Efficient construction of linear models in materials modeling and applications to force constant expansions

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Linear models such as force constant (FC) and cluster expansions, play a key role in physics and materials science. While they can in principle be parametrized using regression and feature selection approaches, the convergence behavior of these techniques, in particular with respect to thermodynamic properties is not well understood. Here, we therefore analyze the efficacy and efficiency of several state-of-the-art regression and feature selection methods, in particular in the context of FC extraction and the prediction of different thermodynamic properties. Generic feature selection algorithms such as recursive feature elimination with ordinary least-squares (OLS), automatic relevance determination regression, and the adaptive least absolute shrinkage and selection operator can yield physically sound models for systems with a modest number of degrees of freedom. For large unit cells with low symmetry and/or high-order expansions they come, however, with a non-negligible computational cost that can be more than two orders of magnitude higher than that of OLS. In such cases, OLS with cutoff selection provides a viable route as demonstrated here for both second-order FCs in large low-symmetry unit cells and high-order FCs in low-symmetry systems. While regression techniques are thus very powerful, they require well-tuned protocols. Here, the present work establishes guidelines for the design of protocols that are readily usable, e.g., in high-throughput and materials discovery schemes. Since the underlying algorithms are not specific to FC construction, the general conclusions drawn here also have a bearing on the construction of other linear models in physics and materials science.

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INTRODUCTION

Linear models such as force constant (FC) and cluster expansions are widely used in materials science, physics, and chemistry to describe the thermodynamic behavior of real materials. Their computational efficiency and mathematical simplicity are also appealing for applications in high-throughput calculations and machine learning, which requires methods for efficient and automatized model construction. In this context, regression techniques are particularly appealing as they promise to require fewer computationally demanding reference calculations than approaches based on systematic enumeration of configurations.

Regression techniques in combination with regularization have received a lot of attention for model building, often under the title compressive sensing (CS). The latter is in principle a task in sparse signal recovery that is usually approached by finding solutions to an underdetermined linear system. The problem of solving the linear system is, however, completely independent of CS and itself is not a solver.

The usefulness of regression with regularization for the construction of physical models has been demonstrated, using the least absolute shrinkage and selection operator (LASSO) as well as the split-Bregman technique. There is, however, a much larger pool of potentially useful regression techniques, including various other forms of regularization and feature selection. These models involve one or several hyperparameters the choice of which often has a very direct impact on the results (as shown extensively below). With applications in high-throughput computations but also more conventional situations in mind, it is therefore necessary to conduct a careful analysis of these techniques for the construction of physical models.

The vibrational degrees of freedoms (DOFs) of materials are crucial for numerous thermodynamic properties, including phase stability and thermal conduction. To model these properties one requires an efficient representation of the potential energy surface (PES). In crystals the vibrational atomic motion can be conveniently described in terms of phonons, quasi-particles that represent periodic and quantized excitations. Phonon theory is commonly formulated by starting from a Taylor expansion of the total energy, in which the expansion coefficients are referred to as FCs. Depending on material and property of interest the FC expansion must be carried out to different orders. Generally, it is preferable to keep the order as low as possible since the number of independent coefficients quickly increases with expansion order, decreasing symmetry, and number of sites in the unit cell.

For ideal materials with comparably small unit cells the FCs up to third order can still be obtained by enumerating displacements and evaluating the derivatives numerically. This direct enumeration scheme becomes, however, tedious or impractical for larger systems (e.g., point defects, interfaces or nanoparticles) and/or materials that require expansions beyond third order (e.g., metastable phases of transition metals or oxides). Accordingly linear regression techniques have been applied including ordinary least squares (OLS), LASSO, and split-Bregman. As noted above, there are various other linear regression techniques and feature selection algorithms that could be suitable for FC regression such as recursive feature elimination (RFE), automatic relevance determination regression (ARDR), and adaptive LASSO.
(ad-LASSO). Further analysis of these techniques with regard to their efficiency, accuracy and reliability for constructing FC expansions is therefore in order. 

Here, we use the HIPHIVE package since it is interfaced with machine-learning libraries such as SCIKIT-LEARN that in turn provide efficient implementations of various optimization techniques. In this paper, we present a comparison of linear regression methods and the direct enumeration approach for the extraction of FCs of different order, including second-order FCs for large systems of low symmetry such as defects, third-order FCs for the prediction of the thermal conductivity, as well as higher-order FCs for bulk and surface (see Supplementary Information) systems. This approach enables us to determine the applicability of regression methods in different regimes. We also demonstrate the application of these FC models for studying anharmonic effects, both in the framework of Boltzmann transport theory and molecular dynamics (MD) simulations. The following section provides a concise summary of the underlying theory, while sections thereafter present the different application examples named above.

