requesting the preparation of homogeneous microstructures with tuned composition, size distribution of phases and/or defect lattices. This is a critical issue for microelectronics, nuclear energy and other applications. Indeed, materials for microelectronics should combine electronic properties with mechanical strength [13, 14] whereas duplex stainless steels for nuclear applications should resist hardening and embrittlement due to the δ-ferrite phase decomposing under neutron irradiation (spinodal decomposition) [15–17], possibly related to spinodal hardening increasing the yield stress [8, 18, 19]. In this context, the present study aims at providing a robust methodology for linking the external parameters of irradiation, such as the temperature, the irradiation flux and the energy of incident ions, to the resulting change in the yield stress of the irradiated alloy. To this end, the phase-field method is here adopted [20, 21], referred hereafter to as PF, for it efficiently integrates the very different space and time scales of the processes leading to patterning in irradiated materials and has been shown capable of reproducing the experimental irradiation microstructures [22]. It is worth noting that the present study has been considerably simplified by previous works from the literature, having explored this subject either, via a similar methodology or by using the Kinetic Monte-Carlo method (KMC) [23–29]. These works have shown that experimental facts, such as the formation of cherry pits under irradiation, are qualitatively reproduced [30], without attempting any quantitative comparison of their findings with experimental observations of the size distribution and the composition of stationary phase domains. Moreover, little is known about the mechanical strength of the irradiated alloy and the variety of microstructures resulting from changes in the experimental parameters such as, the nominal composition, the temperature, the flux and the energy of incident ions, although this knowledge is the prerequisite for effectively operating microstructural engineering [31].

Unlike others, the present work shows that at any value of the nominal composition of a binary alloy, stationary irradiation microstructures are precisely located

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within a pseudo-phase diagram spanned by the temperature and the irradiation flux, which knowledge allows for identifying the regions triggering spinodal hardening, while enabling the flexible tailoring of microstructural features connected with the macroscopic mechanical behavior. For the illustration of the methodology developed in the present study, the case is considered of a concentrated random solution of silver (Ag) and copper (Cu) under 1 Mev krypton (Kr) ion irradiation. This alloy possess a large miscibility gap, has been extensively studied in the literature and is well suited for X-ray spectroscopy investigations of the microstructure since the difference between Ag and Cu absorption factors facilitates identifying their distributions [32].

In the following, the previously developed PF model of the alloy under ion irradiation is briefly presented together with the parameterisation of the free-energy functional, appropriately chosen to reproduce the experimental phase diagram at equilibrium [33, 34]. Under ion irradiation, the calculated pseudo-phase diagram is then presented together with the types of emerging stationary states at various temperature and ion flux values. Therefrom, the experimental conditions likely to yield the mechanical behavior needed for operating the microstructural design are selected. Moreover, the size distribution, the composition and the periodicity of precipitates are given in experimentally accessible units as functions of the temperature and the irradiation flux. Finally, an explicit connexion is established between microstructures and spinodal hardening.

II. QUANTITATIVE PHASE-FIELD MODEL

A. Phase-field model under irradiation

Under ion irradiation, the evolution of a binary solution is customarily described at the mesoscopic scale via the kinetic equation [33]:

\[ \frac{\partial c}{\partial t} = M \nabla^2 \left[ \frac{\delta \mathcal{L}(c)}{\delta c} \right], \tag{1} \]

where \( c(r,t) \) represents the coarse-grained concentration of species, \( M \), is the atomic mobility and, \( \mathcal{L}(c) \), is a nonequilibrium effective free energy defined by:

\[ \mathcal{L}(c) = \int_{\Omega} \left[ f_h(c) + \frac{\kappa}{2} |\nabla c|^2 + \frac{\Gamma}{2M} \{ g \ast c \} (r,t)c(r,t) \right] d\Omega. \tag{2} \]

In this equation, \( f_h(c) = a_2(T)(c - c_L)^2/2 + a_3(T)(c - c_L)^3/3 + a_4(T)(c - c_L)^4/4 \) is the Landau polynomial expansion of the bulk free energy near the concentration, \( c_L \), in the liquid. Unlike the standard Cahn-Hilliard equation [35], the presence in, \( f_h \), of the cubic term allows modeling of both, first and second order phase transitions [36]. However, this expression of the free-energy density neglects elastic effects relating to the large difference between effective diameters of Ag and Cu atoms.

