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Using first-principles calculations to predict the mechanical properties of transmuted tungsten under first wall fusion power-plant conditions

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Abstract

Tungsten and tungsten alloys are being considered as leading candidates for structural and functional materials in future fusion energy devices. The most attractive properties of tungsten for the design of magnetic and inertial fusion energy reactors are its high melting point, high thermal conductivity, low sputtering yield and low long-term disposal radioactive footprint. Yet, despite these relevant features, tungsten also presents a very low fracture toughness, mostly associated with inter-granular failure and bulk plasticity, that limits its applications. Significant neutron-induced transmutation happens in these tungsten components during nuclear fusion reactions, creating transmutant elements including Re, Os and Ta. Density functional theory (DFT) calculations that allow the calculation of defect and solute energetics are critical to better understand the behavior and evolution of tungsten-based materials in a fusion energy environment. In this study, we present a novel computational approach to perform DFT calculations on transmuted materials. In particular, we predict elastic and plastic mechanical properties (such as bulk modulus, shear modulus, ductility parameter, etc) on a variety of W–X compositions that result when pure tungsten is exposed to the EU-DEMO fusion first wall conditions for ten years.

Keywords: first-principles calculations, plasma-facing materials, nuclear transmutation, tungsten, mechanical properties, stacking-fault energy

(Some figures may appear in colour only in the online journal)

1. Introduction

Tungsten and tungsten alloys are being considered as leading candidates for structural and functional plasma facing materials in future fusion energy devices. The most attractive prop-

erties of tungsten for the design of magnetic fusion energy reactors are its high melting point and thermal conductivity, low sputtering yield and low long-term disposal radioactive footprint. These advantages are accompanied unfortunately by a very low fracture toughness (mostly associated with inter-granular failure and bulk plasticity), low ductility at room temperature, and high ductile-to-brittle transition temperature, that limits its applications [1–5].

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A deep understanding of both elastic and plastic mechanical properties of tungsten alloys under first wall fusion power plant conditions is then a necessary step toward their consolidation as a viable option for this promising technology. Given the difficulty and the cost to perform experiments in such extreme environments, the use of computational modeling to provide insights and enrich the experimental knowledge of materials response has received much attention during the last decades. Among all the available techniques that allow for the prediction of mechanical properties across different temporal and spatial scales, density functional theory (DFT) has emerged as one of the most reliable ones to investigate the electronic structure of condensed matter systems [6].

In the elastic region, these first-principles calculations (also called *ab initio* calculations) have been broadly employed to study the elastic constants and the elastic properties of pure crystals and multiple alloys, starting in the 1970s and, particularly, in recent times [7–13].

For its part, plastic deformation in body-centered cubic (bcc) metals like tungsten is peculiar due to the existence of nonplanar dislocations with screw character and thermally-activated mobility [14–17]. This results in various kinds of complex phenomena such as pencil glide, asymmetry of the critical stress in the twinning and antitwining glide directions, asymmetry of the critical stress under tension/compression loading, or anomalous slip [18–30]. Among all the mechanical properties that can be determined in this plastic regime, ductility of tungsten and tungsten alloys has been the subject of much research and discussion over the last decades (both experimentally and computationally) given their well documented brittleness [5, 31]. Despite the scalability limitations of *ab initio* methods to characterize directly dislocation glide, grain boundaries, hardening, and other mesoscopic phenomena governing the plasticity of bcc metals, several approaches have proposed measures of ductility based on its relationship with material parameters that can be determined via DFT simulations. Going back to the 1950s, before computational modeling and *ab initio* calculations became effective methods to study the mechanical properties of materials, Pugh [32] formulated an empirical criterion that characterizes the ductility of materials by the ratio between the bulk modulus B and the shear modulus G . Cauchy pressure, defined in terms of the elastic constants, has also been employed to evaluate the ductility of metals since it includes the angular character of atomic bonding [33, 34]. Another criteria for isotropic polycrystalline materials is the Poisson's ratio, which is found to be proportional to the intrinsic ductility of crystals [35]. During analysis of dislocation nucleation from a crack tip, Rice [36] proposed that the ductile–brittle behavior of a material can be associated with the ratio between the surface energy and the so-called unstable stacking fault energy, a new solid state parameter identified in this analysis that represents the maximum energy encountered in the block-like sliding of a slip plane. Note that true ductility also depends on temperature and the loading strain rate; here we are calculating, via DFT, a measure that allows for an assessment of the relative variation in ductility as a material's composition changes.

Despite the numerous efforts in the literature to investigate the effects of alloying elements on various properties of tungsten such as phase stability [10, 37–40], elastic properties [10, 13, 37–39, 41, 42], ideal tensile strength [9, 10], ductility [43], radiation defects [38], point defects [37, 44–48], dislocation structure [16, 49], grain boundaries [50], etc, to the best of our knowledge there is a lack of understanding on how the expected first wall fusion power-plant conditions change the mechanical properties of these plasma facing structural materials over time. In this work, we perform first-principles calculations based on DFT to investigate the mechanical response of tungsten exposed to fusion-like environments. In particular, we focus on how the elastic constants, elastic properties, ideal tensile strength, and dislocation-based ductility parameter evolve as the composition of tungsten changes during the course of irradiation due to transmutation.

Our paper is organized as follows. After this introduction, we provide an overview of tungsten transmutation in a predicted fusion-reactor environment. This is followed by section 3, where the first-principles computational methods employed are presented. The results are given in section 4, which includes: (i) the validation exercise, with special focus on the elastic properties of pure tungsten; and (ii) the calculation of the elastic constants and elastic properties, the ideal tensile strength, and ductility parameter measurements for the chemical compositions that appear when pure tungsten transmutes during the course of irradiation. We finalize in section 5 with a brief discussion and the conclusions.

2. Tungsten transmutation

To characterize the transmutation properties of W in a fusion-like environment, calculations have been carried out using the FISPACT-II inventory code developed and maintained by the United Kingdom Atomic Energy Authority over the last 30 years. FISPACT-II [51, 52] solves coupled differential equations describing the rate of change of all possible nuclides and thus evolves a nuclide composition in time. In the simplest case, the differential equation describing the rate of change for one nuclide is:

$$\frac{dN_i}{dt} = \underbrace{-N_i(\lambda_i + \sigma_i\phi)}_{\text{depletion}} + \underbrace{\sum_{j \neq i} N_j(\lambda_{ji} + \sigma_{ji}\phi)}_{\text{creation}}, \quad (1)$$

where N_i is the number of atoms of nuclide i at a given time t . This differential equation is formed from two terms that represent the nuclide depletion (negative contribution) or creation (positive). The loss term, $-N_i(\lambda_i + \sigma_i\phi)$, includes λ_i as the decay constant of the nuclide (only non-zero if the nuclide is unstable), ϕ as the total neutron flux (in units of $\text{n cm}^{-2} \text{s}^{-1}$), and σ_i as the total collapsed cross section for all possible reactions on the nuclide (measured in barns and calculated by combining the energy-dependent specified neutron field with nuclear reaction cross-section data). For its part, the creation term consists of the sum over all other nuclides with the ji subscripts indicating the production of nuclide i from a reaction or decay on j . Single-value, total σ cross sections (in cm^{-2}

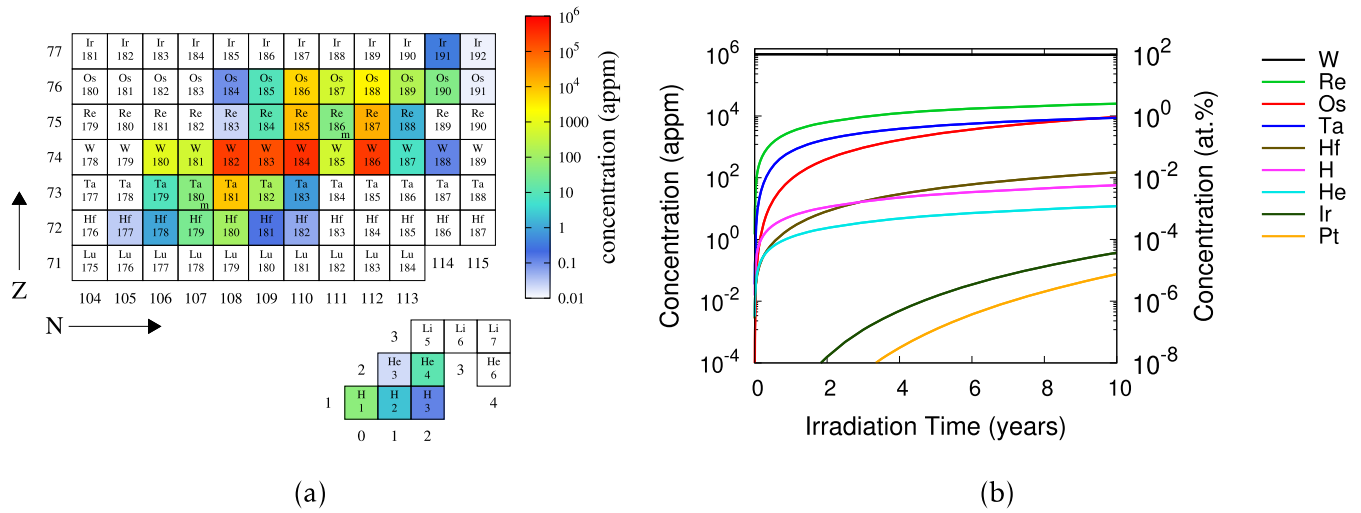


Figure 1. (a) Final nuclide composition of W after 10 years of continuous exposure to EU-DEMO first wall conditions. (b) Transmutation of W during a 10 years irradiation.

units) in equation (1) are obtained from the sum of the energy dependent cross sections values for the reaction weighted by the normalized neutron irradiation spectrum:

$$\sigma = \sum_n \sigma_n \frac{\phi_n}{\sum_m \phi_m}, \quad (2)$$

where the n, m sums are performed over the neutron flux vector.

For the present work, a ϕ vector of fluxes was taken from neutron transport calculations performed for a recent conceptual design (see [53, 54] for details) for EU-DEMO; a demonstration fusion power plant being researched in Europe [55, 56]. Specifically, the spectrum for the outer equatorial first wall of the torus-shaped tokamak has been used, which is predicted to be one of the highest flux regions of a fusion reactor (second only to the inner equator). The total flux ϕ for this spectrum was $2.1 \times 10^{14} \text{ n cm}^{-2} \text{ s}^{-1}$. A FISPACT-II calculation evolved initially pure W in this neutron environment for 10 continuous full-power years. Figure 1(a) shows the final composition of the material after those 10 years in a ‘nuclide map’ [57] tableau.

