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Regression and clustering algorithms for AgCu nanoalloys: from mixing energy predictions to structure recognition†

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The lowest-energy structures of AgCu nanoalloys are searched for by global optimization algorithms for sizes 100 and 200 atoms depending on composition. Even though the AgCu system is very weakly miscible in macroscopic samples, the mixing energy for these nanoalloys turns out to be clearly negative for both sizes, a result which is attributed to the stabilization of non-crystalline Cu@Ag coreshell structures at the nanoscale. The mixing energy is a quantity nowadays unknown in its functional form, so that its prediction may take advantage of machine learning techniques. A support vector regressor is then implemented to successfully predict the mixing energy of AgCu nanoalloys of both sizes. Moreover, with the help of unsupervised learning algorithms, it is shown that the automatic classification of such nanoalloys into different physically meaningful structural families is indeed possible. Finally, thanks to the harmonic superposition approximation, the temperature-dependent probabilities of such structural families are calculated.

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1 Introduction

Since the understanding of their wide range of applications, ranging from catalysis^{1,2} to data storage,³ plasmonics,⁴ biomedicine,⁵ water purification⁶ and others, nanoparticles (NPs) have been studied rather extensively in recent years. For this reason, both experimental and theoretical efforts have been made in order to understand the physical and chemical properties of such systems.⁷ In particular, the obstacles inherent to the theoretical study of a system of a finite number of atoms are often tackled with the help of computer simulations (e.g. global optimization techniques8 and molecular dynamics simulations⁹), which are nowadays an essential partner of experiments.

Only in the past few years, also Machine Learning has been proposed as an effective tool to investigate and solve some problems related to the modeling of nanoparticles, 10 such as energy landscape exploration^{11,12} and dynamic predictions via force field reconstruction,13 but also binding energy14 and free energy¹⁵ predictions, and atom classification.¹⁶ In a very broad general depiction, machine learning algorithms can be divided in two large classes: supervised and unsupervised learning algorithms. The first class deals with data sets consisting of both input and output data, and tries to establish a relation between these sets for fitting or classification purposes. The second class deals with unlabelled data, i.e. only input data, and tries to work out a specific task such as clustering the data into separate sets, feature selection or dimensionality reduction.

Here in particular we decided to implement both supervised and unsupervised learning algorithms, showing indeed their ability to capture different interesting properties of our data set. It is in fact known that a regression algorithm is capable of detecting subtle relationships between input and output data, whereas a clustering algorithm can identify subgroups of an ensemble without any prior knowledge. Our data sets were collected by global optimization searches of the lowest-energy structures of AgCu nanoalloys of sizes N = 100 and N = 200atoms, for different compositions. Machine learning algorithms were then trained in order to make predictions of the mixing energy, a quantity not known in its functional form, and also to find structural families without any previous classification. Finally, we calculated probabilities for such different structural families as a function of temperature by means of the Harmonic Superposition Approximation (HSA). 17-19

AgCu nanoalloys are important for a different number of remarkable reasons. In general, the bi-metallic nature of nanoalloys allows to enrich their range of applicability in real life situations, since the chemical ordering of equilibrium and

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out of equilibrium structures will induce different desirable properties, missing most of the times in their single metal counterparts. For example, AgCu nanoalloys have shown interesting plasmonic, 20 electrical, 21 antibacterial 22 and catalytic 23 properties, but also applications in corrosion resistance²⁴ and solar cells.²⁵ The interest in the theoretical modeling of such nanoalloys is therefore motivated by this impressive variety of experimental results.

AgCu is a weakly miscible system in bulk samples, 26 with positive (endothermic) mixing energy. However, global optimization studies of small nanoalloys (of 34 and 38 atoms)²⁷ showed that the mixing energy at the nanoscale is negative in the whole composition range. It is therefore interesting to check whether this behaviour persists at larger sizes such as those considered in this article (100 and 200 atoms).

The material is organized as follows. The second section includes a brief explanation of the theoretical methods, including Machine Learning (ML) and global optimizations algorithms, as well as more specific topics such as Common Neighbor Analysis (CNA) and Harmonic Superposition Approximation (HSA). The third section is entirely dedicated to the results of the applications of such methods to AgCu nanoalloys of 100 and 200 atoms. Finally in the fourth and last section, the conclusions can be found.

A note on terminology. In order to avoid confusion, aggregates of atoms will be referred to as nanoparticles or nanoalloys. On the other hand, the term cluster will be used to denote a set of nanoparticle structures which are grouped together by a clustering algorithm working in the space of suitable order parameters (i.e. of structural descriptors).

2 Models and methods

2.1 Atomistic force field

Interactions between NPs were modelled by an atomistic potential, in the form proposed by Gupta²⁸ and by Rosato et al.,29 which can be derived from the second moment approximation to the tight-binding model.30 The potential energy E is the sum of one-atom contributions E_i containing a bonding (E_i^b) and a repulsive part (E_i^r) :

$$E = \sum_{j} E_j = \sum_{j} \left(E_j^b + E_j^r \right), \tag{1}$$

where

$$E_{j}^{b} = -\sqrt{\sum_{i \neq j} \xi_{sw}^{2} \exp\left[-2q_{sw} \left(\frac{r_{ij}}{r_{sw}^{0}} - 1\right)\right]},$$
 (2)

and

$$E_j^r = \sum_{i \neq j} A_{sw} \exp\left[-p_{sw} \left(\frac{r_{ij}}{r_{sw}^0} - 1\right)\right]. \tag{3}$$

 r_{ij} is the distance between atoms i and j. s(w) refers to the chemical species of the atom i(j). If s = w, r_{sw}^0 is the nearestneighbor distance in the corresponding bulk lattice, while for $s \neq w$, r_{sw}^0 is taken as the arithmetic mean of the distances of