The term \( \kappa |\nabla c|^2/2 \), referred hereafter to as the Ginzburg term [20], expresses the additional energy associated with emerging chemical interfaces during phase separation, with stiffness coefficient, \( \kappa \). The last term in \( \mathcal{L} \) represents the impact of compositional changes due to the relocation of species during irradiation and is therefore temperature independent [37]. The Green function, \( g \), satisfies \(-\nabla^2 g = \delta - p_R \), where, \( p_R \), is the probability distribution of displacements of species occurring in sub-cascades with relocation frequency, \( \Gamma [23] \) and \( \delta \), represents the Dirac impulse. This frequency, is usually taken as position independent within the reasonable approximation that cascades homogeneously overlap and that the volume fraction of cascades covered by sub-cascades exceeds the percolation threshold [38]. Since collision cascades last a hundred of picoseconds, whereas patterns evolve slowly at the time scale of diffusion, the microstructure is sensitive only to the time-averaged ballistic effects of cascades. Accordingly, \( \Gamma = \sigma^* \Phi \), where \( \Phi \) is the irradiation flux, and \( \sigma^* \), is the effective cross-section of relocation events characterizing the efficiency of ion irradiation to inducing ballistic relocations. Under the same assumptions, the probability distribution, \( p_R \), is a one variable function of the relocation distance, \( r \), customarily taken as a decreasing exponential with decay constant, the average relocation distance, \( R \). Equation 1 is easily put in adimensional form with characterist time and length constants, \( t_0 = \kappa/\alpha V_{at}^4 \), and \( l_0 = \sqrt{\kappa/\alpha V_{at}} \), where \( \alpha \simeq 0.5 \) at low temperatures, and \( V_{at} \) is the average atomic volume in the alloy. PF simulations consisted in solving equation 1 in reduced coordinates by means of the Spectral-Eyre (SE) scheme [39] on a 1000 \times 1000 grid, with reduced space and time steps, \( \Delta x = 0.5 \) and \( \Delta t = 0.1 \). PF simulations have been performed in two dimensions, assuming that no determinant insight might be added by 3D simulations. However, implementing the 3D PF procedure from the existing 2D model presents no major challenge. Afterwards, the conversion of the results in real units is made with values of the above parameters representative of AgCu under 1 MeV krypton ion irradiation.

B. Phase-field parametrization

In this section, the parametrization procedure of the adimensional PF model is recalled. For further details and numerics, are given in [33].

1. Bulk free energy

PF modeling of AgCu at equilibrium, consisted in numerically determining the coefficients of the bulk free energy density, \( f_h(c) \), such as to fit the experimental phase diagram of the alloy [34, 40]. To this end the stationary
numerical solutions of equation 1 have been determined at several concentration and temperature values along the solubility limit [33].

