High Throughput Discovery and Design of Strong Multicomponent Metallic Solid Solutions

Francisco G. Coury1, Kester D. Clarke1, Claudio S. Kiminami2, Michael J. Kaufman1 & Amy J. Clarke1

High Entropy Alloys (HEAs) are new classes of structural metallic materials that show remarkable property combinations. Yet, often times interesting compositions are still found by trial and error. Here we show an “Effective Atomic Radii for Strength” (EARS) methodology, together with different semi-empirical and first-principle models, can be used to predict the extent of solid solution strengthening to discover and design new HEAs with unprecedented properties. We have designed a Cr45Ni27.5Co27.5 alloy with a yield strength over 50% greater with equivalent ductility than the strongest HEA (Cr33.3Ni33.3Co33.3) from the CrMnFeNiCo family reported to date. We show that values determined by the EARS methodology are more physically representative of multicomponent concentrated solid solutions. Our methodology permits high throughput, property-driven discovery and design of HEAs, enabling the development of future high-performance advanced materials for extreme environments.

Multicomponent equiatomic alloys have garnered considerable interest in the literature over the past decade. One of the main focus areas is to fundamentally understand the high strengths and toughnesses exhibited by some of them1-4. Although the properties are extremely promising4, the compositional landscape (i.e., the number of possible compositions), is massive, and an almost infinite number of compositions is possible. One of the so-called “core effects”5,6 for HEAs suggests that the total compositional complexity (the configurational entropy) correlates directly with strength in single phase alloys. Indeed, several single phase quinary or senary alloys display attractive mechanical properties7-10. However, single phase alloys with fewer elements have proven to be stronger than the “parent”, higher entropy alloys that contain 5 elements or more6. A notable example is the ternary equiatomic Cr33.3Co33.3Ni33.3 alloy3,11-13, which is stronger and tougher than the quinary equiatomic Cr20Mn20Fe20Ni20Co20 alloy. To date, this Cr33.3Ni33.3Co33.3 alloy is the strongest and toughest face centered cubic (FCC) single phase solid solution from the quinary CrMnFeNiCo system out of hundreds of possible compositions, including several conventional stainless steels. The ternary Cr33.3Co33.3Ni33.3 alloy, which is the benchmark for strength in this work, was discovered in a study that involved producing all possible equiatomic alloys from the CrMnFeNiCo family with an FCC structure13. This alloy was effectively discovered by trial and error13.

The paradigm of HEA alloy design has shifted significantly since their inception6. The concept was mostly based on the aforementioned four “core effects”, where multicomponent, equiatomic HEAs are single phase and exhibit high strengths. Nowadays, these core effects have been challenged6. The main interest in HEA alloy design lies in the flexibility of these compositions with respect to phase stability and microstructural design14. If high-throughput tools are developed for predicting properties as a function of composition15,16, it then becomes possible to perform targeted alloy design, which means designing an alloy with ideal property combinations for a given application. Since there are endless possible alloys to be discovered, it is almost impossible to think there is not a composition best suited for a given application that exists today. Experimental mapping is prohibitive, as surveying the entire composition space would involve the production and characterization of thousands of samples. Finding idealized alloys cannot be a product of luck and extensive trial-and-error experimentation; the compositional landscape is too big for the time and resources that humankind has available. Targeted alloy design must be used to unlock the potential of new HEA compositions and properties for performance in extreme environments.

1Center for Advanced Non-Ferrous Structural Alloys, George S. Ansell Department of Metallurgical and Materials Engineering, Colorado School of Mines, Golden, CO, 80401, USA. 2Departamento de Engenharia de Materiais, Universidade Federal de São Carlos, Rodovia Washington Luís, km 235, São Carlos, SP, 13565-905, Brazil. Correspondence and requests for materials should be addressed to A.J.C. (email: amyclarke@mines.edu)
Although the high-throughput idea is not new for HEAs\(^{(4,7)}\), CALPHAD simulations, for example, are a high-throughput method for predicting phase equilibria that are used in HEA alloy design\(^{(8,10,18)}\). Alloying\(^{(9)}\) and microstructural evolution have also been linked to strength, but we could not find an example of targeted alloy design for the prediction of HEA solid solution strengthening, the key strengthening mechanism in these alloys, in which a significantly stronger composition was designed based upon prediction. Studies exist that show stronger compositions are not necessarily the ones with the highest entropy, and these alloys are also not necessarily equiatomic\(^{(21)}\). Varvenne et al. have suggested the strongest compositions should be those with the highest volumetric misfit between the atoms and/or the ones with the highest shear moduli\(^{(22,23)}\). If these ideas are correct, it should be possible to design significantly stronger alloys than existing ones if accurate atomic radii and elastic constants can be determined. By modifying two existing models in the literature with our new EARS methodology, we designed and made a single-phase FCC alloy (from the CrMnFeCoNi family) significantly stronger than the Cr\(_{1/3}\)Co\(_{1/3}\)Ni\(_{1/3}\) alloy.