Table 1 Parameters of the interaction potential. From ref. 31

	p	q	A (eV)	ξ (eV)	r_0 (Å)	r_{c1} (Å)	r_{c2} (Å)
Cu-Cu	10.55	2.43	0.0894	1.2799	2.56	4.08707719 3.62038672 4.08707719	4.43405007

pure metals. Cutoff distances on the interactions are imposed as follows. The exponentials in eqn (2) and (3) are replaced by fifth-order polynomials, of the form $a_3(r-r_{c2})^3 + a_4(r-r_{c2})^4 +$ $a_5(r-r_{c2})^5$, between distances r_{c1} and r_{c2} , with a_3 , a_4 , a_5 fitted in each case to obtain a function which is always continuous, with first and second derivative for all distances, and goes to zero at r_{c2} . The parameters of the potential were taken from ref. 31 and are reported in Table 1.

This interaction potential has been used previously to model the structures of AgCu nanoalloys and compared to Density Functional Theory (DFT) calculations and experimental results,32-35 obtaining quite good agreement with DFT data on the behaviour of the composition-dependent excess energy and on the structures of lowest-energy magic polyicosahedra of sizes 34, 38 and 45,32 on the energetics of the placement of Cu impurities in Ag icosahedra and truncated octahedra (see ref. 7, Tables 5.1 and 5.3), on the relative stability of Mackay, anti-Mackay and chiral Cu@Ag icosahedral structures (see ref. 33). Moreover, the agreement between the predictions of this model and the experimentally observed structures is quite good³⁴ as discussed in ref. 35. Therefore we believe that this model, although approximate, is able to catch the relevant structural aspects of AgCu nanoalloys, with the advantage of allowing such a thorough exploration of the nanoalloy energy landscape that would be unfeasible by DFT calculations.

Global optimization and data collection

Each data set consisted of nanoparticle structures obtained by global optimization searches. Therefore, all structures considered in the following correspond to local minima in the energy landscape, and, where specified, to global minima (that are the lowest-energy local minima). These searches were made by the Basin Hopping (BH) algorithm, 36 using our own code. 37-39 For each composition, three independent simulations of 200 000 BH steps each were made, using different seeds and parameters. Generally speaking, each of these three simulations had different proportions for Brownian, exchange, shake and bonds moves. 37,38 Two simulations over three were using a single Monte Carlo walker (standard BH algorithm), whereas the third simulation used three different walkers running in parallel (parallel excited walker algorithm³⁷). The use of different global optimization algorithms usually allows a more thorough exploration of the energy landscape.³⁷

The mixing energy E_{mix}^{7} was used to analyze the energetic stability of the nanoalloys depending on composition. E_{mix} is defined as follows

$$E_{\text{mix}}(m,n) = E(m,n) - \frac{m}{N}E(N,0) - \frac{n}{N}E(0,N)$$
 (4)

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(b) 555 (d) 422 (e) 421

Fig. 1 Some CNA signatures. 555 signature is typical of atom pairs along 5-fold symmetry axes, such as those of icosahedra and decahedra. 421 signature is that of a perfect FCC crystal fragment, whereas 422 signature is typical of FCC fragments with local HCP zones (stacking faults, twin planes).

where E(m, n) is the binding energy of a nanoalloy with m silver atoms, and n = N - m copper atoms, N being the total number of atoms. The mixing energy vanishes for pure nanoparticles, *i.e.* when m = 0, n = N and *vice versa*. When $E_{\text{mix}}(m, n) < 0$, then

$$NE(m, n) < mE(N, 0) + nE(0, N)$$
 (5)

that is, a mixture of m pure NPs of the first element plus n of the second element is higher in energy than N nanoalloys of such composition.

Each nanoparticle structure was described with the aid of two other parameters, coming from the Common Neighbor Analysis⁴⁰ (CNA). For each pair of nearest-neighbor atoms, the CNA defines a signature consisting of a triplet of integer numbers *rst*:

- \bullet r the number of common nearest neighbors of the pair.
- s the number of bonds between those r atoms.
- t the length of the longest chain of bonds that can be made out of the s bonds present.

For a given nanoalloy and signature, we define its signature order parameter as the number of nearest-neighbor pairs presenting such signature divided by the total number of nearest-neighbor pairs in the nanoalloy. Some typical signatures are shown in Fig. 1. In particular, here we decided to use first a two-dimensional description based on the (555, 422) pair of signatures, and the to compare the results to those obtained by means of different choices of the variables.