Note that 10 years of continuous operation is longer than that envisaged for first wall or divertor armour components in current fusion reactor concepts; for example, EU-DEMO operational scenarios consider around 5 years of pulsed operation lifetime for divertors—corresponding to ~ 2 full power years (fpy), while in the second, longer phase of EU-DEMO operation, blanket and first wall components are expected to experience about 14 years of pulsed operation (~ 6 fpy) [58]. In this respect, using the maximum transmutation results (at 10 years) presented here to analyze the mechanical impact on transmutation is an upper-bounding case relative to the end-of-life variations that might be expected in the next (first)

generation of fusion reactors. On the other hand, the lesson from the nuclear fission industry is that life extension of components is critical to the commercial viability of power plants, and thus 10 fpy may eventually be more representative of the lifetime requirements for the first wall armour of commercial reactors.

Figure 1(b) shows how the composition of W changed (transmuted) during the 10 years irradiation. The graph shows the concentrations, in atomic parts per million (appm), on a logarithmic scale of the elements created during the course of the irradiation. The concentrations of each element are the sum over all nuclides of that element (i.e. from one row of the nuclide map in figure 1(a) and table 4 in appendix A presents the numerical concentrations of the elements at 1 year intervals that were used in subsequent calculations.

Table 1 shows time-averaged concentration error estimates for each element, which are derived solely from the nuclear data uncertainties in the TENDL-2017 [59] libraries used by FISPACT-II to perform the calculations. FISPACT-II obtains errors for a given nuclide/isotope (e.g. ¹⁸⁶W, ¹⁸⁵Re or ²H) by summing (in quadrature) the nuclear data uncertainties on each reaction in the production chains of individual nuclides N_i of the inventory. Production chains are found using a tree search algorithm (see [51] for details of the method). The errors shown here for a transmutant element are the sums (again, in quadrature) over the predicted errors for each nuclide of that element at 1 year intervals (see table 4).

Even after 10 years and for the (relatively) highly-transmuting W, the material would still be more than 95% W. Notice that the profile of growth of each transmutant element varies—Os, for example is a secondary transmutant whose rate of production increases as the concentration of Re, a primary transmutant, increases.

Table 1. Time-averaged % errors in transmutant concentrations predicted by FISPACT-II for each transmutant element as W transmutes during the course of the 10 years power-plant first wall irradiation (see main text for details).

| | W | Re | Os | Ta | Hf | H | He | Ir | Pt |
|-----------|------|------|------|-------|-------|-------|-------|-------|-------|
| Error (%) | 0.13 | 6.47 | 7.54 | 36.14 | 31.29 | 12.65 | 42.44 | 31.14 | 37.83 |

3. First-principles computational methods

DFT calculations were performed by using the open-source software distribution QUANTUM ESPRESSO [60, 61]. The local-density approximation (LDA) with the Perdew–Burke–Ernzerhof (PBE) parametrization [62] was employed in the formulation of the exchange correlation functional. Optimized norm-conserving Vanderbilt pseudopotentials [63] compatible with the virtual crystal approximation (VCA) [64] method were chosen as the most suitable ones to study the variety of transmuted tungsten compositions that result from the exposure to the first wall conditions described in section 2. By using this VCA approach, the mixed pseudo-potential for any composition at a specific irradiation time in figure 1(b) can be generated by defining ‘virtual’ atoms at concentrations appropriate for the composition. The selection of the LDA with the PBE parameterization was justified given its compatibility with the VCA method used in this work. Future works will explore hybrid functionals to study transitional metal compounds.

The k -points were sampled using the Monkhorst–Pack method [65] by a shifted $30 \times 30 \times 30$ grid for the two-atom bcc supercell used in the calculation of the elastic constants and the ideal tensile strength. A shifted $28 \times 28 \times 1$ grid was selected for the 12-atom bcc supercell employed in the calculation of the stacking fault energies needed to obtain Rice’s ductility parameter. The planewave cutoff energies were ~ 2041 eV (150 Ry) for the elastic constants and the ideal tensile strength simulations and ~ 544 eV (40 Ry) for ductility simulations. The convergence tests to choose these values are provided in appendix B. Further details on how to extract the elastic and plastic properties of transmuted tungsten from the energies calculated via DFT simulations are given below.

3.1. Elastic constants and elastic properties

The change in the total energy per unit volume ($\Delta U/\Omega$) of a system subjected to a general deformation can be written, in the contracted *Voigt notation*, as

$$\frac{\Delta U}{\Omega} = \frac{1}{2} C_{ij} u_i u_j \quad \text{with} \quad u_i = \begin{cases} \epsilon_i & \text{if } i = 1, 2, 3 \\ 2\epsilon_i & \text{if } i = 4, 5, 6 \end{cases}, \quad (3)$$

where Ω is the volume and U is the total energy, respectively, of the crystalline unit cell, C_{ij} are the components of the stiffness matrix, and u_i are the engineering strain vectors. In the particular case of a cubic crystal such as bcc tungsten, the elastic tensor gets simplified due to the symmetries of the lattice [66],

i.e.:

$$C_{ij} = \begin{pmatrix} C_{11} & C_{12} & C_{12} & & & & & & & \\ C_{12} & C_{11} & C_{12} & & & & & & & \\ C_{12} & C_{12} & C_{11} & & & & & & & \\ & & & C_{44} & & & & & & \\ & & & & C_{44} & & & & & \\ & & & & & C_{44} & & & & \end{pmatrix} \quad (4)$$

where all the empty entries correspond to $C_{ij} = 0$. Then, equation (3) can be rewritten as

$$\frac{\Delta U}{\Omega} = \frac{1}{2} C_{11} (u_1^2 + u_2^2 + u_3^2) + C_{12} (u_1 u_2 + u_1 u_3 + u_2 u_3) + \frac{1}{2} C_{44} (u_4^2 + u_5^2 + u_6^2). \quad (5)$$

Following [67–69], the three non-zero elastic constants C_{11} , C_{12} , and C_{44} can then be obtained by evaluating this equation (5) under three deformations: isotropic ($u_1 = u_2 = u_3 = \eta, u_4 = u_5 = u_6 = 0$), tetragonal ($u_1 = u_2 = -\eta, u_3 = -2\eta, u_4 = u_5 = u_6 = 0$), and trigonal ($u_1 = u_2 = u_3 = 0, u_4 = u_5 = 0, u_6 = \eta$), where η denotes the distortion parameter.

Since C_{11} , C_{12} , and C_{44} constitute the entire set of elastic constants for a cubic system, other elastic properties of interest for single crystals can be extracted from these energy calculations. For example, the bulk modulus B and the tetragonal shear elastic constant C' can be determined as

$$B = \frac{1}{3} (C_{11} + 2C_{12}) \quad (6)$$

$$C' = \frac{C_{11} - C_{12}}{2}. \quad (7)$$

Additionally, mechanical properties of isotropic polycrystalline materials can also be determined from the already calculated elastic constants. By applying the Hill average [70], the shear modulus G can be calculated as

$$G = \frac{G_V + G_R}{2}, \quad (8)$$

where $G_V = \frac{3C_{44} + C_{11} - C_{12}}{5}$ and $G_R = \frac{5(C_{11} - C_{12})C_{44}}{4C_{44} + 3(C_{11} - C_{12})}$ are the Voigt and Reuss bounds, respectively. Then the Young’s modulus E and the Poisson’s ratio ν can be obtained as:

$$E = \frac{9BG}{3B + G} \quad (9)$$

$$\nu = \frac{3B - 2G}{2(3B + G)}. \quad (10)$$

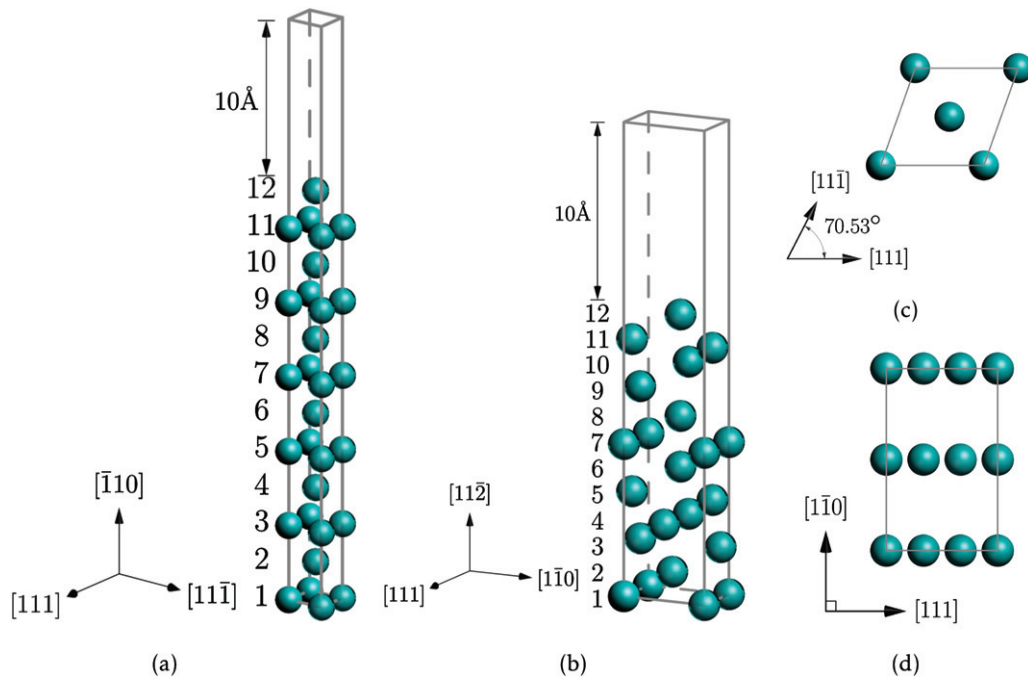


Figure 2. Atomic arrangement of the surface models used to calculate γ_{us} of transmuting tungsten on both (a) $\langle 111 \rangle \{ \bar{1}10 \}$ and (b) $\langle 111 \rangle \{ 11\bar{2} \}$ slip systems.

3.2. Ideal tensile strength

The ideal tensile strength of a material σ_m , also called theoretical strength, is defined as the maximum stress that the material can sustain.