Regardless of the mechanism for strain-hardening after yielding, the yield strength determines the baseline strength potential for a given alloy and is controlled by two primary factors: grain size and composition. Refining the grain size always results in higher strengths\(^{(40,41)}\), and can be accomplished via thermomechanical processing (i.e., rolling, forging, etc.). On the other hand, solid solution strengthening is not as straightforward, and is linked to atomic size and elastic moduli mismatch\(^{(24)}\). The former is usually considered the most important. Two general models have been published in the literature to predict the strength of multicomponent solid solutions. One model is proposed by Varvenne et al.\(^{(22)}\) for FCC solid solutions, while the other applies to FCC and BCC structures and is proposed by Toda-Carballo (TC)\(^{(25,26)}\). The Varvenne model is derived from first principles, whereas the second model is semi-empirical and adapts an equation valid for concentrated binary solid solutions. Both models predict composition-yield strength trends, and are thus suitable for high-throughput modeling. The final formulation of these models considers only the size mismatch contribution for strengthening. Therefore, the outcomes of both models are extremely dependent on atomic size input. The elastic constants are also important. The EARS methodology developed here provides accurate and physically significant values of these two properties.

**Results**

Strength predictions by EARS and the TC and Varvenne models guided the selection of four experimental HEAs in this work. One alloy (Alloy B) is the ternary equiatomic Cr\(_{0.33}\)Co\(_{0.33}\)Ni\(_{0.33}\) alloy, which serves as the benchmark for the strongest single-phase FCC alloy reported to date\(^{(3,4,27)}\) from the CrMnFeCoNi family. The reasons for selecting the other alloys will be further discussed below. The tensile stress vs. strain behaviors of the four alloys are shown in Fig. 1. Our EARS methodology predicts the strength to increase from Alloy A to D. With EARS, yield strength of each of the four alloys is predicted. Alloys C and D are 13 and 53% stronger, respectively, than Alloy B. These numbers rise to 25 and 70%, respectively, if we consider only the solid solution strengthening contribution, which can be determined by extracting the other contributions from the total yield strength, as described in the Methods section and shown visually in Fig. 1. Alloy B (the benchmark) has a remarkable combination of strength and toughness\(^{(3,11,27,28)}\), and has been suggested as a potential candidate for cryogenic applications, due to\(^{(2)}\) nanotwinning during deformation. The same nanotwins are observed in Alloys C and D, two of the stronger alloys, as shown in Fig. 1. A simple estimate of toughness, calculated by the area under the stress-strain curve, suggests that the strongest Alloy D has higher toughness than Alloy B, which to our knowledge represents the toughest HEA alloy ever produced (including HEAs outside the CrMnFeCoNi family).\(^{(19)}\) As mentioned previously, this composition was arrived at by prediction using the EARS methodology. Although there are several alloys in the literature with higher yield strengths than that of the Cr\(_{45}\)Co\(_{27.5}\)Ni\(_{27.5}\) alloy designed here, our alloy is a single-phase, coarse grained FCC solid solution, which typically would be expected to have low strength. The absolute yield strength of this alloy can be significantly increased, for instance by grain size reduction, as discussed later. In this work, we attempted to limit this effect. The most impressive property of Alloy D is not its absolute yield strength, but the solid solution strengthening component.

The predicted strength values using the EARS methodology can be seen in Fig. 2, in which pseudo-ternary phase diagrams are presented. The rationale behind picking Alloys A and D is that they are in the same compositional dimension as Alloy B (Cr\(_{3-x}\)Ni\(_{50-x}\), Co\(_{50-x}\)), but with different amounts of lattice distortion due to Cr content (Cr is a larger atom). The EARS methodology predicts increasing strength from Alloys A to D, with Alloy D as the strongest alloy. The fact that subsequent experimentally measured strength increases in order from Alloy A to D strongly supports the methodology used here. The strength calculations also predict that lattice distortion can be increased by adding small amounts of Mn, as shown in Fig. 2, with a resulting increase in yield strength. By adding Mn and removing Ni and Co, the lattice distortion is increased, since we approach a 50% ratio of big and small atoms (Alloy C). This occurs since Mn is bigger than both Co and Ni, but is smaller than Cr (Table 1). Also, Cr is determined to have a larger elastic modulus than Mn in the FCC structure, as shown in the Supplementary Materials section, while Cr-containing alloys have higher shear modulus than Mn-containing ones. On the other hand, Fe is smaller than Mn, so the same effect is observed at reduced magnitude, which explains our selection of Mn over Fe. In the plot of configurational entropy over the same composition space in Fig. 2, it is clear that the composition with the maximum configurational entropy, which is the equiatomic mixture between four elements (Alloy C), deviates completely from that predicted to have the maximum strength using the EARS methodology. This contradicts the original definition of HEAs\(^{(8)}\).

The robustness of the EARS methodology is further shown by comparing experimental and predicted strengths for a large number of alloys reported in the literature\(^{(19)}\), calculated with different atomic radii from Table 1. These results are presented in Fig. 3 for both the TC and Varvenne models. The main reason why EARS is so effective in the prediction of the solid solution strengthening is how it is determined. It is not trivial to determine accurate atomic radii for concentrated, multicomponent solid solutions. They can be determined by first principles methodologies, as performed by Okamoto\(^{(30)}\). This atomic radii set is called here the “Okamoto Radii”.