2.3 Machine learning models

In order to find a suitable approximation of the mixing energy, regression models were trained and tested. For a classification of nanoalloys into structural families, unsupervised learning algorithms were also implemented. Both tasks were achieved thanks to the open source software Scikit-Learn. 41

2.3.1 Regression. Support Vector Machines^{42–44} (SVMs) are very powerful tools, and given their flexibility they can adapt to a very large variety of data sets. In addition, they benefit from the property of continuity, which is not shared among other powerful models such as, for example, decision trees and random forests. The general formulation of the regression problem for SVMs (in this case the model is referred to as Support Vector Regressor, SVR), can be stated as follow.⁴⁵ Given

a training set of n pairs of data $(x_1, y_1), ..., (x_n, y_n)$, the algorithm solves⁴⁶ the following problem:

$$\underset{\alpha_{i}, \alpha_{i}^{*}}{\operatorname{argmax}} \left[-\frac{1}{2} \sum_{i,j=1}^{n} (\alpha_{i}^{*} - \alpha_{i}) (\alpha_{j}^{*} - \alpha_{j}) k(x_{i}, x_{j}) \right.$$

$$\left. -\varepsilon \sum_{i=1}^{n} (\alpha_{i}^{*} + \alpha_{i}) + \sum_{i=1}^{n} (\alpha_{i} - \alpha_{i}^{*}) y_{i} \right]$$

$$(6)$$

subject to

$$\begin{cases} \sum_{i=1}^{n} (\alpha_i^* - \alpha_i) = 0\\ 0 \le \alpha_i, \quad \alpha_i^* \le C \end{cases}$$

where ε and C are two positive hyperparameters and $k(x_i, x_j)$ is a kernel function. The parameters that have to be found by the algorithm are α_i and α_i^* for each i. Here we used the radial basis function (RBF) kernel, that is $k(x_i, x_j) = \mathrm{e}^{-\gamma ||x_i - x_j||^2}$. The constant γ is the third positive hyperparameter. Given a set of hyperparameters and a solution to the minimization problem $\{\alpha_i, \alpha_i^*\}_{i=1}^n$, the functional relation between x and y can be expressed as

$$f(x) = \sum_{i=1}^{n} (\alpha_i - \alpha_i^*) k(x_i, x) + b$$
 (7)

where b is a constant calculated during the minimization process. To avoid overfitting,⁴⁷ hyperparameters are tuned after V-fold cross validation. Finally the model capability is evaluated on the test set $(x_1, y_1), \ldots, (x_m, y_m)$, using the R^2 metric

$$R^{2} = 1 - \frac{\sum_{i=1}^{m} (y_{i} - f(x_{i}))^{2}}{\sum_{i=1}^{m} (y_{i} - \bar{y})^{2}}$$
(8)

where \bar{y} is the average of all y_i in the test set. The score takes values in the range $(-\infty, 1]$, where 1 means a perfect model.

2.3.2 Clustering. Unsupervised learning algorithms group a set of unlabelled points in space. Here we implemented K-means^{42,48} and Gaussian mixture model^{42,49,50} (GMM), in a two dimensional space description given by the (555, 422) CNA signatures. These signatures have been chosen because they single out local fivefold symmetry points and stacking faults in the fcc lattice, respectively. The *K*-means algorithm solves iteratively the minimization problem of finding the Voronoi tessellation of the data set, according to

$$\min_{m_1,\dots,m_K} \sum_{i=1}^n \min_{j=1,\dots,K} ||x_i - m_j||^2$$
 (9)

where K is the number of clusters, n is the number of points x_i , and m_j are the centers of each cluster. The number of clusters is not known a priori, so that a score based on that number must be provided to evaluate the quality of the result. Here we used the silhouette score, 51 which is the mean of all the silhouette coefficients assigned to each point in the set:

$$s_i = \frac{b_i - a_i}{\max(a_i, b_i)} \tag{10}$$

where a_i is the mean distance to the other instances in the same cluster of x_i , and b_i is the smallest mean distance to the other instances in each cluster that does not contain x_i . The optimal value of K is the one that maximizes the silhouette score, whose range is [-1, 1]. GMM is instead a probabilistic model: it assumes all points to be generated by K Gaussian distributions with unknown weights and parameters. The solution is provided by the Expectation-Maximization (EM) algorithm, which iteratively maximizes the expectation over the current parameters of the log-likelihood of the mixture model. By assumption, all clusters can have ellipsoidal shape, which is a generalization of K-means "hard" clustering. Again, since the number of mixture components (i.e. the number of clusters) is not known a priori, there must be an objective method to deduce it. Here we used the Bayesian Information Criterion 42,51 (BIC), which is defined as

$$BIC = \log(n)p - 2\log\hat{L} \tag{11}$$

where n is the number of points, p the number of parameters and \hat{L} is the maximized value of the likelihood function. The optimal value of K is the one that minimizes the BIC.

2.4 Harmonic superposition approximation (HSA)

Probabilities for nanoparticles as a function of temperature were calculated thanks to the Harmonic Superposition Approximation. The Given a pool of n_{\min} structures that locally minimize the potential energy surface for a fixed number of atoms N, this framework allows to calculate the probability of each structure. For a local minimum s with potential energy E_s , it is assumed that the partition function can be decomposed into translational, rotational and vibrational terms:

$$Z_s = \frac{1}{h_s} Z_s^{\text{tr}} Z_s^{\text{rot}} Z_s^{\text{vib}} e^{-\beta E_s}$$
 (12)

where h_s is the order of symmetry group of the local minimum s and $\beta = 1/k_BT$, where T is the temperature of the canonical ensemble considered. In particular we have that:

$$Z_s^{\rm tr} = V \left(\frac{Mk_{\rm B}T}{2\pi\hbar^2}\right)^{3/2} \tag{13}$$

is the translational term, where V is the volume of the box containing the nanoparticle, M the total mass;

$$Z_s^{\text{rot}} = \left(\frac{2\pi k_{\text{B}} T \bar{I}_s}{\hbar^2}\right)^{3/2} \tag{14}$$

is the rotational term, where \bar{I}_s is the average moment of inertia of local minimum s: $\bar{I}_s = (I_s^{xx} I_s^{yy} I_s^{zz})^{1/3}$ and I_s^{xx} , I_s^{yy} and I_s^{zz} are the principal moments of inertia;

$$Z_s^{\text{vib}} = \prod_{i=1}^{3N-6} \frac{1}{2\sinh(\beta\hbar\omega_{s,i}/2)} \tag{15}$$

is the vibrational term in the harmonic approximation, where $\omega_{s,i}$ is the *i*-th non zero normal mode of the local minimum s.