RESULTS
FC extraction
The PES can be expanded in a Taylor series in the atomic displacements \( \mathbf{r}_0 \) relative to a set of reference positions \( \mathbf{r}_0 \):

\[
V = V_0 + \Phi^{(2)} u_i^2 + \frac{1}{2} \Phi^{(3)} u_i u_j u_k + \frac{1}{6} \Phi^{(4)} u_i u_j u_k u_l + \ldots,
\]

where \( \Phi \) are the FCs, Latin indices enumerate the atoms, Greek indices enumerate the Cartesian coordinates, and the Einstein summation convention applies.

The number of FC components scales as \( \mathcal{O}(N^2) \), where \( N \) is the number of atoms and \( n \) is the expansion order. There are, however, multiple constraints that reduce the number of free parameters, such as lattice symmetries and sum rules. Yet in the case of large systems, low symmetry, and/or higher expansion orders the number of parameters is still very large.

Direct approach. The conventional way of extracting FCs relies on the systematic evaluation of numerical derivatives. For example, for the second-order terms

\[
\Phi^{(2)}_{ij} = \frac{\partial^2 V}{\partial u_i \partial u_j} \approx \frac{F_i^0 - \Phi^{(0)} u_i^2}{\Delta u_i^2},
\]

where \( F_i^0 \) denotes the force on atom \( i \) along \( \alpha \) and \( \Delta u_i^\alpha \) is a small displacement of atom \( j \) along \( \beta \), typically between 0.01 and 0.05 Å. This direct approach is implemented in several software packages, including PHONOPY for second-order and PHONOPPY for third-order FCs as well as ALAMODE for an arbitrary expansion order. This method has been used with great success for predicting vibrational properties of many common materials. The number of reference calculations, however, quickly becomes a limiting factor for systems with many sites in the unit cell, in the case of low symmetry, and/or higher-order FCs. In fact, it is usually impractical to compute any term beyond third-order using the direct approach except for rather simple cases.

Regression approach. The information density, here taken as the number of force components that are sizable, in supercells with only one or two displaced atoms such as the ones used in the direct approach is relatively low. Instead, one can consider general displacement patterns, involving many (or all) atoms in the supercell, and then employ regression techniques to reconstruct the underlying FCs. This approach has been shown to produce accurate higher order FCs, but can also be used to construct effective FC models.

The force acting on atom \( i \) along \( \alpha \) can be written as

\[
F_i^\alpha = -\Phi^{(0)}_{ij} u_j^\beta - \frac{1}{2} \Phi^{(2)}_{ijk} u_j^\beta u_k^\gamma - \ldots,
\]

which can be cast in linear form:

\[
F_i^\alpha = \mathbf{A} \cdot \mathbf{x}.
\]

Here, \( \mathbf{x} \) are the free parameters of the FC model while the rows of the fit matrix \( \mathbf{A} \) encode the displacements with symmetry transformations as well as constraints imposed by the sum rules. The vector comprising all forces in a supercell \( \mathbf{F} \) can thus be expressed as

\[
\mathbf{F} = \mathbf{A} \mathbf{x}.
\]

The construction of the fit matrix \( \mathbf{A} \) is described in ref. and can be trivially generalized to multiple reference structures.

Truncating the expansion. The number of free parameters, i.e., the dimension of \( \mathbf{x} \), can still be very large even for systems with high symmetry. The FC expansion is therefore often truncated. Firstly, as with most Taylor expansions, only few orders are usually needed to obtain an accurate representation of the PES in the range of relevant displacements. Secondly, the atomic interactions often decay rather quickly with interatomic distance, meaning a cutoff can be imposed. Thirdly, pair interactions are often stronger than three-body interactions, which in turn are often stronger than four-body interactions etc., i.e.,

\[
||\Phi_{ij}|| > ||\Phi_{ijk}|| > ||\Phi_{ijk}||.
\]

Linear regression techniques. Equation (1) can be solved by minimizing the objective function \( ||\mathbf{Ax} - \mathbf{F}_{\text{target}}||_2 \), where \( \mathbf{F}_{\text{target}} \) denotes the reference forces. In the overdetermined limit this can be achieved by OLS, which can, however, lead to overfitting. One should emphasize that the rows of the sensing matrix \( \mathbf{A} \) in Eq. (1) are always to some extent correlated due to atomic interactions as each row corresponds to the Cartesian force component of one atom in one structure. This should be kept in mind when using the terms overdetermined and underdetermined in the usual sense that is based on the relation between the number of rows and columns of the sensing matrix.