### Table 1

| Alloy | Composition | EARS Predicted Yield Strength | Experimental Yield Strength |
|------|-------------|------------------------------|-----------------------------|
| A    | Cr\(_{2/3}\)Co\(_{2/3}\)Ni\(_{2/3}\) | 3.4 GPa | 3.4 GPa |
| B    | Cr\(_{0.33}\)Co\(_{0.33}\)Ni\(_{0.33}\) | 4.5 GPa | 4.5 GPa |
| C    | Cr\(_{0.25}\)Co\(_{0.25}\)Ni\(_{0.25}\)Fe\(_{0.25}\) | 5.2 GPa | 5.2 GPa |
| D    | Cr\(_{0.125}\)Co\(_{0.125}\)Ni\(_{0.125}\)Fe\(_{0.125}\) | 6.0 GPa | 6.0 GPa |
They can also be determined by experimentally evaluating the lattice parameters of the pure materials, the “Pure Metals Radii” or by extracting the atomic radii from solid solutions and extrapolating the atomic size of the pure components, the “Solutions Radii”. The atomic radii calculated from first principles, or the “Okamoto Radii” and the “Solutions Radii” are more accurate in calculating the experimentally observed strengths of these alloys, but the strength values can still be overpredicted by as much as 100%. The EARS methodology predicts the strength of all the experimental alloys with the least error, for both the TC and Varvenne models. The EARS methodology significantly increases the accuracy of the predictions for all of the alloys examined, including our experimental alloys and those from the literature.

Efforts were made in this work to assure that all of the compositions were single-phase solid solutions with equivalent grain sizes. As noted previously, grain size may significantly impact strengthening. The four alloys were cast, rolled and annealed, as described in the Methods section. To ensure the FCC single-phase nature of the final microstructures after annealing, the alloys were investigated by XRD and electron backscattered diffraction (EBSD) as shown in Fig. 4 and, for Alloys C and D, transmission electron microscopy (TEM), as shown in Fig. 1 and also in the Supplementary Materials (Fig. S1-2). These analyses confirmed a single-phase solid solution for all four alloys presented here, and that Alloys B, C and D have similar grain sizes. The Hall-Petch (grain size) influence on the strength of these alloys was subtracted from the strength predictions, as described in the Methods section. This is paramount for the accuracy of the predictions, since the TC and the Varvenne models both predict the solid solution strengthening component.

**Discussion**

Considering the CrMnFeCoNi family, only Ni is FCC at room temperature. The exact atomic size changes based upon the crystal structure and the elements in solution. For this reason, the “Pure Metals Radii” will be inaccurate in a solid solution, which will compromise strength model predictions based on lattice distortion. Radii predicted
by Density Functional Theory (DFT), or the “Okamoto Radii”, will also have a reasonable degree of uncertainty, due to various factors such as the accuracy of the database, temperature effects and magnetism. The “Solutions Radii” is potentially the most accurate, but reasonable error still exists when predicting the radii of multicomponent solutions, as will be shown. For the EARS method, instead of directly inputting the atomic radii into the TC and Varvenne models, available strength values from the literature were used to calculate the best atomic radii values that match the experimental data. We assume the main factor for solid solution strengthening is indeed atomic size mismatch and the models considered here are accurate. The output are the EARS values, but the EARS methodology also includes the grain size contribution and the elastic moduli of all compositions (see Methods section).

Figure 2. Pseudo-ternary phase diagrams in at% for the Ni$_{50}$Co$_{50}$-Mn-Cr system. (a) The 1130 °C isotherm predicted by thermodynamic simulations using the software Thermo-Calc$^\text{TM}$ and the TCHEA1 database. (b) Predicted strengths from the TC model and EARS atomic radii (Table 1). (c) Calculated configurational entropy plotted over the same composition space. Clearly, the compositions with maximum configurational entropy do not correspond to those predicted (and shown) to have the maximum strength. The four alloys produced in this study are plotted in all three diagrams. The compositions are given in the legend. The predicted strengths for the alloys A-D are 223, 235, 243 and 258 MPa, respectively, considering only the $\sigma_{ss}$ contribution.
lattice parameters by X-ray (XRD) diffraction of the 4 alloys produced in this work. The lattice parameters deter-
achieve a high degree of accuracy, since very small changes in an atomic radii set will significantly change the
be necessary. The caveat is that experimental lattice parameter determinations must be performed with care to
here, although we believe that further understanding of short range order in concentrated solid solutions may
measured lattice parameters obtained from multicomponent alloys, instead of from strength values as was done
in concentrated multicomponent alloys. This also suggests that EARS radii might also be derived from carefully
mined by EARS are even more accurate than the Solutions Radii, showing that the EARS values are not only more
accurate for the prediction of strength, but are also more physically significant for representing atomic sizes
in concentrated multicomponent alloys. This also suggests that EARS radii might also be derived from carefully
measured lattice parameters obtained from multicomponent alloys, instead of from strength values as was done
here, although we believe that further understanding of short range order in concentrated solid solutions may
be necessary. The caveat is that experimental lattice parameter determinations must be performed with care to
achieve a high degree of accuracy, since very small changes in an atomic radii set will significantly change the
strength predictions. EARS of HEAs is a powerful methodology that predicts lattice strain and solid-solution strengthening. It
was shown that the EARS methodology was applied in conjunction with thermodynamic and strength models
to improve compositions and strength predictions. The atomic radii are also more physically significant for
concentrated solid solutions. Here we apply these ideas for the first time to perform targeted alloy design in the
multicomponent landscape, with the goal of maximizing solid solution strengthening. The resulting alloy
was 52% stronger than the strongest multicomponent alloy measured from the CrMnFeCoNi family, or 75%
stronger if only the solid solution strengthening contribution is considered. This work represents a first step
toward property-driven alloy discovery and design of multicomponent alloys. Given the framework of the meth-
ology, it is also possible to include additional property predictions in the future to target optimal compositions
for a given application (e.g. strength plus oxidation resistance). The results from this work highlight a new
opportunity for the development of strong alloys with different combinations of properties, driving us closer
toward high-throughput, property-driven alloy design needed to realize the full potential of HEAs for enhanced
performance in extreme environments.