The probability as a function of temperature of the local minimum s in the ensemble is then given by

$$p_{s} = \frac{Z_{s}}{\sum_{\sigma=1}^{n_{\min}} Z_{\sigma}}$$
 (16)

Eqn (16) gives the relative probability for a minimum s in a pool of n_{\min} isomers as a function of temperature. In fact clustering algorithms (see Section 2.3.2) separate structures into different structural motifs, so that their temperature dependent probability can be calculated by the HSA, as done in Section 3.4.

3 Results

In Sections 3.1 and 3.2 we report the results of both regression and clustering algorithms applied to the global minima found by our global optimization searches for sizes N=100, and N=200. For N=100, the data set consists of the global minima of Ag_mCu_n for all compositions, *i.e.* for m=0,1,2,...,100. For size 200, where calculations are more cumbersome, the data set consists of the global minima of Ag_mCu_n with even m, *i.e.* m=0,2,4,...,200. The global minima were searched for by the BH algorithm, which has proven its efficiency in the optimization of AgCu nanoalloy structures. ^{33,52-54} For both N=100 and 200 we provide the results for the regression of the mixing energy, as given in eqn (4), as well as for the clustering of the data sets in different structural families. Such structural families are then described in terms of their main geometrical features and chemical ordering.

In Section 3.3, we discuss different possible choices of the order parameters for the application of the clustering algorithms. Finally, in Section 3.4, we consider a specific composition, $Ag_{64}Cu_{36}$. For that composition we consider a set of low-energy local minima collected by our BH searches, we use the clustering algorithm to separate these structures into families, and we calculate the temperature-dependent probability of the families by means of the HSA.

3.1 N = 100

The mixing energy, as a function of the number of silver atoms m is shown in Fig. 2. The mixing energy is negative in the whole composition range. Its profile is evidently rather complicated, with different stationary points.

For a selected set of compositions, we checked the behavior of the mixing energy by DFT calculations. The DFT results (reported in the ESI†) are in good agreement with those of the Gupta potential, confirming the overall behavior of the mixing energy, and even giving somewhat more negative values. The negative values of the mixing energy in nanoalloys such as AgCu, AgNi, AgCo, AuCo and others was attributed to the efficient stress relaxation achieved by core@shell structures in which the element with lower surface energy and larger atomic size is segregating at the surface.³⁵ The DFT data confirm this result and show that, at least for AgCu, the electronic effects not included in the Gupta model, such as directional terms in

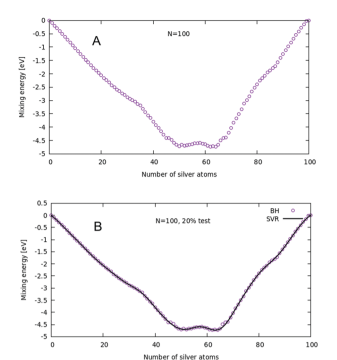


Fig. 2 (A) Mixing energies calculated after global optimization. (B) SVR (20% test) predictions for every composition, along with mixing energies calculated from global optimization.

bonding between atoms or charge transfer, are likely to be of minor importance.

For this size, the mixing energy is calculated for every possible composition, so that in principle there is a complete knowledge about this function. However, for larger nanoalloys (as we shall see later) it can be difficult and rather expensive to compute the mixing energy for every possible composition, since it requires a full global optimization for each composition, so that in principle it is useful to have a method capable of making predictions where they are needed, i.e. on "unseen" data.

To make such predictions, we trained and tested a SVR splitting randomly the data set into an 80% training set and a 20% test set, using the number of silver atoms m as the only variable. In order to avoid overfitting, hyperparameters were tuned from the following ensemble, which spans different orders of magnitudes, after 5-fold cross validation:

$$\begin{cases} \gamma : [0.001, 0.005, 0.01, 0.05, 0.1, 0.5, 1, 3, 5, 8, 10] \\ C : [0.001, 0.005, 0.01, 0.05, 0.1, 0.5, 1, 5, 10, 50, 100, 500, 1000] \\ \varepsilon : [0.001, 0.005, 0.01, 0.05, 0.1, 0.5, 1, 5, 10] \end{cases}$$

The best triplet was found to be composed by $\gamma = 0.01$, C = 50and ε = 0.01. Given this result, the model was then trained on the full training set (we remember that during cross validation a fifth of it was set a part for each validation), and tested on the test set, giving the following scores:

$$R_{\text{training}}^2 = 0.999825 \qquad R_{\text{test}}^2 = 0.999657$$

that are indeed very high. Other details regarding the SVR model (e.g. parameters in eqn (7)) can be found in the ESI.†

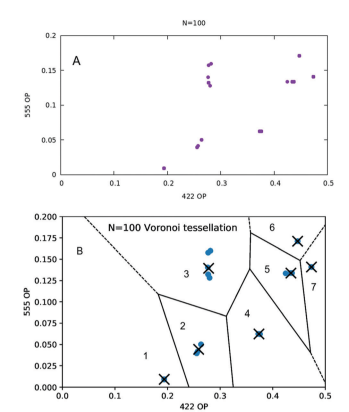


Fig. 3 (A) Data set representation in the two dimensional space of 422 and 555 order parameters. For each nanoalloy, we calculated its 422 and 555 order parameters as the fractions of nearest-neighbor pairs presenting such signatures. (B) Voronoi tessellation and cluster centers (means), for K = 7, when N = 100.