Overfitting can be overcome by regularization, i.e. inclusion of additional penalty terms in the objective function, usually related to the \( \ell_1 \) or \( \ell_2 \) norm of the solution vector. For example in the case of elastic net regularization one has

\[
\mathbf{x}_{\text{opt}} = \arg\min_{\mathbf{x}} \left\{ ||\mathbf{Ax} - \mathbf{F}_{\text{target}}||_2^2 + \alpha ||\mathbf{x}||_1 + \beta ||\mathbf{x}||_2^2 \right\}.
\]

With \( \alpha = 0 \) this expression reduces to Ridge regression, while with \( \beta = 0 \) one recovers the objective function for the LASSO method. As discussed below, LASSO is known to over-select features. To overcome this deficiency the ad-LASSO approach has been proposed, in which the regularization term is modified to

\[
\mathbf{x}_{\text{opt}} = \arg\min_{\mathbf{x}} \left\{ ||\mathbf{Ax} - \mathbf{F}_{\text{target}}||_2^2 + \alpha \sum_i w_i ||x_i|| \right\},
\]

where \( w_i \) are individual weights for each parameter. Here, we employ the iterative update described in ref.

To evaluate the performance of a model obtained by solving Eq. (1) one can employ cross-validation (CV). To this end, the available reference data set is split into training and validation sets. After the former has been used for fitting the parameter vector, one can evaluate the root-mean-square error (RMSE)

\[
\text{RMSE} = \sqrt{\frac{1}{N} \sum_i (\mathbf{F}_{\text{model}} - \mathbf{F}_{\text{target}})^2}.
\]
where the summation extends over the $N$ force components in the validation set. To reduce the statistical error, the RMSE is then averaged over several different splits of the reference data, yielding the CV score.

Efficient and generally applicable implementations of these methods are available, e.g., via the Python machine learning library SCIKIT-LEARN17.

**Feature selection.** In machine learning feature selection refers to the task of isolating the most important parameters (or features) during model construction. Reducing the number of parameters yields a less complex model, which in turn often leads to less overfitting and improved transferability. It can also reduce the computational cost of sampling the model. Feature selection is especially interesting for FC models, for which only some interaction terms may be of importance.

Several feature selection methods are available for linear problems. One can employ for example a simple pruning condition based on the magnitude of the parameters. This is particularly effective in combination with regression techniques that include regularization, typically via the $\ell_1$ or $\ell_2$-norm of the parameter vector (see Eq. (2)). One can also employ matching pursuit algorithms such as orthogonal matching pursuit, which allows one to impose a constraint on the number of non-zero coefficients in the solution vector, or techniques such as RFE. In the latter case, a solution is determined using a fit method of choice, the weakest or least important parameters are removed, and the procedure is iteratively repeated until the target number of features is reached. In some cases, the optimal sparsity of a model can be determined by combining the above techniques with Bayesian optimization6.

It must be noted that the different methods can differ considerably with respect computational effort as well as memory requirements. OLS is the least demanding procedure in both regards. It is difficult to provide general guidelines with respect to the demands of different methods, since the effort can differ dramatically with the choice of hyperparameters and the conditioning of the sensing matrix. As a rough guideline, RFE with OLS typically requires about 100–1000 OLS fits depending on how accurately one wishes to perform the feature elimination. LASSO is comparable to RFE-OLS with respect to computational effort.

Second-order FCs: Large low-symmetry systems
The second-order FCs of systems with many atoms and/or low symmetry can be tedious to obtain by the direct approach. This applies in particular to the FCs of defect configurations, which are needed for example for computing the vibrational contribution to the free energy of defect formation19, analyzing the impact of defects on the thermal conductivity 18,38,39 or predicting the vibrational broadening of optical spectra40,41. In this section, we therefore analyze the extraction of second-order FCs for the vacancy in body-centered cubic (BCC) Ta as a prototypical case, using both the direct approach as implemented in PHONOPY25 and the regression approach as implemented in HIPHIVE15.

**Scaling of regression methods.** Several different methods were considered for constructing FC models for the Ta vacancy models, including OLS, RFE-OLS, LASSO, and ARDR. OLS is by far the computationally least expensive method and exhibits a favorable scaling with training set size (including both computational effort and memory requirements), which prevents its effective application for large sensing matrices including the present case.