### Methods

The four alloys were prepared by non-consumable arc melting from pure elemental components (99.2% Cr, 99.9%
Fe, 99.9% Mn, 99.9% Co and 99.9% Ni). The buttons were flipped and remelted four times to ensure chemical
homogeneity; the weight loss of each ingot was determined by making measurements before and after prepa-
ration, and was always less than 0.5%. Two buttons from each composition were cross rolled to a thickness of
2.5 mm (~70% cold work), annealed at 1130 °C for 2 h, and then cold-rolled an additional 40% to approximately
1.5 mm. Tensile samples were cut from the resulting plates by electro-discharge machining (EDM), annealed at
1130 °C for 30 min and then water quenched.

X-Ray Diffraction (XRD) was performed with a PanAnalytical Empyrean diffractometer using Mo Kα, radia-
tion for background signal minimization. Electron Backscattered Diffraction (EBSD) data was collected with a FEI
Helios FESEM using an accelerating voltage of 30 kV; the samples were prepared by grinding through 1200 grit SiC
papers, polishing sequentially with 6, 3 and 1 μm diamond suspensions for 5 minutes each, and then polishing for 4 h
with a Vibromet vibratory polisher and 0.05 μm colloidal silica. The diffraction patterns and inverse pole figure maps
are presented as raw data, without any refinement or cleaning performed. For the lattice parameter determination, a
Si standard was run at the exact same height as the samples to correct for potential systematic errors. Lattice param-
eters were calculated using the software HighScore Pro. The deviation of the Si lattice parameter was used to baseline
the measured lattice parameters of all the experimental alloys produced in this work.

For TEM analysis, samples were taken from the uniform elongation region of the gauge length of the tensile
samples after deformation. They were prepared by grinding to 100 μm and then electropolishing with a 10% sulfuric
acid in methanol solution at 30 V and around –20 °C using a Fischione dual jet instrument. The annealed Alloy B
sample shown in Fig. S1-2 in the Supplementary Materials was prepared by Focused Ion Beam (FIB) with the FEI
Helios and a final cleanup step at 2 kV. The TEM was performed using a FEI Talos microscope operating at 200 kV.

| Atomic Radii (pm) | Cr     | Mn     | Fe     | Co     | Ni     |
|------------------|--------|--------|--------|--------|--------|
| Pure Metal Radii | 124.90 | 130.55 | 124.12 | 125.10 | 124.59 |
| Okamoto Radii    | 126.90 | 123.50 | 121.90 | 121.90 | 123.90 |
| Binary Solid Solutions | 129.44 | 130.60 | 128.81 | 125.26 | 124.59 |
| TC-EARS (This work) | 129.25 | 127.52 | 126.81 | 124.46 | 123.28 |
| Varvenne-EARS (This work) | 130.09 | 129.41 | 127.86 | 124.30 | 122.66 |

Table 1. Atomic radii of the elements used as inputs to the TC model. The “Pure Metal Radii” were calculated
using the lattice constants of the pure metals in their original crystal structures, as given by Pearson’s
handbook26 for Cr, Fe, Co and Ni. The Mn value is given by Linus Pauling37 for the room temperature crystal
structure. For both Cr and Mn, since the nearest neighbors do not have the exact same distances, the total radii
were calculated as half of the weighted average of the nearest neighbor interatomic spacings. The “Okamoto
Radii” were calculated by first-principles for CrMnFeCoNi alloys30. The “Solutions Radii” were extracted form
binary FCC solid solutions by Varvenne27. The “TC-EARS” were calculated in this work. The fact that the TC-EARS values differ from the Varvenne-EARS values is expected, since the setup of each model is different. Using either models yields with EARS provides better strength predictions than any of the other atomic radii sets.
A standard sub-size ASTM E8 tensile sample geometry was used. Tensile tests were performed at a strain rate of $10^{-3}$ s$^{-1}$, up to failure with a 25.4 mm (1-inch) gage, 50% extensometer. A total of two samples were extracted from each plate of each alloy, resulting in a total of 4 samples per alloy. All the tensile curves are shown in Fig. S1-3 in the Supplementary Materials.

**Figure 3.** Comparison between the experimental and predicted yield strengths for the four alloys studied in this work using (a) the Toda-Caraballo Method and (b) the Varvenne method, plus additional alloys from the literature, determined with three different sets of atomic radii (Table 1). The prediction using the “Pure Metal Radii” data is poor, primarily because the size of the Mn atoms is larger than the other atoms. Therefore, all the compositions with Mn are predicted to be strong, whereas the other compositions are predicted to be weak. This is possibly the result of the uncommon crystal structure of Mn at room temperature, which most likely does not represent the atomic bond length that Mn has when it is in an FCC structure with other elements in a concentrated solid solution. Using the “Okamoto Radii”, it is possible to predict the strength vs. composition trends for these alloys; however, the values are generally overpredicted by a factor of 2 or 3. The EARS methodology developed here provides the most accurate strength predictions of the alloys. The alloys from this work are indicated by their respective letter A full table with the compositions plotted in this figure is provided in the Supplementary Materials section. (c) The calculated atomic radii of the four alloys produced in this work are compared to experimentally measured lattice parameters determined by XRD (see Methods section). EARS values are not only more accurate in performing strength predictions, but also give the most accurate lattice parameter predictions for HEAs, suggesting EARS values are more physically significant.