For the uniaxial tensile test, the tensile stress σ relates to the total energy of the system U , the tensile strain ϵ , and the volume at a given strain $\Omega(\epsilon)$ as

$$\sigma = \frac{1}{\Omega(\epsilon)} \frac{\partial U}{\partial \epsilon}, \quad (11)$$

which allows for the calculation of the ideal tensile strength using first-principles simulations. To do so, the total energy for the unloaded material is firstly calculated. Second, a uniaxial tensile strain ϵ is imposed on the system, finding the minimum of the total energy by relaxing the atoms at the directions perpendicular to the loading axis. This process is repeated for different values of the strain (along the same loading direction) and the total energy is obtained as a function of the applied tensile strain. At each strain step (0.02 at the beginning and 0.01 as the results approach the region of the ideal strength), the force tolerance during the energy minimization was set to a value of $2.57 \times 10^{-3} \text{ eV } \text{\AA}^{-1}$ (10^{-4} Ry/bohr) for all five components except for the one where the uniaxial tensile strain is applied. Equation (11) is evaluated to calculate the stress σ for each particular strain, and the ideal tensile strength σ_m is obtained as the maximum in the stress-strain curve, which also relates to the inflection point in the total energy-strain curve [9]. Next, equation (11) is evaluated for each value of applied strain to determine the dependence of the stress σ with the strain. Finally, the ideal tensile strength σ_m is obtained as the maximum in the stress-strain

curve, which also relates to the inflection point in the total energy-strain curve [9].

3.3. Dislocation-based ductility parameter

According to the dislocation theory formulated by Rice [36], the brittle-ductile behavior of a crystalline solid can be characterized by the so-called ductility parameter D that accounts for the competition between the nucleation of dislocations from the crack tip and crack cleavage, i.e.:

$$D = \frac{\gamma_s}{\gamma_{us}}, \quad (12)$$

where γ_s is the surface energy and γ_{us} is the unstable stacking fault energy. In the present work, we focus on this Rice-criterion of ductility instead of the other three criteria described in section 1 since the generalized stacking fault energies (GSFE) that are needed in equation (12) can be determined directly by using first-principle simulations. Additionally, the calculation of the GSFE could also provide physical insights into the movement of dislocations, a mechanism that governs plastic deformation in bcc metals like tungsten [71–74].

The first step in the calculation of the GSFE is the definition of the geometry. A supercell containing twelve surface layers with a unit surface of 1×1 and 10 (\AA) vacuum is created as shown in figure 2. This is done for both $\langle 111 \rangle \{ \bar{1}10 \}$ and $\langle 111 \rangle \{ 11\bar{2} \}$, the two primary slip systems found in these bcc structures [75–78]. Next, a displacement along the $[111]$ direction is imposed on the upper half (7th to 12th layers) of the surface model while keeping the lower half (1st to 6th layers) fixed. After introducing the slip, the entire system is allowed

to relaxed along the z direction. The GSFE (also called γ -surface) is then defined as the energy cost per unit area of the cut incurred as a result of the shift, i.e.:

$$\gamma(x) = \frac{E(x) - E_0}{A}, \quad (13)$$

where x is the displacement along the shift vector (given as a fraction of the Burgers vector $\mathbf{b} = \frac{1}{2} [111]$), $E(x)$ is the total energy of the system after imposing a slip x , E_0 is the total energy of the surface model before the slip, and A is the area of the surface. The unstable stacking fault energy γ_{us} is obtained as the maximum value of the GSFE curve and it corresponds to the maximum energy that comes upon sliding the top half along a slip plane [36].

For its part, the surface energy γ_s is the energy required to create a new surface during the cleavage of an infinite crystal in two. Despite its importance to understand surface structure, reconstruction, roughening and crystals' equilibrium shape [79], there are some challenges to determine it both experimentally [80] and computationally [81]. In this work, we use the method proposed by Fiorentini and Methfessel [82] that calculates the surface energy as

$$\gamma_s \approx \frac{E_{\text{slab}}^N - NE_{\text{bulk}}}{2}, \quad (14)$$

where E_{slab}^N is the total energy of an N -layer slab and E_{bulk} is the bulk total energy. By using this approach, the divergence of the surface energy with the slab thickness is eliminated. Additionally, it does not require the calculation of the bulk energy term on a separate system since E_{bulk} can be taken as the slope of the total energy of the slab over the slab thickness.

4. Results

In this section, we present results of first-principles calculations to explore the dependence of the elastic constants, ideal tensile strength, and ductility-parameter on the time-dependent composition of transmuted tungsten under first wall fusion power-plant conditions.

4.1. Benchmarking—pure tungsten

As a preliminary step in our calculations, the elastic and plastic properties of pure tungsten were obtained and compared with previous works in the literature. Table 2 summarizes the lattice parameter, elastic constants and elastic properties from this work and previous studies. For their part, the values of the ideal tensile strength σ_m , the surface energy γ_s and unstable stacking fault energy γ_{us} are shown in table 3. These results are consistent with their counterparts from previous works (the relative differences are less than 15% in table 2 and less than 12% in table 3). Additionally, given the variety of methods compared, convergence studies were also performed on several properties of interest. The reader is referred to appendix B for more details about these convergence tests.

4.2. Elastic behavior of transmuted tungsten

With the confidence conferred by the benchmarking exercise of pure tungsten, next we proceed to calculate the elastic and plastic properties for a number of tungsten compositions that result during the first ten years of continuous exposure to EU-DEMO first wall conditions. The chemical compositions of transmuted tungsten shown in figure 1(b) and table 4 were used to generate the required pseudopotentials via the VCA method [64]. In addition to studying the current composition (i.e. all transmutants in table 4) at each irradiation time, two other scenarios were considered: (i) chemical composition at each irradiation time with tungsten and the primary transmutant (Re); (ii) chemical composition at each irradiation time with the top three transmutants (Re, Os, Ta). The purpose of investigating these two supplementary scenarios is to understand how the mechanical properties change as specific transmutants are added.

Figure 3 shows the time dependence of the lattice parameter and the elastic constants as the chemical composition of the tungsten-based materials change due to irradiation. This linear behavior has been confirmed in previous experimental [91] and computational [10, 49] measurements of W–Re alloys.

In figure 4 we provide detailed results on the evolution of elastic properties such as the bulk modulus B , the tetragonal shear elastic constant C' and mechanical properties of polycrystalline materials like the shear modulus G and the Young's modulus E . As described in section 3.1, the above properties can be obtained directly from relations between elastic constants, by applying the Hill average, or by imposing Voigt and Reuss bounds. The monotonic behavior observed in our calculations is also in agreement with previous computational works that studied W–Re alloys [10].

In advance of discussing these results and their implications in detail in the following section, we note the following features from the figures: (i) the lattice constant and C' are the only measurements decreasing as irradiation time (and therefore the relative concentration of transmutants) increases. All other properties monotonically increase with irradiation time; (ii) the magnitude of the slope when only the primary transmutant (Re) is considered, (no matter whether it is positive for some properties and negative for others) is always smaller than the one for the current composition with all transmutants or the scenario that includes the top three transmutants; and (iii) the effect of adding all remaining transmutants is negligible when compared with the impact of the chemical compositions that includes the top three transmutants (Re, Os, Ta).

4.3. Ideal tensile strength of transmuted tungsten

The stress–strain curve is one of the most fundamental tools to understand the mechanical behavior of materials. It reveals many of the elastic and plastic properties of interest, including but not limited to Young's modulus, yield strength, ultimate tensile strength, etc. Here we perform a systematic study of the ideal tensile strength of pure W, transmuted W (with

Table 2. Theoretical and experimental equilibrium lattice parameters a_0 , bulk modulus B , elastic constants C_{ij} , tetragonal shear modulus C' , shear modulus (G) and Young's modulus (E) of pure tungsten from our calculations and previous works.

| Software package, pseudopotential ^a | a_0 (Å) | C_{11} (GPa) | C_{12} (GPa) | C_{44} (GPa) | B (GPa) | C' (GPa) | G (GPa) | E (GPa) |
|--|-----------|------------------|------------------|----------------|--------------------|--------------------|--------------------|--------------------|
| EMTO, PAW [13] | 3.195 | 536.7 | 179.9 | 168.6 | 298.8 | 178.4 | 172.5 | 434.0 |
| QE, NCPP [49] | 3.187 | 499 ^b | 201 ^b | 149 | 300 | 160 | 154 ^b | 394 ^b |
| VASP, PAW [83] | 3.1755 | 529.94 | 211.19 | 140.59 | 317.44 | 159.38 | 148.11 | 384.52 |
| VASP, PAW [84] | 3.17 | 536.3 | 202.2 | 138.7 | 313.6 ^b | 167.1 ^b | 149.4 ^b | 386.8 ^b |
| QE, USPP [17] | 3.1903 | 518 | 197 | 141 | 304 | 160 | 160 ^b | 408 ^b |
| Exp. (0 K) [85] | 3.165 | 532.55 | 204.95 | 163.13 | 314.15 | 163.80 | 163.40 | 417.76 |
| Exp. (300 K) | — | 530.25 | 201.9 | 160.92 | 311.35 | 164.18 | 160.18 | 410.09 |
| QE, NCPP | 3.1835 | 512.501 | 197.906 | 142.461 | 302.771 | 157.298 | 148.22 | 382.669 |
| [this work] | [6%] | [5%] | [6%] | [15%] | [5%] | [12%] | [14%] | [12%] |

^aAcronyms used for software packages and pseudopotentials: exact muffin-tin orbitals (EMTO) formalism [86], projector augmented wave (PAW), norm conserving pseudopotentials (NCPP), ultra-soft pseudopotentials (USPP).

^bValues are obtained by substituting the elastic constants calculated via first-principles calculations into equations (6)–(9).

Table 3. Theoretical ideal tensile strength σ_m , corresponding strain ϵ_m , surface energy γ_s , and unstable stacking fault energy γ_{us} of pure W from our calculations and previous works.

| Software package, pseudopotential ^a | σ_m (GPa) | ϵ_m (%) | $\langle 111 \rangle \{ \bar{1}10 \}$ slip system | | $\langle 111 \rangle \{ 11\bar{2} \}$ slip system | |
|--|------------------|------------------|---|------------------------------------|---|------------------------------------|
| | | | γ_s (J m ⁻²) | γ_{us} (J m ⁻²) | γ_s (J m ⁻²) | γ_{us} (J m ⁻²) |
| CPMD, GTH [9] | 26.7 | 11.7 | — | — | — | — |
| QE, PAW [10] | 29.6 | 14 | — | — | — | — |
| VASP, PAW | 29.1 [84] | 14 [84] | 3.181 [43] | 1.633 [43] | 3.367 [43] | 1.830 [43] |
| VASP, PAW [88] | — | — | 3.197 | 1.6983 [87] | — | 1.7464 [87] |
| QE, NCPP [this work] | 27.763 [6%] | 14 [19%] | 3.3083 [4%] | ~1.680 | ~3.377 | ~1.764 |
| | | | | 1.6870 [3%] | 3.7620 [12%] | 1.8755 [7%] |

^aAcronyms used for software packages and pseudopotentials: Car–Parrinello molecular dynamic (CPMD) code [89], Goedecker–Teter–Hutter pseudopotentials [90].

all transmutants), W with primary transmutant (Re), and W with top three transmutants (Re, Os, Ta) as a function of the chemical compositions that result when pure tungsten is exposed to the EU-DEMO first wall conditions for ten years. While studying the effect of temperature on the stress–strain curve is outside the scope of this paper, it is still of interest to calculate the theoretical ideal tensile strength of transmuting tungsten in its ground state and relate our findings to the larger mechanical behavior of these candidate materials if possible.