2. Interface and Ginzburg term

The energetic cost of diffuse interfaces forming in the alloy during phase separation is controlled by the stiffness coefficient, $\kappa$, which value is not available from experiments (see equation 2). In substitution, its value has been estimated from fits of PF composition profiles across a (100) semi-coherent interface between crystalline Ag and Cu on to counterparts obtained via Grand Canonical Monte Carlo (GCMC) simulations of this heterophase interface. This last has been taken as a convenient average representative of interfaces in the phase-separated alloy in which phases at equilibrium are expectedly bounded by the following low-index interfaces sorted in ascending excess energy order, (111), (100) and (110). GCMC calculations at different temperatures included relaxations of atom positions, of the system volume and of the overall composition at fixed the difference between chemical potentials of the two species, $\delta \mu$. The each time temperature dependent $\delta \mu$ has been set to the value driving a bulk AgCu solid solution to the solubility limit [41]. Upon convergence, interfacial and bulk regions reach the respective equilibrium compositions, revealing that the concentration of species at the interface is practically temperature independent, whereas bulk regions in this bi-crystalline model evolve toward the experimental solubility limits at each temperature. This finding shows that interfacial composition profiles are tightly related to the segregation of species, weakly changing on increasing the temperature, which justifies using as an input for the PF simulations the constant value, $\kappa = 1.5 \text{ eV.nm}^{-1}$ obtained as indicated above [33, 41]. An additional approximation is made by maintaining the interfacial stiffness under irradiation at the same value, which is equivalent to admit that effects of thermal segregation of species at interfaces and species relocation under irradiation superimpose linearly. In the PF modeling, this value of $\kappa$ has served for fixing the spatial correlation length of the coarse-grained compositional field, $\xi$, making this procedure the pivot connecting the scale of space of GCMC simulations (atomic) to that of PF simulations (mesoscopic). This is the ground of the multiscale modeling of phase separation phenomena investigated in the present work. In this context, it is worth noting that within the PF framework, $\xi \simeq \sqrt{\kappa/\sigma_2}$, yielding, $\xi \sim 0.2 \text{ nm}$, a length comparable to the atomic spacing in AgCu much inferior to the correlation length of concentration fluctuations allowed by the Landau-Ginzburg phenomenology. Though singular, this result is the price to pay for fitting the composition values of the atomic scale modelling, in the bulk and at the interface, via the continuous and derivable functions solving the PF equations. Thereby, atomic modelling and PF lead to compositions in bulk phases identical to these obtained in the experiment. However, the quality of the fit does not attest for applicability of the PF modelling at the atom scale.

A final remark is about the deliberate choice made in this work, to consider that interfaces between phases are semi-coherent, which is justified given the mismatch between Ag and Cu lattices, but is certainly not true at the very beginning of the alloy decomposition and could not apply to states close to equilibrium, where incoherent relationships between phases may exist. However, the work focuses on stationary phases under irradiation (disrupted growth) that can be reasonably considered in a semi-coherent relationship whereas no experimental evidence has been found in the literature reporting on observations of incoherent precipitation in the range of temperatures and irradiation flux values used in the present study.

3. The irradiation term

Displacement cascades have been simulated within the Binary Collision Approximation (BCA) via the MARLOWE package [42]. Although displacement cascades thereby investigated do not account for atom vibrations and multiple collisions, such simulations have proven yielding on average a realistic description of atom relocation in the ranges of PKA energy and irradiation flux covered by ion mixing experiments in the ballistic regime. In particular, estimations of the effective cross-section of relocation events, $\sigma^*$, and the ballistic displacement probability, $p_R$, have proven consistent with MD simulations of copper irradiation in the range of energies of Primary Knock-on Atoms (PKA) accessible by this technique. The proof has also been given that the relocation range, $R$, is ion energy and flux independent, involving exclusively the chemical characters of incident ions and of the atom species present in the target. The detailed justification of the irradiation model used in this work can be found in [43].

4. Equilibrium and irradiation mobilities

Under irradiation, the atomic mobility, $M$, has been evaluated via an hybrid numerical and analytical method embodying both the mobility terms, at thermal equilibrium and under irradiation where additional diffusion relates to the increase of point defect concentrations. At low and intermediate temperatures, the first term is predominantly vacancy-controlled and, since values of vacancy migration energies in Ag and Cu are close one to each other, has been conveniently approximated by the equilibrium mobility of vacancies in copper. For evaluating the second term, the average number of Frenkel pairs produced in displacement cascades has been obtained by using the MARLOWE package without recombinations. Since the time scales of point defect migration and of phase growth are very different, stationary irradiation
defect concentrations have been estimated by using the chemical model of Sizmann, in the recombination regime, assuming that recombination is exclusively taking place at the interfaces between phases [44]. Elimination of defects at sinks consists in this model in a linear relation between recombination and the respective concentrations of point defects with the proportionality coefficient relating to the average size of precipitates [45]. The method would require downscaling of microstructural data into the point defect model yet, due to the symmetry of Sizmann’s model, this coefficient does not appear in the stationary expression of defect concentration, which considerably simplifies the evaluation of defect concentrations under irradiation. Finally, it is worth noting that due to the large bias in the mobilities of vacancies and interstitials, the last eliminate rapidly at phase limits and volume diffusion is predominantly vacancy mediated.