The performance of the model on the entire data set is shown in Fig. 2B. The same data set was then split in 50% training set and 50% test set. For this setting we found $\gamma = 0.005$, C = 1000and ε = 0.005. The two scores are again both very high:

$$R_{\text{training}}^2 = 0.999875 \quad R_{\text{test}}^2 = 0.999468$$

As it can be seen in the ESI,† the two results are very similar in terms of predictions, even though hyperparameters are quite different.

When the data set is represented in the two dimensional space given by the 422 and 555 CNA signatures, all nanoalloys automatically separate into different groups. However, it is still not clear at this stage how many are there, as shown in Fig. 3A. After K-means is implemented, and the silhouette score is plotted as a function of the number of clusters K, one obtains the plot given in Fig. 4A. The optimal number of clusters, according to this criterion, is then K = 7. The Voronoi tessellation, along with each cluster center, is shown in Fig. 3B. A representative structure for each family can be found in Fig. 4B. Further details and images are given in the ESI.† Here we give a description of the different structural families found, specifying the number of silver atoms m.

• (1)
$$m = 0-4$$
, $m = 100$

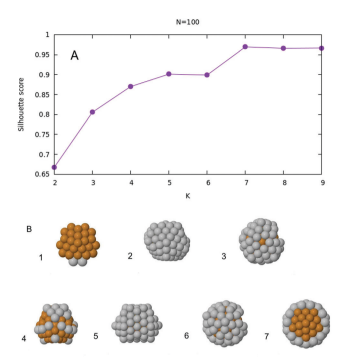


Fig. 4 (A) Silhouette score as a function of K for the case N = 100(B) Seven representative structures, one for each family. Labels are referred to Fig. 3B. The seven structures have 2, 82, 62, 23, 72, 56 and 45 silver atoms respectively.

These are the only six structures with Marks decahedral⁵⁵ symmetry.

• (2) m = 79-83

Five asymmetric icosahedral structures.

• (3) m = 59-66

These eight nanoalloys have a 55-atom perfect Mackay icosahedron covered by an incomplete Mackay crust, which is slightly distorted and presents a small rotation.

 \bullet (4) m = 5-34, m = 84-99

These forty six structures, which compose the largest cluster, have icosahedral symmetry. They are basically composed of a 55-atom perfect Mackay icosahedron covered with an incomplete Mackay icosahedral shell (which is part of the surface of the 147-atom icosahedron). However, even if they share this common feature, there are two subgroups that can be identified. The first subgroup is that of m = 5-34, where the silver atoms are at the nanoalloy surface, typically occupying icosahedral vertices and the external shell. The second subgroup can be identified with the remaining nanoalloys. Those structures have all of their copper atoms in the core, typically forming part of the 13-atom perfect icosahedron.

• (5) m = 67-78

These twelve structures are polyicosahedra³² resulting from three joined 55-atom icosahedra sharing some atoms. The third icosahedron is incomplete because of an insufficient total number of atoms.

 \bullet (6) m = 54-58

These five structures are again core-shell as those in group 4. However their crust is of anti-Mackay type.³³ Moreover, they have part of such crust removed at the boundary and some silver atoms are placed at the fivefold vertices. We remark that this small deviation (their 422 signature differs 0.0262 from the seventh group) was detected by the clustering algorithm.

•
$$(7) m = 35-53$$

All nineteen these structures are similar for their icosahedral symmetry to those already described in group 4 and 6, however they have the 55 perfect Mackay icosahedron covered with a partial anti-Mackay crust to form a ball-and-cup structure. ⁵⁶ Perfect symmetry is achieved when m = 45.

With very few exceptions, the global minima of 100-atom particles belong to some type of icosahedral family. Exceptions are found at a few extreme Cu-rich compositions and for pure Ag, where the best structures are decahedral. All structures with the lowest mixing energy are icosahedral, and the best ones (belonging to clusters 3, 6 and 7) present a 55-atom Mackay icosahedron covered either by a Mackay or an anti-Mackay incomplete shell. These results show that icosahedral and polyicosahedral structures take advantage from stress relaxation in Cu@Ag structures much more efficiently than decahedral ones.

We note also that the transition between neighboring clusters of structures often corresponds to a change in slope or to an inflexion point in the mixing energy curve (see the ESI,† Fig. S2).