First, we study the stress–strain relations of these materials as we increase the irradiation time. By way of example, the results for the chemical composition of transmuting tungsten that appears by the sixth year of irradiation are presented in figure 5(a). These reveal that the overall stress–strain curve drops as the number (and amount) of transmuting elements increases. Next, we analyze the impact on the ideal tensile strength σ_m , defined as the maximum in the stress–strain curve. The inset of figure 5(a) shows that while σ_m also decreases with the presence of transmuting elements, the related strain ϵ_m for which σ_m is achieved remains constant and ~14% for all the chemical compositions studied. Figure 5(b) summarizes the evolution of σ_m during the first ten years of irradiation in increments of one year. As the figure shows, σ_m monotonically decreases with the irradiation time for all

the scenarios considered. This is consistent with previous works that studied the ideal tensile strength of tungsten and tungsten alloys by first-principles calculations [9, 10]. Furthermore, adding all transmutants results only in a slight deviation from the chemical compositions that includes the top three transmutants (Re, Os, Ta). The observed higher values of σ_m for W including primary transmutant (Re) with respect to all-transmutants composition also aligns with the results obtained when studying the elastic behavior of transmuting tungsten, where the magnitude of the slope for W with primary transmutant was always smaller than the one for the current composition with all transmutants.

4.4. Dislocation-based ductility parameter of transmuted tungsten

In this section, we investigate the effect of transmuting elements on the ductility parameter of tungsten for $\langle 111 \rangle \{ \bar{1}10 \}$ and $\langle 111 \rangle \{ 11\bar{2} \}$ slip systems. We decided to focus here on pure W, the chemical composition that includes only the primary transmutant (Re), and the chemical composition that includes all transmutants. This choice is justified by (i) the increased computational cost to calculate γ_{us} (the reader is referred to appendix B for more details about the convergence tests shown there), and (ii) the small variations observed when studying the elastic properties and the ideal tensile strength

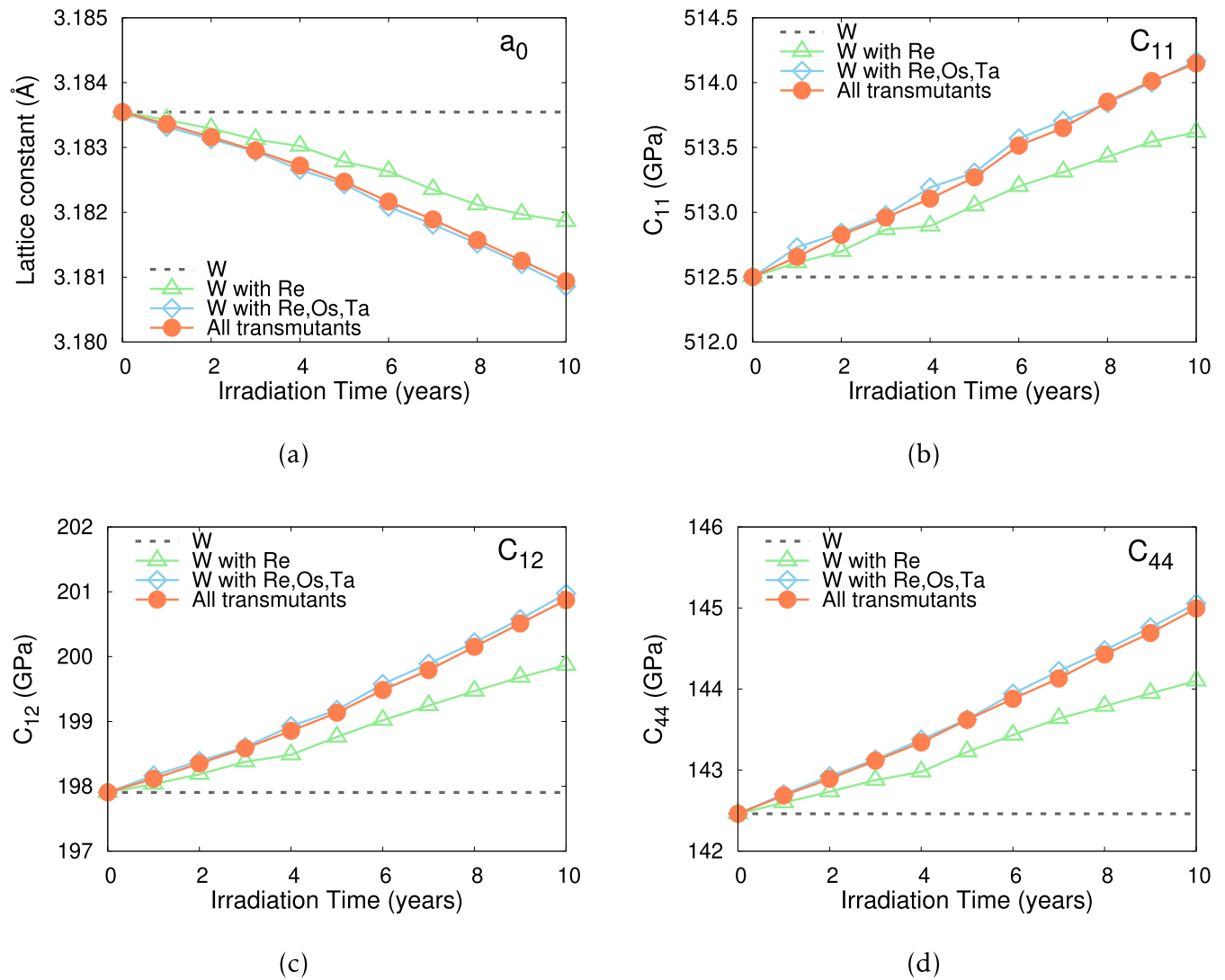


Figure 3. Evolution of the equilibrium lattice parameter a_0 and the elastic constants C_{ij} of transmuting tungsten during the first ten years of continuous exposure to EU-DEMO first wall conditions.

of the chemical composition including top three transmutants (Re, Os, Ta) and the chemical composition with all the transmutants.

As noted in section 3.3, the dislocation-based ductility parameter D formulated by Rice [36] can be obtained in terms of the surface energy γ_s and the unstable stacking fault energy γ_{us} . Next, we describe the results obtained when calculating each of these measurements, as well as the resultant ductility parameter.

γ_{us} is defined as the maximum value of the GSFE curve. As such, we need to calculate the GSFE curve for each of the chemical compositions and irradiation times considered. Figures 6(a) and (b) show the obtained GSFE curves of both $\langle 111 \rangle \{ \bar{1}10 \}$ and $\langle 111 \rangle \{ 11\bar{2} \}$ slip systems for the chemical compositions that results after irradiating pure tungsten at fifth year. The figures reveal several interesting trends that are consistent with previous works in the field. Firstly, γ_{us} is always in the middle of the energy path from one equilibrium position to another on a $\{ \bar{1}10 \}$ plane while there is certain asymmetry toward the first equilibrium position on a $\{ 11\bar{2} \}$

plane [43, 78, 87, 88]. Secondly, adding transmuting elements lowers γ_{us} of both slip systems [13, 39, 92]. Thirdly, γ_{us} is higher for the $\langle 111 \rangle \{ 11\bar{2} \}$ slip system [17, 38, 43, 87, 88]. The evolution of γ_{us} for both slip systems during the first ten years of irradiation is summarized in figures 6(c) and (d). These results show higher values of γ_{us} of the $\langle 111 \rangle \{ 11\bar{2} \}$ slip system over the entire time domain, and a monotonic decrease of γ_{us} as the irradiation time (and therefore the amount of transmuting elements) increases.

For its part, the calculation of γ_s in terms of the chemical composition and the irradiation time requires a preliminary step: the confirmation that γ_s does not diverge with increasing slab thickness [82, 93]. Following the method proposed by Fiorentini and Methfessel [82], figure 7(a) shows the calculated surface energies as a function of the slab thickness for the chemical compositions (all transmutants) that appear after 4 years of irradiation in tungsten. It can be seen from these results that convergence of γ_s is achieved with increasing the number of layers and a slab thickness of 12 layers assures a relative difference of less than 0.02%. With the confidence