Taking into account CV, the computational effort required for the largest supercells considered here becomes notable for LASSO and RFE-OLS. In the remainder of this section, we therefore primarily consider OLS, which will be demonstrated to work very well if combined with cutoff selection to avoid overfitting.

Furthermore, it is possible to tune parameters such as the rate at which the number of features is reduced in the RFE algorithm in each step, which effectively reduces the pre-factor of RFE. The present comparison focuses, however, on the ability of these methods to recover physically correct solutions and their convergence behavior, whence aspects concerning their computational performance are not further explored here.

**Cutoff selection via cross-validation.** The number of DOFs in a FC model grows rapidly as the cutoff increases. At the same time, as discussed above, interaction strength and hence the magnitude of the FCs decay with increasing interatomic distance. At some point an increase in cutoff will therefore lead to negligible improvement in accuracy but merely an increase in model complexity. Specifically in the absence of regularization terms in the objective function ($\alpha = \beta = 0$ in Eq. (2)), one can therefore observe a deterioration of model quality with the inclusion of more terms in the expansion due to overfitting. In this case one should therefore evaluate the performance of models with different cutoffs.

We employed CV using the shuffle-and-split method with 5 splits and 15 training structures for a system size of $N = 6$ and constructed a series of second-order FC models with increasing cutoffs (Fig. 2). While the RMSE over the training set continues to decrease with increasing parameter space, the CV-RMSE has a minimum around 6–7 Å. For standard OLS the validation score increases for larger cutoffs due to overfitting. This behavior can be counteracted by using RFE, which yields a slight improvement of the CV-RMSE. As discussed above RFE is, however, computationally substantially more expensive whence OLS with a judicious choice of cutoffs is preferable. All subsequent analysis was therefore carried out using OLS and a second-order cutoff of 6 Å.

![Figure 1](image-url)  
**Fig. 1 Computational cost of different optimization algorithms.** Relative timings for the computational effort for carrying out a single optimization of second-order force constant models for a vacancy in Ta ($N = 6$) with a cutoff of 6.0 Å using different regression techniques. Time is normalized using the time for ordinary least squares (OLS) with a training set size of 5000 force components (~3 s). Calculations were carried out on an Intel Xeon E5-2650 V3 processor with 10 cores (20 threads). ARDR automatic relevance detection regression, LASSO least absolute shrinkage and selection operator, OLS ordinary least squares, RFE recursive feature elimination.
where parameters in the model. Comparison with the measure based about 3000 force components, which is equivalent to two to convergence behavior. Both quantities reach convergence at

\[ \| \Delta \Phi \| = \left\| \Phi_{\text{regression}} - \Phi_{\text{direct}} \right\| / \left\| \Phi_{\text{regression}} \right\| \]  

(6)

where \( \Phi \) denotes the Frobenius norm.

The frequencies and free energies exhibit very similar convergence behavior. Both quantities reach convergence at about 3000 force components, which is equivalent to two to three configurations and corresponds to the number of parameters in the model. Comparison with the measure based on the FC matrix itself, Eq. (6), suggests that \( \Delta \Phi \leq 5\% \) is sufficient to achieve convergence of the frequency spectrum and the free energy. Considering the convergence of \( \Delta \Phi \) itself suggests a more conservative threshold of 2%.

The comparison includes both models with only second-order FC terms and models with additional very short-ranged third-order FC terms using a cutoff of 3.0 Å. The latter perform consistently better than the second-order-only models. The inclusion of a few third-order terms thus stabilizes the extraction of the second-order FCs, an observation that has also been made in other situations. These terms enable one to account for anharmonicity in the vicinity of the reference positions that would otherwise be effectively included in the second-order FCs.

Third-order FCs: thermal conductivity

Calculating the thermal conductivity using the linearized Boltzmann transport equation requires knowledge of the second and

Convergence of thermodynamic properties. In order to evaluate how accurately the regression approach reproduces the correct second-order FCs, we considered three different measures. First, we evaluated the absolute error of the zone-center (I) frequencies obtained by regression relative to the direct approach (Fig. 3a)

\[ \Delta \omega = \sqrt{\frac{1}{3N} \sum \left( \omega_{\text{regression}} - \omega_{\text{direct}} \right)^2}, \]  

(4)

where \( \omega_i \) is the frequency of mode \( i \). Secondly, we considered the absolute difference in the harmonic free energy at 2250 K corresponding to 75% of the calculated melting temperature (Fig. 3b)

\[ \Delta F = \left| F_{\text{vib, regression}} - F_{\text{vib, direct}} \right|. \]  

(5)