3.2 N = 200

The mixing energy profile, as a function of m (the number of silver atoms), along side with the SVR fit is shown in Fig. 5. The behaviour is similar to the first case, that is rather complicated. Mixing energy values are again negative, with very few exceptions that are found in the extreme Ag-rich limit, where there are some slightly positive values. The model was trained on the 80% of the data, and tested on the remaining 20%. The same set of hyperparameters for the case N = 100 was considered during 5-fold cross validation. The best triplet was found to be given by $\gamma = 0.001$, C = 50 and $\varepsilon = 0.01$. As before, the model was finally trained on the full training set, and tested on the test set, with the following scores:

$$R_{\text{training}}^2 = 0.999013 \quad R_{\text{test}}^2 = 0.997736$$

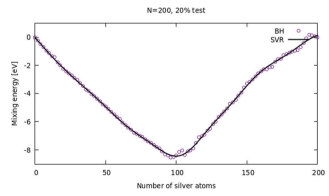


Fig. 5 Mixing energy profile for AgCu nanoalloys with N = 200 atoms in total, and SVR fit for the case of 80-20 split.

Other details regarding the SVR best model can be found in the ESI.† Similarly to the case N = 100, the data set was then split in 50% training set and 50% test set. For this setting the best triplet found is $\gamma = 0.005$, C = 10 and $\varepsilon = 0.001$. The two scores are again both very high:

$$R_{\text{training}}^2 = 0.999591 \quad R_{\text{test}}^2 = 0.998537$$

The model performance can be found in the ESI.†

The representation of these nanoalloys in the same two dimensional space described by the 422 and 55 CNA signatures is given in Fig. 6A.

When K-means is implemented, and the silhouette score is plotted as a function of the number of clusters K, one obtains the plot given in Fig. 7A. The optimal number of clusters, according to this criterion, is then K = 5. The Voronoi tessellation, along with each cluster center, is shown in Fig. 6B. A representative structure for each family is shown in Fig. 7B. Further details and images are given in the ESI.† The different structural families can be described as follows

• (1) m = 0-30, m = 196-200

Marks decahedra. In Cu-rich decahedra, Ag atoms fill the vertices first, then they start filling (100) facets. In Ag-rich decahedra, the few Cu are atoms placed in the fivefold axis.

 \bullet (2) m = 104-108

Core-shell FCC-HCP fragments.

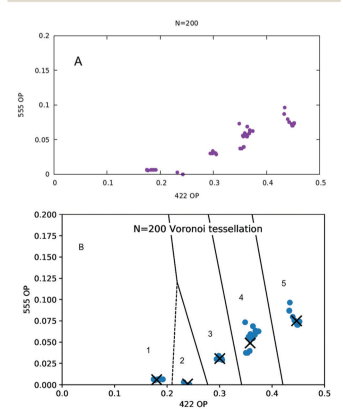


Fig. 6 (A) Data set representation in the two dimensional space of 422 and 555 order parameters. For each nanoalloy, we calculated its signature 422 and 555 order parameters as the fractions of nearest-neighbor pairs presenting such signatures. (B) Voronoi tessellation and cluster centers (means), for K = 5, when N = 200.

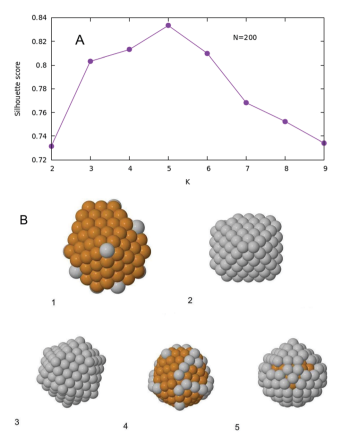


Fig. 7 (A) Silhouette score as a function of K for the case N = 200. (B) Five representative structures, one for each family. Labels are referred to Fig. 6B. The five structures have 10, 106, 168, 42 and 80 silver atoms respectively.

• (3) m = 160-174

Core-shell asymmetric icosahedra.

• (4) m = 32-54, m = 102, m = 110-152, m = 176-194

Incomplete core-shell structures, with a Cu icosahedron of N = 147 atoms covered by a Mackay Ag crust. Ag atoms initially fill the vertices, and later the edges. Finally they become part of the surface of the core 147 icosahedron. Again as in the case with N = 100, this is the largest cluster of structures.

 \bullet (5) m = 56-100, m = 154-158

As group (4), but with an anti-Mackay crust.

Also for size 200, a vast majority of global minima is of icosahedral structure. The exceptions are the decahedra found for Ag-rich and Cu-rich compositions and few FCC-HCP structures at intermediate compositions. These results confirm the key role of stress relaxation in determining the most stable structures also at this larger size of 200 atoms.

Alternative choices for the clustering of structures

Here we analyze how clustering into structural families is sensitive to the choice of the nanoparticle descriptors, i.e. of our 555, 422 and 421 variables. This analysis was made by K-means, which was applied to different choices of variables besides the (522, 422) choice used so far.

First of all, we applied the clustering algorithms in the threedimensional (3D) space of (555, 422, 421) signatures. Then we applied clustering in the spaces of two signatures at a time: (555, 421) and (422, 421) besides (555, 422), and finally to 1D spaces of one signature at a time, i.e. (555), (422) and (421) separately. We compared all clustering choices finding the following results:

- For size 100, only the pair (555, 422) was able to reproduce exactly the same clustering as the 3D description, while (555, 421) and (422, 421) were giving somewhat different clusters. On the other hand, the clustering obtained by 1D descriptions was significantly different.
- For size 200, only the pair (422, 421) was able to reproduce the same clustering as the 3D description. At variance with the (555, 422) description, (422, 421) gave six clusters instead of five, due to the splitting of cluster 4 of Section 3.2 into two new clusters corresponding to Cu-rich and Ag-rich parts. These two parts differ by a small distortion of the Ag crust in the Ag-rich part which eliminates the few 421 signatures that are present in Cu-rich nanoparticles.