**Figure 4.** Characterization of as-tested Alloys A-D. (a) XRD patterns of the four alloys indicates that each is single-phase FCC solid solution. (b) Electron back scattered diffraction (EBSD) inverse pole figure maps are provided, along with an orientation color legend. As can be seen, Alloy A has a slightly more uniform grain size distribution than the other alloys, with an average grain size of approximately 20 µm; the other three alloys have a similar, but slightly wider, distribution of grain sizes, centered around 10 µm. The scale bar corresponds to 100 µm.
The thermodynamic simulations were performed using Thermo-Calc™ software with the TCHEA1 and Gheno31 (Fig. S1-1 in the Supplementary Materials) databases.

The Toda-Caraballo model (TC) was used to calculate the strength and lattice distortion. The first calculation necessary to implement the model is the lattice parameter (a) of the alloy. It can be calculated using the following equation (1).

\[ a = \frac{1}{f} \begin{pmatrix} x_1 & \ldots & x_n \end{pmatrix} \begin{pmatrix} s_{11} & \ldots & s_{1n} \\ \vdots & \ddots & \vdots \\ s_{n1} & \ldots & s_{nn} \end{pmatrix} \begin{pmatrix} x_1 \\ \vdots \\ x_n \end{pmatrix} \]

(1)

where \( f \) is the packing factor of the crystal structure, \( x_i \) is the atomic fraction of element \( i \), and \( s_{ij} \) is the interatomic spacing between the atom \( i \) and atom \( j \). This interatomic spacing is estimated by equation (2).

\[ s_{ij} = \frac{(2r_i)^2 K_i x_i + (2r_j)^2 K_j x_j}{2r_i K_i x_i + 2r_j K_j x_j} \]

(2)

where \( r_i \) is the atomic radii of atom \( i \) and \( K_i \) is the bulk modulus. The strength of the alloy is estimated as given by equation (3).

\[ \sigma = \sigma_0 + M Z \mu \left( \frac{r_i a}{\xi} \right)^{\frac{2}{3}} \]

(3)

The \( \sigma_0 \) term is a base strength for the material and was considered to be 60 MPa based upon literature values for pure Ni13 and \( M \) is the Taylor factor. A value of 3 was used here25,26. \( Z \) is a fitting parameter used by TC, and \( \alpha \) is a parameter that accounts for the nature of the dislocations. The TC model uses 16 to account for a mixture of edge and screw dislocations. The \( \delta^* \) is the estimated lattice distortion, and can be calculated by equation (4).

\[ \delta^* = \begin{pmatrix} x_1 \\ \vdots \\ x_n \end{pmatrix} \begin{pmatrix} 0 & \frac{da}{x_1} & \ldots & \frac{da}{x_n} \\ \frac{da}{x_1^2} & 0 & \ldots & \frac{da}{x_n} \\ \vdots & \vdots & \ddots & \vdots \\ \frac{da}{x_n} & \frac{da}{x_1} & \ldots & 0 \end{pmatrix} \begin{pmatrix} x_1 \\ \vdots \\ x_n \end{pmatrix} \]

(4)

where each term in the matrix estimates the distortion caused by substituting one element from the mixture for another. The simplest way of calculating each of these terms is by considering how much the lattice parameter changes by making a \( dx \) change in composition. Such as given by equation (5).

\[ \frac{da}{x_i} = \begin{pmatrix} x_1, x_2, \ldots, x_n \end{pmatrix} \begin{pmatrix} s_{11} & s_{12} & \ldots & s_{1n} \\ s_{21} & s_{22} & \ldots & s_{2n} \\ \vdots & \vdots & \ddots & \vdots \\ s_{n1} & s_{n2} & \ldots & s_{nn} \end{pmatrix} \begin{pmatrix} x_1 \\ \vdots \\ x_n \end{pmatrix} \]

\[ -(x_1 - \delta x, x_2 + \delta x, \ldots, x_n) \begin{pmatrix} s_{11} & s_{12} & \ldots & s_{1n} \\ s_{21} & s_{22} & \ldots & s_{2n} \\ \vdots & \vdots & \ddots & \vdots \\ s_{n1} & s_{n2} & \ldots & s_{nn} \end{pmatrix} \begin{pmatrix} x_1 - \delta x \\ x_2 + \delta x \\ \vdots \\ x_n \end{pmatrix} \frac{1}{\delta x} \]

(5)

The value of \( \delta x \) can be a small value. Here, 0.001 was used.

For determining the EARS values, the TC equation was used to calculate the yield strength of all the alloys in Table S1-1. The atomic radii of the elements were varied until the difference of the squares between the calculated strength and experimental strength from the literature13,32 were minimized. The “Solutions Radii” were used as a starting point, and the maximum allowable variation of the atomic radii was set to be 3%.