Here, the free energies were computed within the harmonic approximation using PHONOPY. Lastly, in order to obtain a computationally cheaper measure, we computed the relative error of the second-order FC matrices (Fig. 3c), defined as follows:

\[ \Delta \Phi = \left\| \Phi_{\text{regression}} - \Phi_{\text{direct}} \right\| / \left\| \Phi_{\text{regression}} \right\| \]  

(6)

Scaling with system size. Following the analysis in the previous section, computing \( \Delta \Phi \) as a function of the training set size allows one to determine the number of training structures needed for recovering the second-order FCs at the accuracy level of the direct approach (Fig. 4). Using the more conservative threshold of \( \Delta \Phi < 2\% \) (Fig. 3), we thereby determined the necessary number of supercell calculations as a function of system size (Fig. 4).

While in the direct approach the number of necessary calculations increases steeply with system size, in the regression approach it is constant or decreases slightly with system size. This is possible due to the introduction of a cutoff but more importantly since the regression takes advantage of the increase in information content with supercell size. In this context, it is important that all atoms are displaced by least a small amount. By contrast, the information content in configurations employed in the direct approach decreases substantially with system size as only one atom is displaced at a time. One can anticipate this scaling effect to be even more pronounced for third or higher-order FCs due to the exponential increase of the number of parameters with order.
third-order FCs\(^4\), providing a sensitive test for the extraction of higher-order FCs. Here, we analyze different regression methods for obtaining FCs and the resulting thermal conductivity in silicon. Specifically, we consider OLS, which has been used for the same purpose in ref.\(^1\), LASSO, ad-LASSO, RFE-OLS, and ARDR as implemented in SCIKIT-LEARN. This comparison enables us to demonstrate the importance of studying convergence with respect to the choice of cutoffs and number of training structures as well as the selection of a suitable fit method.

Reference second and third-order FCs were calculated for 250-atom supercells (5 × 5 × 5 primitive unit cells) via the direct approach using PHONOPY\(^25\) and PHONO3PY\(^26\), respectively. No cutoff was imposed during the calculation of the third-order FCs, which therefore required 801 individual force calculations.

For the regression approach, we generated a total of 20 reference structures based on 250-atom supercells (5 × 5 × 5 primitive unit cells) with displacements drawn from a normal distribution yielding an average displacement amplitude of 0.03 Å. For the computation of CV scores we used the same splits throughout to enable a one-to-one comparison of the regression methods.

Since ARDR exhibits a stronger scaling with system size than the other methods (Fig. 1), the pruning hyper-parameter \(\lambda_t\) was not optimized but set to a constant value of \(\lambda_t = 10^4\).

For ARDR we first constructed a second-order-only model and then trained a second and third-order to the remaining forces. This \(\alpha\)-approach led to a significant improvement in accuracy and numerical stability for ARDR but did not improve the performance of the other fit methods.

For ad-LASSO the hyperparameter \(a\) was optimized once using CV and five training structures, after which the same value was used for all training set sizes. While one could thus possibly obtain slightly better results for larger training sets, the present choice allows a substantial reduction of the computational effort.

**Optimization with cutoff selection.** The number of DOFs associated with the second-order FCs is very small and thus we used the maximum cutoff range of 9.65 Å that the 5 × 5 × 5 supercells employed here can support. The nearest-neighbor fourth-order interaction, corresponding to a fourth-order cutoff of 2.5 Å, was included in order to improve the accuracy of the second and third-order FCs (as demonstrated in the second-order FC example below). The third-order cutoff was then treated as a tunable parameter.

When using five training structures the accuracy of the model is already converged for a third-order cutoff of 4.0 Å (Fig. 5a) regardless of fit method, which yields a total of 80 DOFs. RFE-OLS and ARDR have, however, the distinct advantage of selecting fewer parameters and thus avoid overfitting for large cutoffs (Fig. 5b).

Using a third-order cutoff of 4.0 Å, the thermal conductivity converges to within 2% of the PHONOPY values using as few as 14 structures (Fig. 5). For comparison, the PHONO3PY calculation requires 801 structures if no cutoff is imposed and 57 structures when including a pair cutoff of 4.0 Å (two neighbor shells). In the latter case, one must, however, take into account that convergence testing would require including at least one more shell, which increases the number of calculations to 95 (three neighbor shells).

For example, in the case of high-throughput studies one is often content with a less accurate estimate of the thermal conductivity. In this context, it is noteworthy that when using ARDR or RFE-OLS one converges to within 10% of the reference value already with four structures, while being able to test for convergence.