III. RESULTS

A. Stationary state microstructures and patterning

Under irradiation, the forces driving the evolution of the alloy originate from ordering and disordering processes. Ordering proceeds by means of vacancy-mediated bulk diffusion and drives the alloy toward phase separation, because of the repulsive interactions between atom species. Disordering is caused by the ballistic displacements occurring within the collision cascades and tends homogenizing the alloy locally. It has been established that under the parallel action of the two processes non equilibrium stationary states can emerge when ordering and disordering come to balance one another. Example is given by the the dissolution of precipitates occurring under irradiation whenever the disordering rate overcomes that of thermal coarsening [24, 25]. In the following, attention is given to patterning, that is the emergence of stationary, periodic distributions of precipitates in a decomposing alloy.

1. Stationary state criterion

In decomposing alloys, the thermal coarsening of precipitates is disrupted by ion irradiation and leads to a variety of stationary microstructures that differ from these emerging under thermodynamical equilibrium conditions [25]. These out of equilibrium states correspond to the minima of the Lyapunov function, \( L \), embodying thermodynamical and ballistic effects (equation 2, figure 1). This legitimates considering this function as an effective free energy of the system and the microstructure emerging upon convergence to a minimum as the signature of the corresponding stationary state \( t \gtrsim 560 \text{s} \) in figure 1.

2. Flux and temperature as controlling parameters of stationary microstructures

Demange et al. [33] have shown recently that the temperature and the irradiation flux are the parameters controlling exclusively the kind of stationary microstructures forming under ion irradiation. Thereby, a pseudo-phase diagram can be drawn in the temperature-flux plane revealing that all the possible stationary microstructures are organised within domains delimited by frontiers, which crossing would trigger microstructural transitions (figures 2-3). At low temperatures and high ion fluxes, where ballistic disordering predominates, the stationary state is a fully disordered solid solution, whereas high temperature and low ion flux values trigger phase separation since, in this case, ballistic effects do not counterbalance thermal diffusion. At intermediate flux and temperature values, coarsening is disrupted and a variety of microstructures are expected forming. In all these situations, the competition between irradiation and diffusion is the controlling factor as is shown by molecular statics studies where, in absence of thermal diffusion, the irradiation-induced mass transport triggers the formation of complex ordered structures. Previous diffraction studies by Wei at al. [26] of the patterned microstructures forming in a concentrated AgCu alloy submitted to 1 MeV krypton ion irradiation reveal consistent with the above statements.

3. Stationary microstructures in the patterning domain

With the flux values given in figure 2 and temperatures lying within the patterning domain, ranging from 271 K to 366 K, the various stationary microstructures forming
are displayed in figure 3. Disk-like phases are precipitates made principally of silver within a copper matrix represented by the background carrying a different contrast [24, 25, 33]. Moreover, this figure shows that at the lower temperatures (figures 4(a), 4(d) and 4(g)), phase nanoparticles are practically mono-disperse in diameters, which provides a basis suitable for manufacturing materials with the desired mechanical response, provided the correspondence is established between the topology of the microstructure and the appropriate mechanical observable. On increasing the temperature, the dispersion of precipitate diameters increases as does the distance between them (figures 4(c), 4(f) and 4(i)). In total, this behavior is compatible with the theoretical and numerical investigations by Chen and Khachaturyan [46] and by Abraham [47].

B. Microstructural design

1. Size of precipitates

The qualitative topological trends of stationary microstructures illustrated in figure 3 have also been quantitatively studied as functions of the flux and of the temperature by computing the evolution of $n = 200$ initial random configurations of the solid solution at the critical composition, $c_{Ag} = 0.39$. Thereby, statistically meaningful averages of stationary particle diameters and composition fields have been obtained (figure 4). The former can be conveniently fitted by gaussians in confirmation of the statement made above that, at low temperatures, the size distributions of particles are mono-disperse. Such microstructures are fully characterized by specifying the average values of the composition and of the diameter of precipitates. At higher temperatures, by comparison to the case above, diameter distributions are enlarged, asymmetric, and heavily tailed, indicating that the relative proportion of the large precipitates increases on increasing the temperature (figures 3 and 4, right column). Considered all together, these results suggest that, under ion irradiation, composition modulations amplifying to form stationary microstructures extend over a pretty narrow range of wavelengths in difference with radiation-induced dissolution of precipitates during annealing, where coarsening is disrupted above a critical size value [48, 49]. Thus, ion irradiation reveals well adapted to tailoring of quasi-mono-disperse distributions of precipitates as needed for microstructural design.