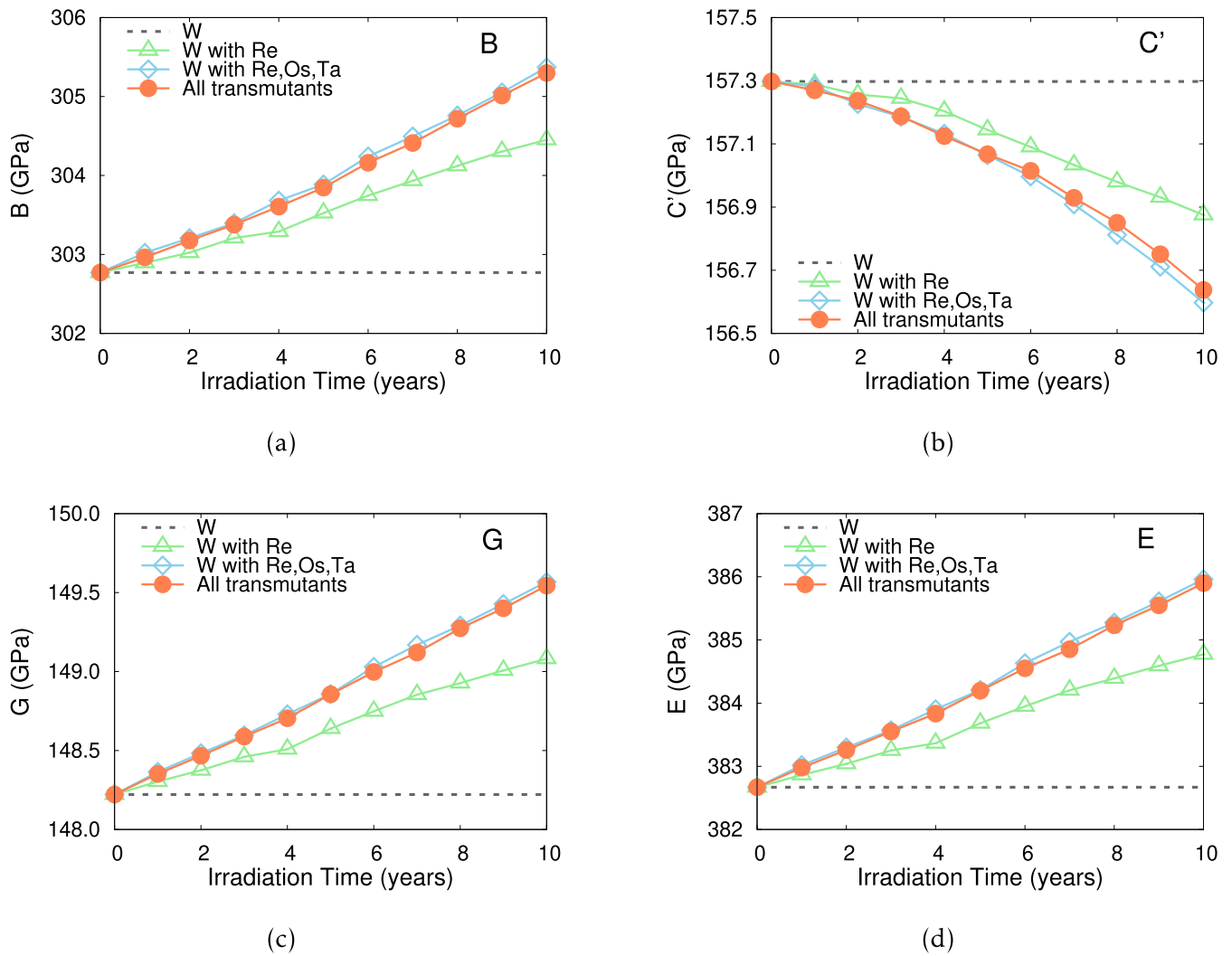


Figure 4. Evolution of the bulk modulus B , tetragonal shear modulus C' , shear modulus (G), and Young's modulus (E) of transmuting tungsten during the first ten years of continuous exposure to EU-DEMO first wall conditions.

conferred on our approach by the convergence tests, next we proceed to calculate the surface energy for a number of chemical compositions, irradiation times, and both slip systems. The results are summarized in figure 7(b). Despite the higher values of γ_s for the $\langle 111 \rangle \{11\bar{2}\}$ slip system, there is a small variation in terms of the chemical composition and the irradiation time studied, specially when compared with the evolution of γ_{us} shown in figures 6(c) and (d).

Once both γ_s and γ_{us} are calculated in terms of the chemical composition and the irradiation time, the ductility parameter D is obtained by substituting these terms into equation (12). We present results for the two slip systems of interest in figure 8, where the linear dependency with irradiation time is clearly distinguished. It can also be observed that D is higher on the $\langle 111 \rangle \{11\bar{2}\}$ slip system and the positive impact of adding more transmuting elements for a given irradiation time. This is in agreement with the well accepted improvement of the ductility of W when adding Re [17, 38, 43, 87, 88, 94].

5. Discussion

5.1. Nuclear transmutation

Section 2 presented typical transmutation results for pure W in a fusion reactor first wall (plasma-exposed) environment. While tungsten is a relatively highly transmuting element (for example, compare the relative burn-up of W to other elements in figure 6 of [95]), figure 1(b) and table 4 showed that W will remain relatively pure even after 10 years of exposure, with only 4–5 atomic % of transmutation impurities. However, even at such low concentrations, transmutants such as Re, Os, and Ta could still have significant impact on the engineering performance of W-based components in fusion systems. For example, 5% Re in tungsten can cause a 10%–20% reduction in the thermal diffusivity (and hence thermal conductivity) [96]. In a first wall fusion reactor having such a large variation of thermal performance in armour

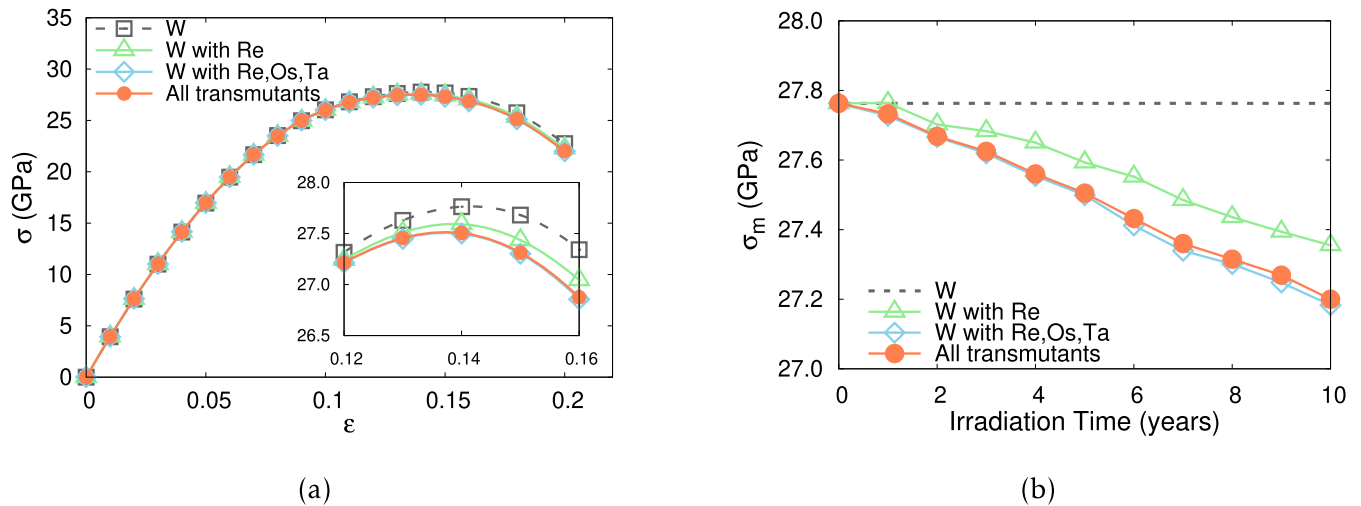


Figure 5. (a) The stress–strain curves under [001] tensile strain after six years of irradiation. (b) Evolution of the ideal tensile strength of transmuting tungsten during the first ten years of continuous exposure to EU-DEMO first wall conditions. These calculations are performed in their ground state and therefore they can not be compared directly to experimental stress–strain curves.

(where W is the preferred choice) would pose an engineering design challenge.

Even more problematic for the engineering of fusion components is uncertainties in predictions associated with nuclear response. In the present work, we have shown (table 1) a rare attempt to quantify uncertainties in transmutation predictions—in codes such as FISPACT-II [51] uncertainty quantification (UQ) is typically focused on radiological responses, which is clearly important as safety margins must be well understood to plan maintenance, handling, and dismantling operations where human (or even robotic) workers may be involved. However, where mechanical, structural, and thermal behavior may be impacted by the complete burn-up (transmutation) response (i.e. not merely from the typically small concentrations of radioactive products of nuclear reactions) it will be vital to provide UQ for the complete inventory evolution response.

The calculations performed in this work showed errors on the main transmutation products (Re and Os) of less than 10%, which is reasonable and typical of errors originating purely from nuclear data—see, for example, [97, 98], where decay-heat predictions by FISPACT-II for important fusion materials show good agreement with experimental measurements and where the computational errors generally encompass the experimental values. However, the errors shown in table 1, are not the complete picture—full UQ for the results would require, in addition to the errors originating from the nuclear data used in the inventory simulations, assessment of the errors associated with the neutron transport simulations used to generate the neutron spectrum.

In the present, the spectrum comes from a neutron transport simulation (performed using MCNP [99]) of a highly conceptualized model for EU-DEMO [55, 56]. As well as the errors inherent in the neutron transport simulations, which originates from the same nuclear data used for the inventory simulations with FISPACT-II, there is the unquantified—and

likely quite large—uncertainty in the fusion reactor model itself.

A key challenge for the future development of fusion, which will never have the wealth of experimental devices afforded the fission industry during its development, is to assess these engineering uncertainties, perhaps using Monte-Carlo-based sensitivity studies on the reactor design parameters or otherwise.

5.2. Effects of transmutant elements on the mechanical properties

In sections 3.1–3.3 we have provided calculations of the lattice parameter, elastic behavior, ideal tensile strength, and ductility parameter as new transmutant elements are added to the initial composition of pure tungsten due to irradiation. Furthermore, to better understand the effect of the most predominant elements produced in W under fusion-neutron irradiation, we also studied chemical compositions corresponding W with transmutant Re (W–Re) and also with Re, Os, and Ta (W–Re–Os–Ta) for each irradiation time. Here we consider the most important implications of the above calculations.

The results shown in figures 3 and 4 reveal that the lattice constant a_0 and the tetragonal shear modulus C' are the only two elastic measurements decreasing with increasing the amount of transmutant elements. These effects are closely related to the variation of the number of valence electrons per atom. For example, the reduction of the lattice constant (for a given irradiation time) with increasing the amount of transmutant elements is in agreement with experimental measurements and d -band filling predictions [91]. The decrease of C' with increasing impurity elements also suggests how transmuting tungsten becomes structurally unstable as the number of valence electrons increases. The variation of other elastic constants and properties such as C_{11} , C_{44} , B and E is also proportional to their solid solute concentrations and the number of valence electrons. These observations are in agreement with