**Optimization with generic feature selection.** Cutoff selection can be thought of as a feature selection approach, in which the cutoffs are pruning hyperparameters. In the present case, in which we
They were, for example, used recently to evaluate alloy cluster parameters of the model. These types of measure may serve as a compliment to the CV score when evaluating FC models. Absolute shrinkage and selection operator, RFE recursive feature elimination.

focus on the third-order FCs we effectively obtain only one such hyperparameter, namely the third-order cutoff. The number of cutoff parameters increases with the expansion order and in more complex materials can require fine tuning e.g., between different atomic species or crystallographic sites. In such situations it can be advantageous to employ generic feature selection algorithms. In this section, we consider the suitability of LASSO, ad-LASSO, RFE-OLS as well as ARDR for this purpose. For comparison, we also include the performance of OLS is included. Cutoffs of 9.65, 9.65, and 2.5 Å for second, third, and fourth-order clusters, respectively, were used throughout, which yields 2525 DOFs.

In the case of RFE-OLS and ARDR the thermal conductivity converges to within 10% of the reference value using about five structures (Fig. 6b; see Supplementary Fig. 2 for temperature dependence). Moreover, with ARDR one achieves convergence within 2% with about 12 structures, which is even better than in the case of cutoff selection. In the case of LASSO and OLS the convergence rate is considerably lower and neither method achieves convergence with respect to the reference value with the number of structures considered here.

We note that based on the convergence of the CV-RMSE scores (Fig. 6a) and the comparison with the cutoff selection study (Fig. 5a), one could expect LASSO to yield a reasonably converged thermal conductivity when using about 15 structures. This is not the case, demonstrating that CV-RMSE scores alone are insufficient for assessing the quality of a model. Information criteria such as alkaline information criteria (AIC) and Bayesian information criterion (BIC) can be used to evaluate models, taking into account the predictive power but also penalizing the number of parameters of the model. These types of measure may serve as a useful compliment to the CV score when evaluating FC models. They were, for example, used recently to evaluate alloy cluster expansion models.

To explore the differences in performance between LASSO, RFE-OLS, and ARDR further we explicitly computed the CV-RMSE as a function of their respective pruning hyperparameter using five training structures, which allows us to obtain the variation of the CV score with the number of features (Fig. 7a). A notable difference is observed in the third-order FCs. While RFE-OLS and ARDR produce a small number of short-ranged interaction terms, LASSO yields a large number of spurious third-order FCs terms (Supplementary Fig. 2).

The over-selection in LASSO can be overcome by using, e.g., ad-LASSO (see Eq. (3)). This yields a learning curve comparable to ARDR (Fig. 6a) and a much smaller number of features compared to LASSO (Fig. 7a). Thereby, one obtains a small set of third-order FCs terms (Fig. 8) that properly reproduce the physical properties of the system (Supplementary Fig. 2).

It is striking that all techniques except for LASSO can achieve convergence of the thermal conductivity to within 5% with only five training structures (Fig. 7b). It is also noteworthy that the CV score alone is an unreliable predictor for model quality. This is especially apparent in the case of LASSO, for which the model with the smallest CV score leads to an underestimation of the
moreover has implications for the thermal conductivity 30,33,49.

required and the number of DOFs is very large.

as an example for a system, in which both higher-order FCs are

dependence of vibrational modes associated with Ba 49 and

In this section, we are concerned with the inorganic clathrate

Fourth-order FCs: Strong anharmonicity

We emphasize that the data sets used here are publicly

able choice. In the following, we therefore restrict our analysis to

ability of different optimization methods to recover the

correct third-order force constants. Norm of third-order force

constants in Si obtained using generic feature selection algorithms

and 10 training structures. Cutoffs of 9.65, 9.65, and 2.5 Å

were employed for second, third, and fourth-order clusters, respectively. The cluster radius is defined as $r_c = \sum_{i \in \text{cluster}} ||r_i - \bar{r}_g|| / N$ where $r_{g}$ is the geometrical center of the cluster (compare Supplementary Fig. 2). ARDR automatic relevance detection regression, LASSO least absolute shrinkage and selection operator, RFE recursive feature elimination.

thermal conductivity by 50%. Using the CV score alone for model

selection can hence be very misleading. For this purpose, one
could therefore also consider model evaluation metrics such as AIC and BIC—an aspect that deserves further study.

Finally, we note that overestimating the true number of features

leads to very large errors in the predicted thermal conductivity for all techniques. Underestimation on the other hand, i.e. overly sparse models, give much smaller errors, which indicates that over-regularization is the preferable mode of error.