In complement to figure 4, figure 5 displays the mean diameter of stationary precipitates as a function of the temperature and the ion flux. It can be seen that the values are nanometric, amounting about twice the interfacial thickness at the frontier of phases (Cf section II B 2, [33]), which indicates that the predicted microstructures are closer to the expected stationary modulations of the composition in early stages of the spinodal decomposition.
than to an assembly of bulk phases. Interestingly, the conclusions of atom probe tomography observations in thermally aged stainless steels show that Cr precipitates as small as 0.8 nm form in the ferrite phase [50, 51], akin to these predicted by phase-field simulations of aging effects in a CuNi alloy [52]. KMC simulations from the literature provide qualitative support to the findings listed above and to the trends illustrated by figure 5 showing that the size of precipitates increases on increasing the temperature or by decreasing the irradiation flux [28]. Finally, the inset in this figure shows that the dispersion of precipitate size is thermally activated, a behavior tightly relating to the thermal mobility contribution underlying the phase separation.

2. Composition profiles, amplitudes and periodicity

The phase-field simulations performed in this work yield also informations about the composition and the periodicity of stationary precipitates forming under ion irradiation, which are expected closely relating to the mechanical behavior of the alloy [53]. Figure 6 displays the profiles of Ag concentration as a functions of the temperature at flux value, $\Phi = 10^{13} \text{ cm}^{-2}\text{s}^{-2}$. It appears that the Ag content within precipitates exceeds by far the solubility limit at equilibrium of the bulk alloy [34], whereas the concentration profiles adopt a nearly sinusoidal shape at low temperatures. At high temperatures, the figure indicates the formation of strongly segregated interfaces with Ag concentration close to that predicted for (100) interfaces (Cf section II B 2, [33]). This is in agreement with Monte-Carlo simulations, showing that under irradiation the temperature is the controlling parameter for the position dependent composition across an (100) interface, moving from segregated toward sinusoidal modulation profiles [8]. Moreover, on the experimental side, this behavior is reminiscent of the observations of intermixing produced during irradiation by energetic ions [54] and matches the observations made by Atom Probe Tomography (APT) which showed the formation of quasi-periodical profiles of composition in Ag-40Cu after aging at 473 K [55].

Computed composition amplitudes, $\bar{A}$, and modulation wavelengths, $\lambda$, are respectively displayed as functions of the temperature and ion flux values in figures 7(a) and 7(b). Amplitude maximum values reveal much larger than observed in after annealing studies of spinodal decomposition in AgCu, $\bar{A} = 0.3$ [55]. In addition, the wavelengths of stationary composition modulations investigated in the present work are much smaller the values obtained in studies of thermal aging in alloys [56–59]. It is therefore concluded that ion irradiation yields stationary precipitates with enhanced compositions much finely distributed than these produced by thermal annealing, adding thereby better tuning flexibility to microstructural design.
C. From the irradiation induced microstructure to spinodal hardening