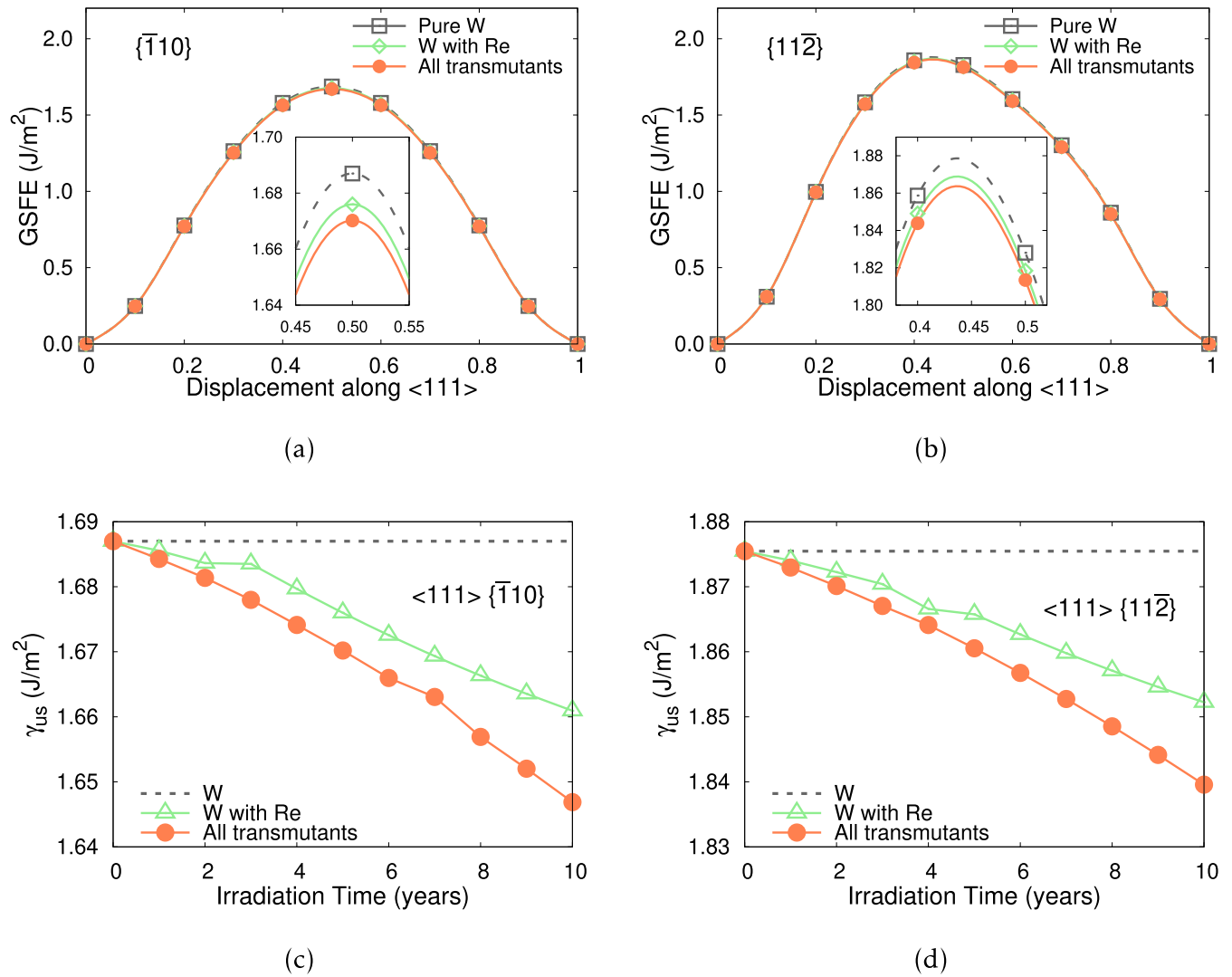


Figure 6. The general stacking faults energy for the slip along $\langle 111 \rangle$ direction in (a) $\{1\bar{1}0\}$ and (b) $\{11\bar{2}\}$ plane after 5 years of irradiation. Evolution of the unstable stacking fault energy γ_{us} for pure tungsten and transmuting tungsten during the first ten years of irradiation under EU-DEMO first wall conditions for both (c) $\langle 111 \rangle \{1\bar{1}0\}$ and (d) $\langle 111 \rangle \{11\bar{2}\}$ slip systems.

previous experimental [91] and computational works [10, 13, 17, 100]. When evaluating the quantitative effects of the chemical composition on the elastic behavior, the small amounts of impurity elements that appear due to transmutation (cf table 1) justify the smooth change of all these elastic parameters (the maximum difference observed is 1.8% for C_{44} after 10 years of irradiation). Still, it is noticeable the major qualitative role that Re plays in the evolution of the elastic behavior, compensating the effect of other transmutant elements like Ta that decrease the number of valence electrons [10, 100]. Finally, it can be observed from these figures that the effect of burn-up creating impurity elements with a relative concentration ≤ 0.87 at.% is negligible. This has computational consequences as adding extra elements significantly increases the computational cost of the simulations.

For its part, the variability of the ideal tensile strength shown in figure 5(a) can be explained by investigating the electronic structures of the chemical compositions compared. When group-VI transition metals like tungsten are deliberately

alloyed with elements with more valence electrons like Re or if those elements grow-in due to transmutation, the elastic shear instability is hindered by the Jahn–Teller distortion [101]. The result is a material that presents a slightly lower ideal tensile strength when compared to the pure element, as it is shown in the aforementioned figure. This is in fact consistent with previous experimental and computational works that describe how the peak stress of W–Re alloys are lower than the corresponding values of pure W [10, 100, 102–104]. Furthermore, similarly to the observations when studying the elastic behavior of transmuting tungsten, the small variations of the ideal tensile strength for a given irradiation time when adding Hf, H, He, Ir, and Pt can be explained by the small relative concentrations of these elements (always ≤ 0.87 at.%) that appear during the course of the 10 years power-plant first wall irradiation.

Another important physical aspect of mechanical deformation in bcc crystals is their ductility. As described in section

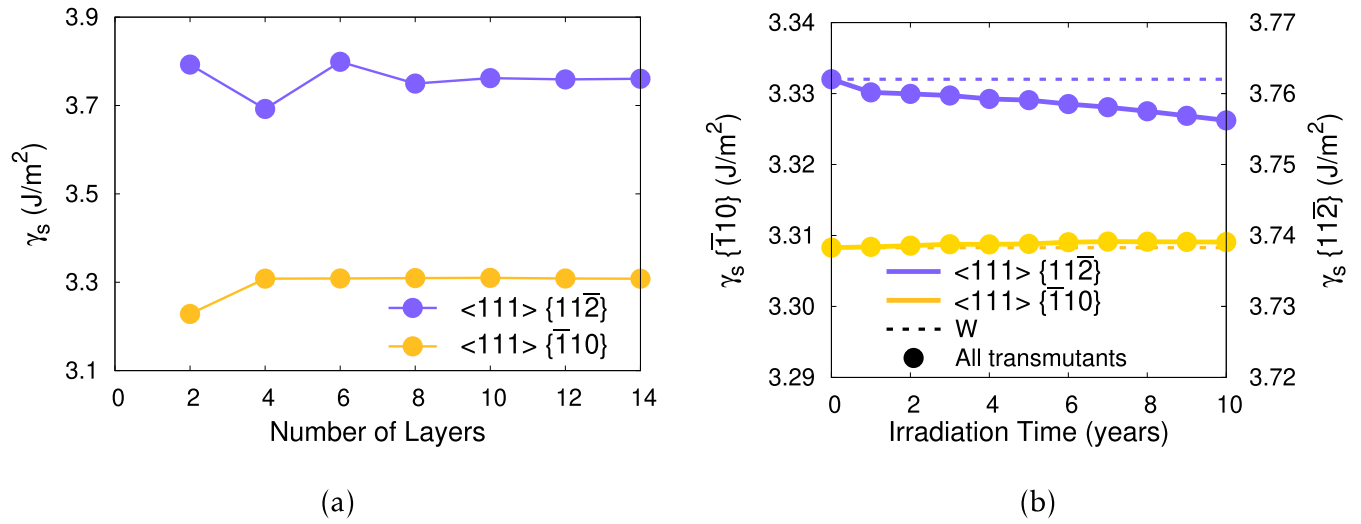


Figure 7. (a) Calculated surface energies for transmuting tungsten (all transmutants) as a function of the slab thickness after four years of irradiation; (b) evolution of the surface energy for pure tungsten and transmuting tungsten during the first ten years of irradiation under EU-DEMO first wall conditions for both $\langle 111 \rangle \{\bar{1}10\}$ and $\langle 111 \rangle \{11\bar{2}\}$ slip systems.

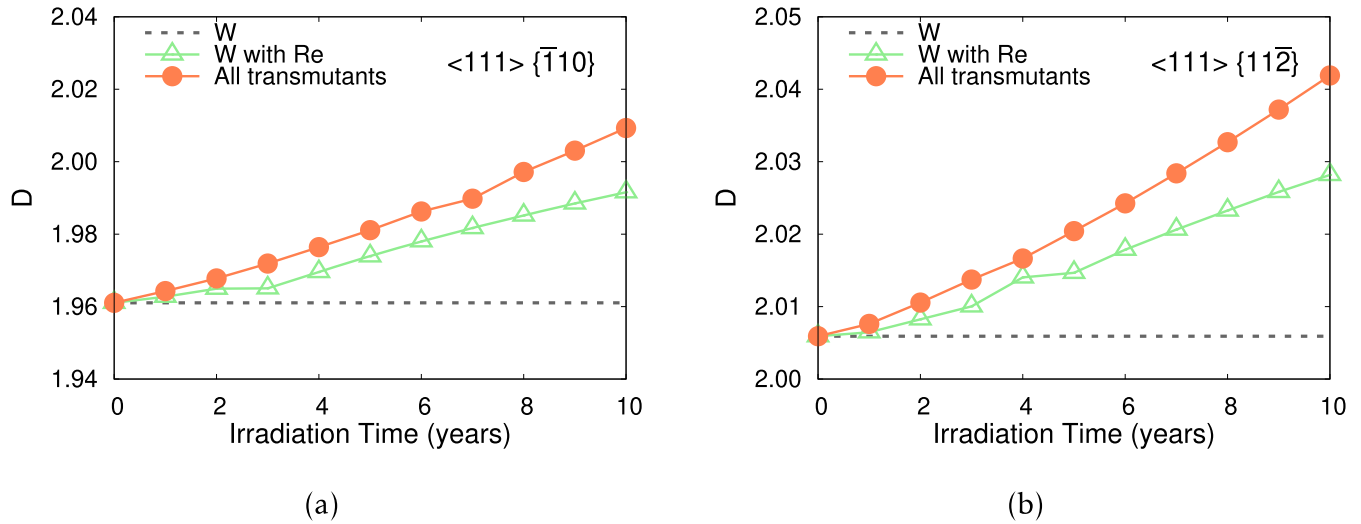


Figure 8. Evolution of the ductility parameter D for transmuting tungsten along the (a) $\langle 111 \rangle \{\bar{1}10\}$ and (b) $\langle 111 \rangle \{11\bar{2}\}$ slip systems during the first ten years of continuous exposure to EU-DEMO first wall conditions.

3.3, the Rice criterion [36] was chosen to estimate the ductility of transmuting tungsten. As such, the dependence of both γ_{us} and γ_s with respect to the chemical composition needs to be investigated prior to the calculation of the dislocation-based ductility parameter D (cf equation (12)). Several features can be discussed from these results, shown in figures 6 and 7. Firstly, the value of γ_{us} for each chemical composition on the $\langle 111 \rangle \{\bar{1}10\}$ slip system is always smaller than its counterpart on the $\langle 111 \rangle \{11\bar{2}\}$ slip system (cf figures 6(a) and (b)). This behavior suggests that slip should predominantly occur on $\{\bar{1}10\}$ planes, an observation with important physical consequences in the plastic deformation of tungsten-based materials [74, 77, 105, 106]. Secondly, the addition of transmuting elements decreases γ_{us} . Previous works have shown a different behavior depending on the number of valence electrons of the add-elements. Adding elements with more

valence electrons than W (such as Re, Os, Ir, Pt) have consistently reduced γ_{us} [43, 103] while adding elements with less valence electrons than W (such as Ta, Ti, Nb, Hf) tend to slightly increase it [43, 103]. However, when comparing the value of γ_{us} in pure W, W–Re and the chemical composition with all transmutants, we observe that the addition of elements always make γ_{us} to shrink. On this basis, we find that the higher relative concentration of Re with respect to other transmutants (cf table 4) plays a crucial role here, hindering the effect that other transmutant elements with less valence electrons than W should have.