The Varvenne et al. model22,23 is derived from first principles. In this model, the interaction energy between a dislocation and a solute atom is calculated. The interaction energy is then inputted into a conventional equation to account for thermal aid to overcome the activation energy. The end result is a model for the critical resolved shear stress (converted into yield strength by the Taylor factor) that accounts for strain rate and temperature dependence. The final formulation considers only size misfit contributions, the activation energy for moving a dislocation (\( \Delta E_a \)) and the Peierls Stress at 0 K (\( \tau_0 \)) as follows in equations (6) and (7).
The $\sigma_0$ term is the proportionality constants between the dislocation line tension and $Gb^2$, with $G$ being the shear modulus and $b$ the Burgers vector. The value $0.123$ was used following Varvenne. The two $f(w_c)$ functions are called minimized core coefficients, and they account for the non-straight character of the dislocations, that bow out to find local energy minima. The values used were $0.35$ and $5.70$ for $f_1(w_c)$ and $f_2(w_c)$ respectively. The locking parameter of 250 MPa $\mu$ was reported locking parameters, so some alloys only had values extracted from hardness tests. The locking parameter is $\Delta \Gamma_n$, which is the average volumetric misfit of each atom, calculated as the volume of the $n$th atom minus the average atomic volume of that mixture. The atomic volume is calculated from the atomic radii given in Table 1. The atomic volume is the volume of the FCC unit cell divided by four, so it is not the atomic volume considering a hard sphere model. The two terms calculated from the equation above are then plugged in the following equation (8) for calculating the yield strength.

$$\tau_f(T, \dot{\epsilon}) = \sigma_0 \exp \left( -\frac{1}{0.51} \frac{kT}{\Delta E_b} \ln \frac{\tau_f}{\dot{\epsilon}} \right)$$

In the equation above, $k$ is the Boltzmann constant and $\dot{\epsilon}_0$ is a reference strain rate state set as $10^{-4}$ s$^{-1}$, again following Varvenne. All the studies performed in this work used a strain rate ($\dot{\epsilon}$) of $10^{-3}$ s$^{-1}$. The calculations were performed to evaluate the strength at 293 K.

For all the elastic constants used as input to both the TC and the Varvenne models, the actual elastic constants measured from ultrasonic techniques reported in the literature were used. For the high-throughput predictions (Fig. 2), the elastic constants were extrapolated from the shear and Poisson moduli to what would be expected from pure single-phase FCC Cr, Mn, Fe and Co. This was performed by fitting a first order polynomial for both moduli values as a function of composition for all of the 10 different alloys used in this work. The final elastic moduli used for a composition were calculated using equations (9) and (10)

$$G_a = 103.5 x_{Cr} + 81.0 x_{Mn} + 51.7 x_{Fe} + 81.0 x_{Co} + 76.0 x_{Ni}$$

$$\nu_a = 0.275 x_{Cr} + 0.056 x_{Mn} + 0.353 x_{Fe} + 0.293 x_{Co} + 0.310 x_{Ni}$$

These values were only used for the high-throughput calculations. Note that the elastic constants used for Ni are those from pure FCC Ni. The elastic modulus and bulk modulus were obtained from these two elastic constants using standard conversion equations.

Both the Varvenne and TC models predict only the solid solution strengthening contribution. Therefore, all other strength terms were subtracted from the experimental values. This practice is commonly performed. The total strength of an alloy was considered to be given by equation (11).

$$\sigma = \sigma_0 + \sigma_{HP} + \sigma_d$$

$\sigma_0$ is a base strength term explained in the following section, $\sigma_{HP}$ is the solid solution strengthening term calculated by the TC and the Varvenne models, and $\sigma_{HP}$ is the grain size contribution, accounted for using the Hall-Petch (HP) equation (12).

$$\sigma_{HP} = k d^{-1/2}$$

The constant $k$ is material dependent and is usually called the locking parameter; $d$ is the grain size. The $k$ values inputted into the HP equation were extracted from the literature. Not all the alloys produced in this work had reported locking parameters, so some alloys only had values extracted from hardness tests. The locking parameter seems to escalate with Cr content. Two alloys with Cr–Ni and Ni$_{60}$Co$_{40}$ have a k of 180 MPa $\mu$. The quinary Cr$_{20}$Mn$_{20}$Fe$_{20}$Co$_{20}$Ni$_{20}$ has a k of 226 MPa $\mu$ for the alloys.

The constant $k$ is the Boltzmann constant and $\dot{\epsilon}_0$ is a reference strain rate state set as $10^{-4}$ s$^{-1}$, again following Varvenne. All the studies performed in this work used a strain rate ($\dot{\epsilon}$) of $10^{-3}$ s$^{-1}$. The calculations were performed to evaluate the strength at 293 K.

For all the elastic constants used as input to both the TC and the Varvenne models, the actual elastic constants measured from ultrasonic techniques reported in the literature were used. For the high-throughput predictions (Fig. 2), the elastic constants were extrapolated from the shear and Poisson moduli to what would be expected from pure single-phase FCC Cr, Mn, Fe and Co. This was performed by fitting a first order polynomial for both moduli values as a function of composition for all of the 10 different alloys used in this work. The final elastic moduli used for a composition were calculated using equations (9) and (10)

$$G_a = 103.5 x_{Cr} + 81.0 x_{Mn} + 51.7 x_{Fe} + 81.0 x_{Co} + 76.0 x_{Ni}$$

$$\nu_a = 0.275 x_{Cr} + 0.056 x_{Mn} + 0.353 x_{Fe} + 0.293 x_{Co} + 0.310 x_{Ni}$$

These values were only used for the high-throughput calculations. Note that the elastic constants used for Ni are those from pure FCC Ni. The elastic modulus and bulk modulus were obtained from these two elastic constants using standard conversion equations.