From these results it turns out that (555, 422) and (422, 421) are the best choices to capture the physically relevant clusters of the full 3D description. However we note that the (422, 421) pair would be unable to separate well nanoparticles with fivefold symmetries and nanoparticles with extended hcp domains or several stacking faults. The latter were not present in our samples of global minima for sizes 100 and 200, but they may appear in samples including also higher energy isomers. For this reason we prefer the (555, 422) pair as the most useful 2D description. We note that this procedure of selecting the minimal set of variables for clustering of structures can be generalized to multi-dimensional cases including many CNA signatures and also other variables. Work in this direction is in progress.

Analysis of N = 100, m = 64, n = 36

Here we consider a specific composition for N = 100, Ag₆₄Cu₃₆, which is in the composition range with the lowest mixing energy. Here we consider all structures collected in the output of our three BH searches. This does not correspond to the much larger set of all visited structures during optimization, because these structures were collected by dividing the space of the 422 signature into small intervals of width 0.01 for the first two simulations and 0.002 for the third and looking for the lowestenergy structure for each interval. In total, our pool consists of 309 structures. In the usual (555, 422) 2D space given by the CNA signatures, the structures are arranged as shown in Fig. 8A. Given the nature of such representation (evidently more complicated than the previous two cases), we used the Gaussian mixture model in order to find a separation into structural families, because the Gaussian mixture model allows clusters to have ellipsoidal shapes (a more general and flexible hypothesis with respect to the hard tessellation found by K-means). The BIC and silhouette scores are then plotted as a function of the number of mixture components, K, in Fig. 9(A) and (B). The minimum of the BIC scores dictates that the optimal separation happens when the number of components

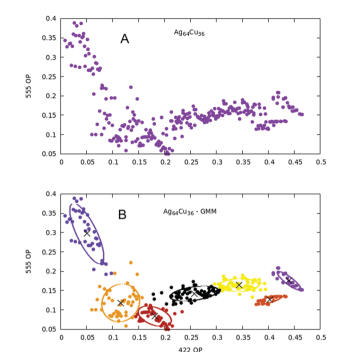


Fig. 8 (A) Data set representation in the two dimensional space of 422 and 555 order parameters. (B) Gaussian mixture model plot with centers and contours for K = 7 distributions

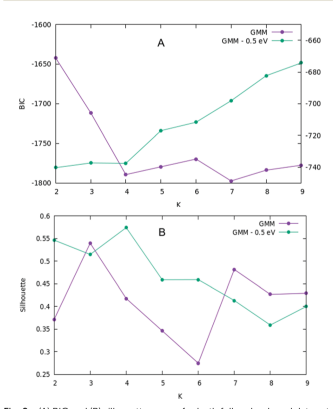


Fig. 9 (A) BIC and (B) silhouette scores for both full and reduced data sets. In (A) the curve for the reduced data set has the y-axis range given on the right part of the plot.

is seven. This optimal number is confirmed by the silhouette score which shows a local maximum at K = 7. However we must

recall that the silhouette score is less and less representative as the elongation of clusters increases. Apparently then, there is a clear distinction between seven different structural families. However, after manual inspection, one finds out that the first three clusters on the left (i.e. the orange, purple and red one) comprehend a large variety of amorphous structures, making very difficult the distinction between them, even for human eye. The reason behind this is that the pool of structures is composed of various local minima. among which a large part (in fact the majority) is very high in energy with respect to the global minimum, and thus represents a sort of noise of non-relevant structures. A simple way to overcome this obstacle is to put an energy cutoff.

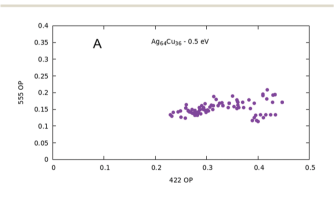
If we consider only structures which differ at most 0.5 eV from the global minimum, we obtain another representation of such nanoalloys, as shown in Fig. 10A.

With such restriction, the amount of structures reduces to 90. BIC and silhouette score suggest (as expected) four clusters to be the optimal subdivision of the reduced data set, as shown in Fig. 9(A) and (B). A representative structure for each family is instead depicted in Fig. 11.

When the HSA is implemented, the probability for these four structural families as a function of temperature can be calculated, as shown in Fig. 12. Specifically, the relative probability P of family \mathcal{F} is given by

$$P = \frac{\sum\limits_{i \in F} p_i}{n_{\min}}$$

$$\sum\limits_{i \in F} p_i$$
(17)



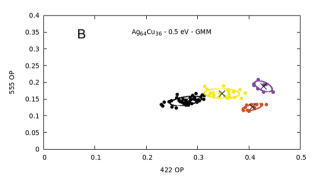


Fig. 10 (A) Data set representation in the two dimensional space of 422 and 555 order parameters, when the cutoff for energy is applied. (B) Gaussian mixture model plot with centers and contours for K = 4distributions.

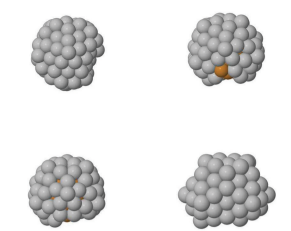


Fig. 11 Representative structures for the four clusters of Fig. 10B. Top left is an icosahedron with a Mackay crust (black cluster), top right is an icosahedron with mixed Mackay and anti-Mackay crust (yellow cluster), bottom left is an icosahedron with full anti-Mackay crust (purple cluster) and finally bottom right is a polyicosahedron (red cluster).