Additionally, the results shown in figure 7(b) reveal that γ_s on the $\langle 111 \rangle \{\bar{1}10\}$ slip system is also smaller than its counterpart on the $\langle 111 \rangle \{11\bar{2}\}$ slip system. Still, the limited variability of γ_s with the chemical composition as irradiation time

increases implies that the main factor governing the dependence of D with the chemical composition is γ_{us} . Connecting the above observations regarding γ_{us} and γ_s , the results shown in figure 8 indicate that, for a given chemical composition, the ductility parameter is higher as the amount of transmutants is increased. This suggests that the addition of Re (which significantly decreases γ_{us} for a relatively constant γ_s) is the main factor responsible for this increment in the ductility.

5.3. Evolution of the mechanical properties with irradiation time

Previous works in the literature have extensively studied the effects of alloying elements on various properties of tungsten such as phase stability [10, 37–40], elastic properties [10, 13, 37–39, 41, 42], ideal tensile strength [9, 10], ductility [43], radiation defects [38], point defects [37, 44–48], dislocation structure [16, 49], grain boundaries [50], etc. However, to the best of our knowledge, there is a lack of understanding on how the expected first wall fusion power-plant conditions change the mechanical properties of these structural materials over time. Furthermore, most of these works have studied the effects when adding single or a reduce number of alloying elements to tungsten. In this work we address this gap by investigating not only the time dependence of the mechanical properties but also the effect of adding a combination of all the elements that appear due to transmutation. Our primary observation from the results shown in figures 3–5(b), 7(b), and 8 is that the time evolution of the mechanical properties is governed by the characteristics of the transmutant element with the highest relative concentration. As described in table 4, this corresponds to Re, an element with more valence electrons than W. The mechanical properties that monotonically increase when Re is added also increase as the exposure to first wall conditions continues, and vice versa. This effect (either increasing or decreasing the specific property) is slightly augmented when all the transmutants are considered, but the trend is governed by the primary transmutant element, no matter whether the other transmutants actually increase or decrease the number of valence electrons. However, we issue this conclusion with caution, as the relative concentrations of the deliberately added alloying elements in other candidate materials such as tungsten-based high-entropy alloys [107] are more similar and their mechanical behavior is expected to be different.

6. Conclusions

To summarize, we have used the FISPACT-II inventory code to calculate with uncertainties how the chemical composition of pure W changed when exposed to the EU-DEMO fusion first wall conditions for ten years. We have also performed first-principles calculations to characterize how the elastic constants, elastic properties, ideal tensile strength, unstable stacking fault energy, surface energy, and ductility parameter evolve as new transmutant elements are added to pure W due to transmutation. The novelty of our work lies in the integration of

nuclear transmutation and first-principles calculations to predict how the mechanical properties of plasma-facing irradiated materials change over time.

Our first conclusion from these investigations is that W will remain relatively pure even after 10 years of exposure to first wall fusion conditions, with only 4–5 at.% of transmutant elements. Secondly, we have observed that the effects of these transmutant elements on the mechanical properties of transmuting tungsten is closely related to the relative variation of the number of valence electrons per atom as the composition changes. In particular, we find that the relative difference of these valence electrons created by the most predominant transmutant element (Re), has an important influence all the properties calculated. Furthermore, the impact of minor transmutants seems negligible. Finally, our analysis of the mechanical properties as exposure to first wall fusion conditions continues over time also suggests that Re plays an important role in the evolution of the mechanical properties, increasing for example the Young's modulus, shear modulus, bulk modulus, and the ductility parameter; and decreasing the lattice constant, tetragonal shear elastic constant, ideal tensile strength, and the unstable stacking fault energy.

Our current and future efforts are directed toward studying the possibility of clustering of the transmutant elements [108]. As such, we are exploring other first-principles approaches such as the special quasirandom structures [109] or more complete alloy theories such as the coherent-potential approximation [110, 111]. Furthermore, we are also considering the selection of hybrid functionals.

Acknowledgments

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Data availability statement

The data that support the findings of this study are available upon reasonable request from the authors.

Appendix A. Concentration of transmuting elements

See table 4.

Appendix B. Convergence of the DFT calculations

Figures 9 and 10 show the convergence of the lattice parameter, elastic constants, and elastic properties with respect to the

Table 4. Relative concentrations that appear as W transmutes during the course of the 10 years power-plant first wall irradiation.

| Year | at% | | | | appm | | | | |
|------|-------|------|------|------|-------|------|------|----------------------|----------------------|
| | W | Re | Os | Ta | Hf | H | He | Ir | Pt |
| 1 | 99.61 | 0.31 | 0.01 | 0.07 | 2.3 | 5.6 | 1.2 | 4.4×10^{-6} | 3×10^{-8} |
| 2 | 99.14 | 0.64 | 0.04 | 0.18 | 8.2 | 11.3 | 2.4 | 1.6×10^{-4} | 3.6×10^{-6} |
| 3 | 98.67 | 0.95 | 0.09 | 0.28 | 17.3 | 17.0 | 3.6 | 1.2×10^{-3} | 5.0×10^{-5} |
| 4 | 98.22 | 1.23 | 0.17 | 0.38 | 29.4 | 22.6 | 4.7 | 4.9×10^{-3} | 3.1×10^{-4} |
| 5 | 97.77 | 1.48 | 0.26 | 0.48 | 44.0 | 28.4 | 5.9 | 1.4×10^{-2} | 1.2×10^{-3} |
| 6 | 97.33 | 1.72 | 0.37 | 0.57 | 60.9 | 34.1 | 7.1 | 3.5×10^{-2} | 3.7×10^{-3} |
| 7 | 96.90 | 1.94 | 0.49 | 0.65 | 80.0 | 39.8 | 8.3 | 7.2×10^{-2} | 9.3×10^{-3} |
| 8 | 96.49 | 2.14 | 0.63 | 0.73 | 101.1 | 45.6 | 9.5 | 1.3×10^{-1} | 2.1×10^{-2} |
| 9 | 96.07 | 2.33 | 0.77 | 0.80 | 124.1 | 51.4 | 10.8 | 2.3×10^{-1} | 4.1×10^{-2} |
| 10 | 95.67 | 2.50 | 0.93 | 0.87 | 148.8 | 57.1 | 12.0 | 3.7×10^{-1} | 7.5×10^{-2} |

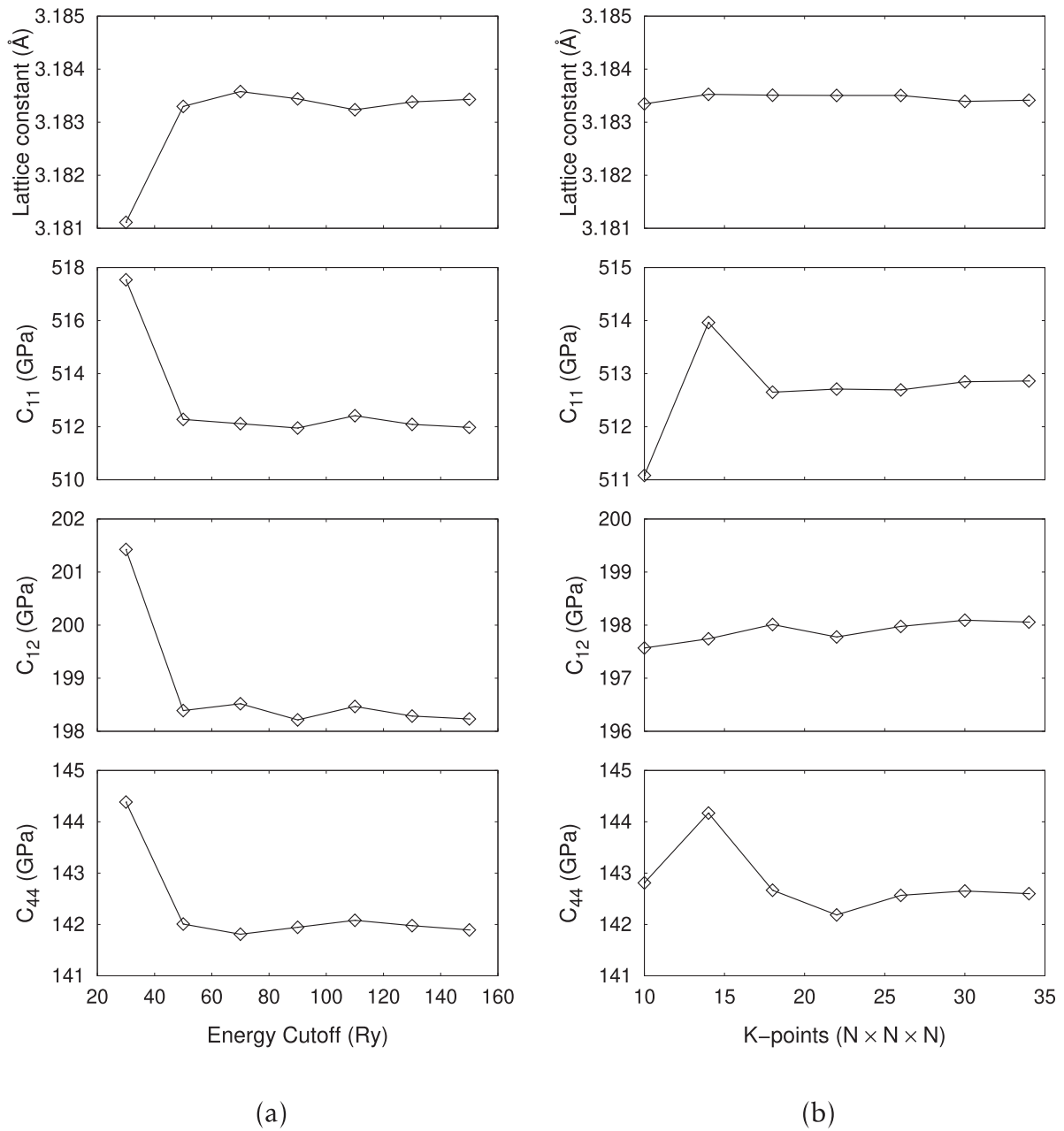


Figure 9. Convergence of the lattice constant and elastic constants C_{11} , C_{12} and C_{44} as a function of the plane-wave energy cutoff (a) and k mesh (b). The energy cutoff convergence (left) is studied keeping the k -points fixed to $20 \times 20 \times 20$. The k mesh convergence (right) is studied keeping an energy cutoff of 80 Ry.