We emphasize that the data sets used here are publicly available48 and can serve as a test bed for a systematic comparison with respect to other fit algorithms and feature selection methods.

Fourth-order FCs: Strong anharmonicity

In this section, we are concerned with the inorganic clathrate $\text{Ba}_8\text{Ga}_{16}\text{Ge}_{30}$, in which the motion of Ba atoms is strongly anharmonic49. This manifests itself in a strong temperature dependence of vibrational modes associated with Ba49 and moreover has implications for the thermal conductivity40,33,49. While perturbation theory formally provides an expression for the temperature induced phonon frequency shifts caused by the third-order FC terms14, one commonly carries the expansion at least to the next higher even order, when analyzing frequency shifts40,33,50. Since $\text{Ba}_8\text{Ga}_{16}\text{Ge}_{30}$ has a large unit cell, it thus serves as an example for a system, in which both higher-order FCs are required and the number of DOFs is very large.

Clathrates are inclusion compounds with a defined lattice structure that can trap atomic or small molecular species. $\text{Ba}_8\text{Ga}_{16}\text{Ge}_{30}$ belongs to the class of type-I clathrates with spacegroup $\text{Pm}$ $\text{3n}$51. In this case, the host lattice is made up of Ga and Ge atoms, which occupy Wyckoff sites 6c, 16i, and 24k, whereas Ba atoms reside inside the cages occupying Wyckoff sites 2a and 6d. Due to the size mismatch between guest species and cages, which is particularly large for the 6d sites, the Ba atoms experience a very wide and flat PES with pronounced anharmonicity. In earlier work, we analyzed the ordering of the host species and extracted the ordered ground state structure of $\text{Ba}_8\text{Ga}_{16}\text{Ge}_{30}$15, which serves as a prototype structure for the analysis of the FCs.

Model construction. First, we generated 50 structures based on the primitive 54-atom unit cell with an average displacement amplitude of 0.28 Å using the Monte Carlo rattle approach described in ref.15. These structures were used to train an initial fourth-order model, which was subsequently sampled by MD simulations at 300 and 650 K for 10 ps. We extracted 50 structures from each runs and generated reference forces via density functional theory (DFT) calculations. Thereby we obtained a total of 150 reference structures and thus a sensing matrix with 24,300 rows. Due to the large number of DOFs in this structure (Table 1) the computational effort that has to be expended for training models becomes very significant. For illustration, a OLS fit with 10-fold CV of the largest model considered below requires approximately one hour on an Intel Xeon E5-2650 V3 CPU. Given the scaling analysis above (Fig. 1) this translates to several days and more when using RFE-OLS or LASSO. Since the computational effort associated with ad-LASSO is yet another order of magnitude larger than for LASSO, we restricted ourselves to OLS, RFE-OLS, and (standard) LASSO, the latter of which has been previously used for a very similar structure23.

We constructed several different models and evaluated their performance by CV in order to identify a suitable combination of expansion order and cutoff parameters (Table 1).

The three optimization methods considered here yield virtually identical CV-RMSE scores (Fig. 9a), which is sensible as even for the largest cutoff parameters the number of DOFs is still significantly smaller than the number of rows in the sensing matrix (Table 1). RFE-OLS and to a very slight extent also LASSO yield a smaller number of features. In practice, one has to balance this advantage (which can also imply shorter run times for force evaluations of the final model) against the additional effort required for construction of the model. Based on the good scores that can already be obtained with OLS and the fact that these cases are commonly easy to move into the overetermined region, even for spaces as large as in the present case, one can argue that OLS is often the more natural choice. In the following, we therefore restrict our analysis to the OLS fits.

The smallest CV-RMSE score is obtained for model 4 (Fig. 9a), which yields a value of 68 meV/Å to be compared with maximum force components of about 4000 meV/Å (Fig. 9b). The fourth-order cutoff for this model is 4.35 Å, which is enough to include all Bag-cage interactions indicating that the anharmonicity of all of these interactions is important. The PES for Ba atoms along different directions calculated is in excellent agreement with DFT calculations (Fig. 10). It is apparent that Ba atoms in 6d sites behave more anharmonic than those in 2a sites, as the cages surrounding the former are larger.