Precipitation in the spinodal decomposition domain changes the mechanical behavior of alloys through the interaction of mobile dislocations with precipitates impeding their motion. This “spinodal hardening” contributes differently to the material strength depending on the degree of advancement of the precipitation reaction: in the early stages of precipitation significant hardening is observed, which is followed by a saturation regime and, sometimes, a strength decrease with further evolving the microstructure [60–63]. Despite longstanding efforts, no widely accepted global modeling of this behavior has emerged as is attested by the numerous models proposed in the literature (Mott & Nabarro [64], Cahn [53], Ardell [65], Friedel [66], Ghista & Nix [67] etc.). At least, Cahn’s theory of spinodal hardening convincingly describes the evolution of the mechanical property in the early stages of precipitation. In this model, applied to cubic structures deforming via dislocation glide on (111)[110] slip systems, the Critical Resolved Shear Stress (CRSS), $\tau$, is shown proportional to $\bar{A}^2\bar{\lambda}$, where $\bar{A}$ and $\bar{\lambda}$ represent the average values of the amplitude and wave length of composition fluctuations computed in figure 7. Upon increasing the size of precipitates, dislocations overcome the obstacles via the Orowan mechanism and the CRSS decay is predicted proportional to $\bar{A}^{1/3}\bar{\lambda}^{-2/3}$. Experimental confirmation of the model has been provided by Schwartz & Plewes [68], who observed that the CRSS increase of a Cu-9Ni-6Sn alloy undergoing spinodal decomposition is strictly proportional to $\bar{A}^2\bar{\lambda}$, whereas the model does not reveal capable of satisfactorily describing the late stages of the microstructure evolution. Similar findings have been reported on Cu-5at%Ti by Miyazaki et al. [69] and by others for various systems [70, 71]. It therefore appears that Cahn’s theory applies strictly to the periodic microstructures emerging during the early stages of spinodal decomposition, made of sinusoidal composition modulations, not to their long term evolution.

A main finding of the present study is that composition variations in the decomposing alloy are nearly sinusoidal in the patterning regime (low temperatures), which strongly suggests that Cahn’s theory can be used for evaluating the CRSS associated to such patterned microstructures. Accordingly, the expression of the CRSS $\tau$ is given by:
In this study, a previously developed phase-field multiscale modeling [33] has been employed for connecting the irradiation conditions, temperature and irradiation flux, to the mechanical behavior of stationary irradiation microstructures in a AgCu alloy. Stationary irradiation microstructures of two different types are found namely, at low temperatures, patterning consisting of periodic composition modulations forming closely packed diffuse particles of nanometric size and a disordered collection of larger precipitates with strongly segregated interfaces emerging at high temperatures. The mechanical response of the former type of microstructures is obtained by applying the model of spinodal hardening proposed by Cahn [53]. The mechanical behavior of the latter microstructures, is still an open question. Ongoing work focuses on short-range order between precipitates in such microstructures, which might constitute the factor controlling the mechanical response of the alloy [47, 72].

IV. CONCLUSIONS

In qualitative agreement with experiments, the graphs representing the evolution of $\tau$ with the temperature are shifted toward the higher temperatures on increasing the irradiation flux. However, it must be emphasized here on that only the contribution of the spinodal decomposition to the strength has been considered and that other causes may further harden the alloy as would expectedly happen when the dislocation density increases in response to a flux increase. This adds to the possible failure at high temperatures of the CRSS prediction via equation 3 since the concentration profile is not sinusoidal in shape anymore as is shown in the present work. Nonetheless, the above results constitute a first attempt relating PF predicted mesoscopic microstructures to the mechanical property. Further effort is required for satisfactorily modelling the evolution of the CRSS under irradiation and for clarifying the origins of hardening regimes observed in irradiated alloys.

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AUTHORS CONTRIBUTION

David Simeone was the one who suggested this study, and the resulting article. Gilles Demange, Laurence Lunéville and David Simeone elaborated the phase field model rooting the study. Gilles Demange performed the phase-field simulations and the BCA simulations. Vassilis Pontikis and Laurence Lunéville developed the MD/MC procedures and ran the corresponding simulations. Sylvain Dépinoy took care of the mechanics aspect of the study. Gilles Demange, Vassilis Pontikis and Sylvain Dépinoy took part in the redaction of the study. Gilles Demange, Laurence Lunéville and David Simeone performed the phase-field simulations and the BCA simulations. Vassilis Pontikis and Laurence Lunéville developed the MD/MC procedures and ran the corresponding simulations. Sylvain Dépinoy took care of the mechanics aspect of the study. Gilles Demange, Vassilis Pontikis and Sylvain Dépinoy took part in the redaction of the study.
manuscript: Sylvain Dépinoy wrote the mechanics section (III-C), Gilles Demange wrote the remaining part of the manuscript, and Vassilis Pontikis corrected and improved the intermediate versions of the manuscript.

**ADDITIONAL INFORMATION**

Declarations of interest: none.