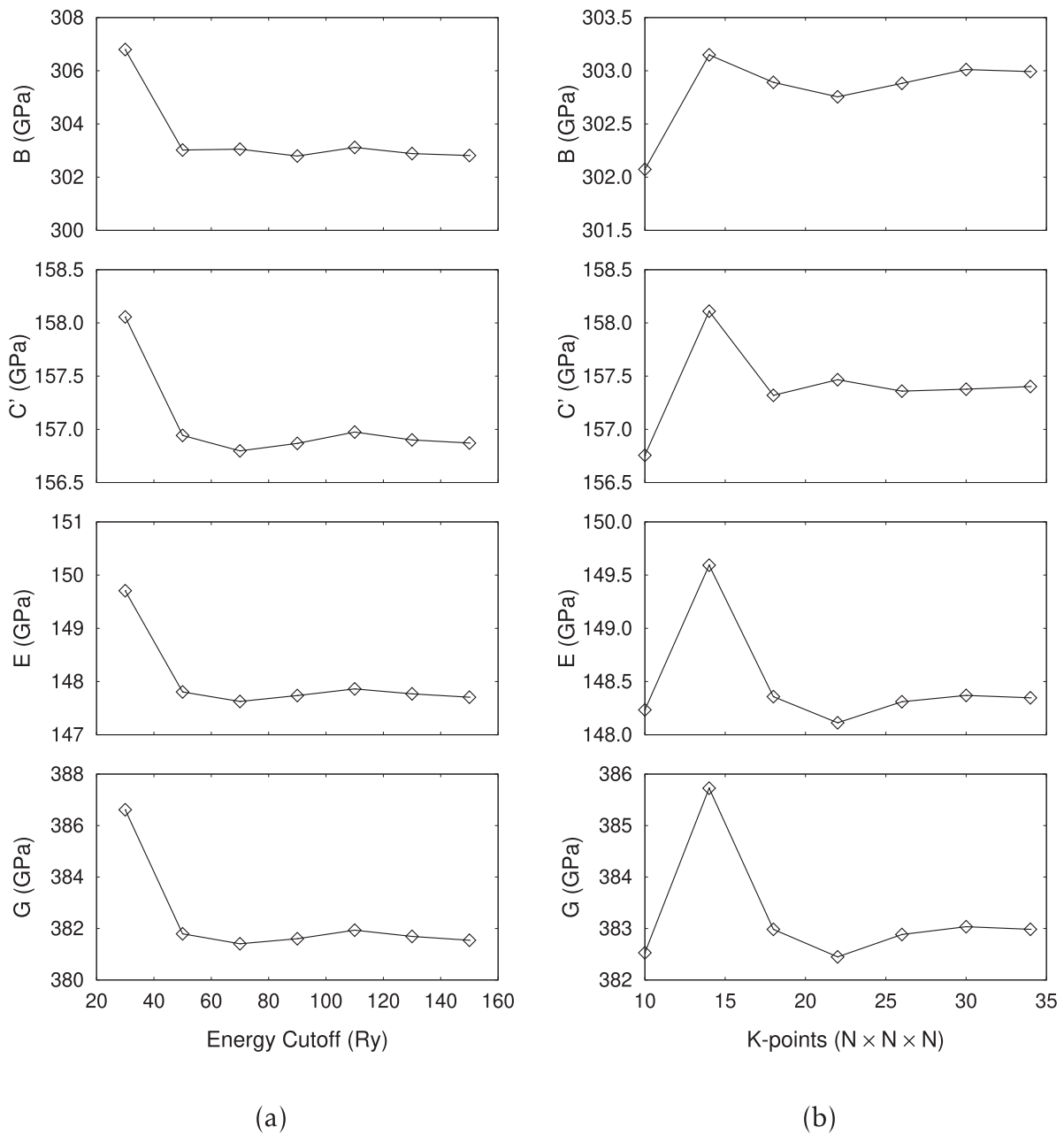


Figure 10. Convergence of the bulk modulus B , the tetragonal shear elastic constant C' , the shear modulus G , and the Young's modulus E as a function of the plane-wave energy cutoff (a) and k mesh (b). The energy cutoff convergence (left) is studied keeping the k -points fixed to $20 \times, 20 \times 20$. The k mesh convergence (right) is studied keeping an energy cutoff of 80 Ry.

plane-wave energy cutoff and the k mesh. These results suggest that an energy cutoff of 150 Ry and a $30 \times, 30 \times 30$ k -point mesh are sufficient to perform our first-principles calculations of the elastic behavior.

The convergence of the unstable stacking fault energy γ_{us} with respect to the energy cutoff, k mesh and slab size is shown in figure 11. First, we studied the convergence of γ_{us} with respect to the energy cutoff and k points, keeping a fixed slab structure of eight layers. In figure 11(a) the k -points were sampled using a fixed $16 \times 16 \times 1$ grid, being the energy cut-off the only parameter allowed to change. The results show that an energy cutoff of 40 Ry turns out to be sufficient, specially considering the computational cost and the number of

GSFE calculations needed. In figure 11(b) we did the opposite, fixing the energy cutoff to 60 Ry and allowing the k -points sample to change. Only small variations of less than 0.03% (for $\langle 111 \rangle \{ \bar{1}10 \}$) and 0.01% (for $\langle 111 \rangle \{ 11\bar{2} \}$) were found when comparing the results between a $28 \times, 28 \times 1$ and a $30 \times, 30 \times 1$ grid. As such, we chose the $28 \times, 28 \times 1$ k -point mesh as a good compromise between computational accuracy and computational cost for both slip systems. Then we fixed the energy cutoff to 40 Ry and the k points sampling to a $28 \times 28 \times 1$ grid, allowing the slab size to change. Previous works in the literature (cf table 3) have shown that γ_{us} for the $\langle 111 \rangle \{ 11\bar{2} \}$ slip system is higher than its counterpart for the $\langle 111 \rangle \{ \bar{1}10 \}$ slip system. The results shown in figure 11(c)

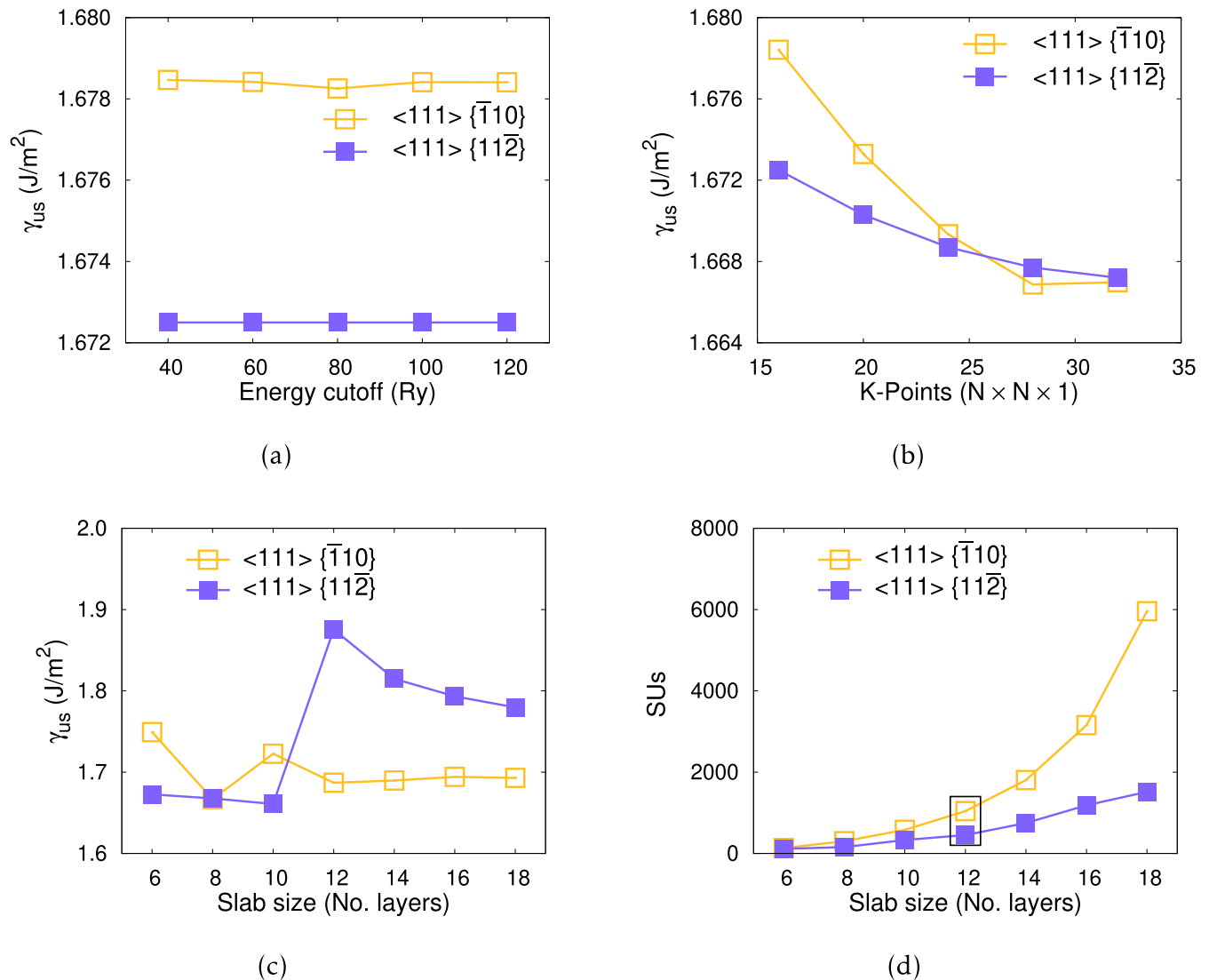


Figure 11. Benchmarking to identify the set of parameters that guarantee accurate and efficient DFT calculations of the unstable stacking fault energy γ_{us} for transmutated tungsten. Figures (a)–(c) show the convergence with the energy cutoff, k -points sampling, and slab size, respectively. Figure (d) shows the total number of SUs needed in Expanse XSEDE cluster to complete each simulation. The highlighted region in figure (d) indicates the total number of SUs for the final set of parameters.

reveal that a minimum of 12 layers is needed to avoid image effects that would reverse this trend. Given the periodicity of the cell (six layers) along the $\{11\bar{2}\}$ plane and the significant computational increase of choosing 18 layers instead of 12 layers shown in figure 11(d), we decided to choose a slab size of 12 layers in our simulations.

These convergence tests and the selection of accurate but reasonable DFT parameters are relevant to our approach given the significant computational cost of the first-principles simulations performed to obtain the GSFE along the entire $\langle 111 \rangle$ path for each of chemical compositions at each the irradiation time.

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