Both the Varvenne and TC models predict only the solid solution strengthening contribution. Therefore, all other strength terms were subtracted from the experimental values. This practice is commonly performed. The total strength of an alloy was considered to be given by equation (11).

$$\sigma = \sigma_0 + \sigma_{HP} + \sigma_d$$

$\sigma_0$ is a base strength term explained in the following section, $\sigma_{HP}$ is the solid solution strengthening term calculated by the TC and the Varvenne models, and $\sigma_{HP}$ is the grain size contribution, accounted for using the Hall-Petch (HP) equation (12).

$$\sigma_{HP} = k d^{-1/2}$$

The constant $k$ is material dependent and is usually called the locking parameter; $d$ is the grain size. The $k$ values inputted into the HP equation were extracted from the literature. Not all the alloys produced in this work had reported locking parameters, so some alloys only had values extracted from hardness tests. The locking parameter seems to escalate with Cr content. Two alloys with Cr–Ni and Ni$_{60}$Co$_{40}$ have a k of 180 MPa $\mu$. The quinary Cr$_{20}$Mn$_{20}$Fe$_{20}$Co$_{20}$Ni$_{20}$ has a k of 226 MPa $\mu$ for the alloys.
with the higher values of k for samples with higher Cr contents, as this is the sample with the highest k and the highest Cr content.

The experimental strength of pure Ni reported by Wu et al. is about 80 MPa for a grain size of 85 μm. After extracting the HP contribution for this material (k = 180 MPa μm⁻¹/₂), 60 MPa is still unaccounted for. This is the σ₀ term that accounts for the base strength of Ni at room temperature, although this value is likely slightly higher than what would be expected for a pure single-phase FCC metal. Since all the alloys used in the EARS determination procedure were characterized in the same study by Wu et al., this σ₀ term also accounts for any systematic experimental deviations characteristic of their specific setup.

Data availability. All data generated or analyzed during this study are included in this published article (and its Supplementary Information files).