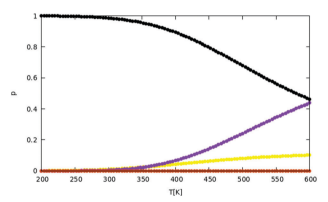


Fig. 12 Probability P of the different structural families as a function of temperature in the range 200-600 K. P is calculated by eqn (17). The curves refer to icosahedra with Mackay crust (black), icosahedra with mixed Mackay and anti-Mackay crust (yellow), icosahedra with full anti-Mackay crust (purple) and polyicosahedra (red). The colors correspond to those of the clusters of Fig. 10B, while representative structures are given in Fig. 11.

where the sum in the numerator is restricted to the minima belonging to family \mathcal{F} , while the sum in the denominator extends to the minima of all families considered in the calculation. The probabilities p_i of individual minima are given by eqn (16). In this way we can compute the probability for each motif as a function of temperature. The black curve in Fig. 12 suggests that the icosahedron with Mackay crust is the most favorite structural family (i.e. the most probable) in the relevant interval of temperature from 0 to 500 K (after which AgCu nanoalloys start melting their surface), coherently with the previous discussion on the structural properties of global minima at all compositions for AgCu nanoalloys with N = 100atoms in total. Icosahedra with anti-Mackay crust are ranked second (purple curve), whereas mixed crust (yellow) and polyicosahedra (red) are ranked respectively third and fourth.

Paper

4 Conclusions

In this article, the lowest-energy structures of AgCu nanoalloys of sizes N = 100 and 200 were searched for by the Basin Hopping global optimization algorithm. For N = 100, all compositions were considered, whereas for N = 200 we optimized one composition over two. These searches revealed a rich variety of structural motifs and a non-parabolic behaviour of the mixing energy.

The mixing energy was found to be negative, for all compositions at size 100 and for the vast majority of compositions of size 200, with very few slightly positive values in the extreme Ag-rich limit. We attribute the negative values of the mixing energy to the possibility of forming non-crystalline structures with core shell Cu@Ag chemical ordering, which is the optimal chemical ordering due to the larger size and lower surface energy of Ag compared to Cu. This chemical ordering, which is possible at the nanoscale with no counterpart in the bulk limit, allows an efficient stress relaxation which helps in stabilizing nanoalloy structures with respect to elemental nanoparticles.32,35 In fact, the structures with the lowest mixing energy are found in the composition range where the inner part and the surface layer are almost completely made of Cu and Ag atoms, respectively. The most efficient stress relaxation is achieved by structures of the icosahedral or polyicosahedral families, even for sizes, such as 100 and 200, that are far from icosahedral magic numbers. The stabilization of icosahedra by stress relaxation was already found for nanoparticles of sizes below 50 atoms, in several systems (AgCo, AgNi, AgCu and others).^{27,32,35} Here we have shown that stress relaxation is still very effective in stabilizing icosahedral structures at significantly larger sizes, and for wide composition intervals.

The global optimization data were used as a benchmark for the use of machine-learning techniques with the aim of answering to the following questions:

- 1. Is it possible to accurately fit the mixing energy by a suitable expression?
- 2. Is it possible to automatically group the nanoalloy structures into physically meaningful families?

As regards question 1, our calculations showed that the SVR algorithm is able to produce very accurate fits, for both N = 100and N = 200. These results indicate that the same procedure is likely to produce accurate functions for interpolating the mixing energy also in cases in which the search of the global minima for all compositions might be very cumbersome (think for example about nanoalloys of sizes 500 or 1000), so that interpolating on the basis of the results of a limited set of compositions may be quite useful. We note also that the interpolation procedure by the SVR method includes a crossvalidation of the result of the fitting that allows to determine the reliability of the fit itself.

As regards question 2, thanks to the clustering algorithms, we successfully divided each data set in distinct groups, thus underlying the variety of structural families present in AgCu nanoalloys of size N = 100 and N = 200 atoms. In particular we found that even without a previous manual classification of

such nanoalloys, it is still possible to recover a physically meaningful as well as detailed separation into different clusters of structures. This result was achieved also thanks to the clever description given by the CNA signatures, which made possible the success of the two algorithms used, in a two dimensional space. We remark the importance of our choice of using unsupervised learning algorithms, for it would be surely possible to train a supervised classification model on a manually labelled data set, but more time consuming and, at some point, subjective. In order to cluster the data sets of the global minima for N = 100 and 200, we used K-means which produced a hard partition (Voronoi tessellation) of the two dimensional space of the two order parameters relative to 422 and 555 signatures. The classification obtained by the (555, 422) pair was compared also to those obtained by using other CNA sets of variables, discussing how the classification depended on the choice of the set.

A different algorithm, the Gaussian mixture model, was instead used to perform the same task on a slightly more complicated case, that is the one offered by the two dimensional representation of some of the local minima found in the global minimization of Ag₆₄Cu₃₆. Here the superior flexibility of such algorithm with respect to K-means was crucial to find a good separation into clusters.

The usefulness of the clustering of the minima of Ag₆₄Cu₃₆ was then demonstrated by calculating the equilibrium probabilities of the different structural families depending on temperature. Finally, we note that the approaches developed in this work do not rely on any specific feature of the AgCu system, so that they can be easily used to treat other nanoalloys.

Conflicts of interest

There are no conflicts to declare.

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