References
1. Tracy, C. L. et al. High pressure synthesis of a hexagonal close-packed phase of the high-entropy alloy CrMnFeCoNi. Nat. Commun. 8, 1–6 (2017).
2. Zhang, F. et al. Polymorphism in a high-entropy alloy. Nat. Commun. 8, 1–7 (2017).
3. Gludovatz, B. et al. Exceptional damage-tolerance of a medium-entropy alloy CrCoNi at cryogenic temperatures. Nat. Commun. 7, 10602 (2016).
4. Zhang, W., Liaw, P. K. & Zhang, Y. Science and technology in high-entropy alloys. Sci. China Mater. 61, 2–22 (2018).
5. Ye, J. W. Recent progress in high-entropy alloys. Ann. Chim. Sci. des Matérl. 31, 633–648 (2006).
6. Miracle, D. B. & Senkova, O. N. A critical review of high entropy alloys and related concepts. Acta Mater. 122, 448–511 (2017).
7. Okamoto, N. L. et al. Size effect, critical resolved shear stress, stacking fault energy, and solid solution strengthening in the CrMnFeCoNi high-entropy alloy. Sci. Rep. 6, 35863 (2016).
8. Couziník, J. P. et al. On the room temperature deformation mechanisms of a Ti50HfNbTa refractory high-entropy alloy. Mater. Sci. Eng. A 645, 255–263 (2015).
9. Y. A. H. et al. Mechanical properties of refractory high-entropy alloys: Experiments and modeling. J. Alloys Compd. 696, 1139–1150 (2016).
10. Senkova, O. N., Senkova, S. V. & Woodward, C. Effect of aluminum on the microstructure and properties of two refractory high-entropy alloys. Acta Mater. 68, 214–228 (2014).
11. Yoshida, S., Bhattacharjee, T., Bai, Y. & Tsuji, N. Friction stress and Hall–Petch relationship in CoCrNi equi-atomic medium entropy alloys processed by severe plastic deformation and subsequent annealing. Scr. Mater. 134, 33–36 (2017).
12. Laplanche, G. et al. Reasons for the superior mechanical properties of medium-entropy CrCoNi compared to high-entropy CrMnFeCoNi. Acta Mater. 128, 292–303 (2017).
13. Wu, Z., Bei, H., Pharr, G. M. & George, E. P. Temperature dependence of the mechanical properties of equiatomic solid solution alloys with face-centered cubic structural acts. Acta Mater. 81, 428–441 (2014).
14. Lu, Z., Pradeep, K. G., Deng, Y., Raabe, D. & Tisan, C. C. Metastable high-entropy dual-phase alloys overcome the strength–ductility trade-off. Nature 534, 227–230 (2016).
15. Ritchie, R. O. The conflicts between strength and toughness. Nat. Mater. 10, 817–822 (2011).
16. Lu, K. The Future of Metals. Science 328, 319–321 (2010).
17. Ye, Y. F., Wang, Q., Lu, J., Liu, C. T. & Yang, Y. High-entropy alloy: challenges and prospects. Mater. Today 19, 349–362 (2016).
18. Senkova, O. N. et al. Microstructure and elevated temperature properties of a refractory TaNbHfZrTu alloy. J. Mater. Sci. 47, 4062–4074 (2012).
19. Chen, H. et al. Contribution of Lattice Distortion to Solid Solution Strengthening in a Series of Refractory High Entropy Alloys. Metall. Mater. Trans. A Phys. Metall. Mater. Sci. 49, 772–781 (2018).
20. Senkova, O. N., Jensen, J. K., Pilchak, A. L., Miracle, D. B. & Fraser, L. H. Compositional variation effects on the microstructure and properties of a refractory high-entropy superalloy AlMo0.5 NbTa0.5 TiZr. Mater. Des. 139, 498–511 (2018).
21. Wilcoxon, P., Field, R. & Kaufman, M. The use of diffusion multiples to examine the compositional evolution of phase stability and hardness of the Co-Cr-Fe-Mn-Ni high-entropy alloy system. Intermetallics 75, 15–24 (2016).
22. Varvenne, C., Luque, A. & Curtin, W. A. Theory of strengthening in fcc high entropy alloys. Acta Mater. 118, 164–176 (2016).
23. Varvenne, C., Leysom, G. P. M., Ghazisaeidi, M. & Curtin, W. A. Solute strengthening in random alloys. Acta Mater. 124, 680–683 (2016).
24. Leysom, G. P. M., Curtin, W. A., Hector, L. G. Jr & Woodward, C. F. Quantitative prediction of solute strengthening in aluminum alloys. Nat. Mater. 9, 750–755 (2010).
25. Toda-Caraballo, I. & Rivera-Díaz-Del-Castillo, P. E. J. Modelling solid solution hardening in high entropy alloys. Acta Mater. 85, 14–23 (2015).
26. Toda-Caraballo, I. A general formulation for solid solution hardening effect in multicomponent alloys. Scr. Mater. 127, 113–117 (2017).
27. Gludovatz, B. et al. A fracture-resistant high-entropy alloy for cryogenic applications. Science 345, 1153–1158 (2014).
28. Zhao, Y. L. et al. Heterogeneous precipitation behavior and stacking-fault-mediated deformation in a CoCrNi-based medium-entropy alloy. Acta Mater. 138, 72–82 (2017).
29. Lu, K., Lu, L. & Suresh, S. Strengthening Materials by Engineering Coherent Internal Boundaries at the Nanoscale. Science 324, 349–353 (2009).
30. Okamoto, N. L., Yuge, K., Tanaka, K., Inui, H. & George, E. P. Atomic displacement in the CrMnFeCoNi high-entropy alloy – A scaling factor to predict solid solution strengthening. AIP Adv. 6, 125008 (2016).
31. Gheno, T., Liu, X. L., Lindwall, G., Liu, Z.-K. & Gleeson, B. Experimental study and thermodynamic modeling of the Al-Co-Cr-Ni system. J. Alloys Compd. 665, 33–36 (2017).
32. Laplanche, G., Kostka, A., Horst, O. M., Eggeler, G. & George, E. P. Microstructure evolution and critical stress for twinning in the CrMnFeCoNi high-entropy alloy. Acta Mater. 118, 152–163 (2016).
33. Clarke, K. D., Comstock, R. J., Mataya, M. C., Tyne, C. J. & Matlock, D. K. Effect of strain rate on the yield stress of ferritic stainless steels. Metall. Mater. Trans. A Phys. Metall. Mater. Sci. 39 A, 752–762 (2008).
34. He, J. Y. et al. A precipitation-hardened high-entropy alloy with outstanding tensile properties. Acta Mater. 102, 187–196 (2016).
35. Wu, Z., Bei, H., Otto, F., Pharr, G. M. & George, E. P. Recovery, recrystallization, grain growth and phase stability of a family of FCC-structured multi-component equiatomic solid solution alloys. Intermetallics 66, 131–140 (2016).
36. Villars, P. & Calvert, L. D. Pearson’s Handbook of Crystallographic Data for Intermetallic Phases (American Society for Metals, 1986).
37. Pauling, L. Atomic Radii and Interatomic Distances in Metals. J. Am. Chem. Soc. 69, 542–553 (1947).
Acknowledgements
This work was supported by CNPq, Conselho Nacional de Desenvolvimento Científico e Tecnológico - Brazil (grant number 233746/2014-5) and the Center for Advanced Non-Ferrous Structural Alloys (CANFSA), a National Science Foundation Industry/University Cooperative Research Center (I/UCRC) (Award No. 1624836), at the Colorado School of Mines (CSM), in Golden, CO, USA, and the Office of Naval Research (Grant #12488790). We also thank Mr. John Copley (CSM) for TEM sample preparation support, Mr. Paul Mason from Thermo-Calc® for providing a software license, Mr. Paul Wilson from Boeing for sharing results that aided in the computational model setup, and Profs. Xerxes Steirer and Branden Kappes (CSM) for assisting with the XRD measurements.

Author Contributions
F.G.C., A.J.C., M.J.K., K.D.C. and C.S.K. conceived the research. F.G.C. carried out the experiments with guidance from the co-authors, analyzed the results, and developed the computational methods. F.G.C., A.J.C., K.D.C. and M.J.K. wrote the manuscript with input from all of the authors.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-018-26830-6.

Competing Interests: The authors declare no competing interests.

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018