Table 1. Fourth-order FC models for $\text{Ba}_8\text{Ga}_{16}\text{Ge}_{30}$ obtained by OLS using different combinations of cutoffs and expansion orders.

| Model | Two-body cutoffs | Three-body cutoffs |
|-------|------------------|--------------------|
|        | 2nd  | 3rd  | 4th  | 5th  | 6th  | 3rd  | 4th order |
| 1     | 5.4  | 3.0  | 3.0  |      |      |      |           |
| 2     | 5.4  | 3.5  | 3.5  |      |      |      |           |
| 3     | 5.4  | 4.0  | 4.0  |      |      |      |           |
| 4     | 5.4  | 4.35 | 4.35 |      |      |      |           |
| 5     | 5.4  | 4.7  | 4.7  |      |      |      |           |
| 6     | 5.4  | 4.35 | 4.35 | 3.0  | 3.0  |      |           |
| 7     | 5.4  | 4.35 | 4.35 | 3.0  | 3.0  | 4.0  | 4.0       |

Table 8 Ability of different optimization methods to recover the correct third-order force constants. Norm of third-order force constants in Si obtained using generic feature selection algorithms and 10 training structures. Cutoffs of 9.65, 9.65, and 2.5 Å were employed for second, third, and fourth-order clusters, respectively. The cluster radius is defined as $r_c = \sum_{i \in \text{cluster}} ||r_i - \bar{r}_g|| / N$ where $\bar{r}_g$ is the geometrical center of the cluster (compare Supplementary Fig. 2). ARDR automatic relevance detection regression, LASSO least absolute shrinkage and selection operator, RFE recursive feature elimination.
To analyze the behavior of model 4 further, we generated an ensemble of FC models that are trained in identical fashion but are based on different training sets. The latter were constructed by selection with replacement (bagging) from the available data set such that the number of reference forces in the training sets equals the total size of the data set. The RMSEs over the training set obtained for these FC models are similar to model 4. This ensemble of models enabled us to estimate the sensitivity for predictions for physical properties such as phonon frequencies (Fig. 11) and thermal conductivity (Fig. 12).

Thermal conductivity. The low thermal conductivity of inorganic clathrates is often attributed to the rattling motion of the guest species\(^{29,49}\). These modes exhibit a notable temperature dependence\(^{49}\) that needs to be accounted for in order to predict the thermal conductivity accurately\(^{30,33}\). The common approach to Boltzmann transport theory, however, only considers terms up to third-order and neglects the temperature dependence of the phonon frequencies. The fourth-order model allows us to investigate the temperature dependence of the frequency spectrum and include this effect when calculating the thermal conductivity via the temperature-dependent FCs approach\(^{30,34,53}\).

Yet instead of training effective third-order models from ab-initio MD simulations, we train them against snapshots and forces generated from MD simulations using the full fourth-order model. The latter simulations were carried out using a supercell of 2 × 2 × 2 primitive unit cells (432 atoms). The systems was first equilibrated for 10 ps using a Langevin thermostat as implemented in ASE\(^{54}\). Subsequently, the simulations were continued in the micro-canonical ensemble for another 5 ps. From the latter part, 100 snapshots were selected to train effective second and third-order FCs.

The density of states (DOS) obtained from the effective second-order FCs reveals a significant temperature dependence of the low-frequency Ba modes around 4 meV (Fig. 11), in line with experimental work\(^{30}\) and our previous study, which was based on ab-initio MD simulations\(^{30}\). The sensitivity analysis shows that the dependence of the DOS on model uncertainty is considerably weaker than the temperature dependence. We also note that the DOS has a slightly weaker temperature dependency if the harmonic FCs are trained without including the third-order FCs. This is in qualitative agreement with the negative frequency shift due to cubic FCs observed in ref.\(^{33}\).

The effective FCs were furthermore used to extract the thermal conductivity at the respective temperature they were trained using the linearized Boltzmann transport equation via SHENG\(^{27}\). The resulting thermal conductivity is in very good agreement with our previous calculations (Fig. 12), which were based on full ab-initio MD simulations and which in turn agrees well with experimental results\(^{30}\). We note that as shown in refs\(^{30,33}\), the thermal conductivity is strongly underestimated when using the temperature-independent (zero-K) second-order and third-order FCs as that approach fails to account for the...
strong renormalization with temperature of the lowest-
frequency heat carrying modes. As in the case of the DOS, the
model sensitivity analysis reveals a very weak variation in
thermal conductivity due to model uncertainty.

The very good agreement with the results obtained using
temperature-dependent interatomic FCs trained using ab-initio
MD simulations demonstrates that our fourth-order model
provides an accurate and transferable description of the PES in
the temperature range of interest. The effective models in the
present work, however, required only a fraction of the computer
time that was needed for the ab-initio MD simulations in ref. 30.
The present results are also in semi-quantitative agreement
with the results from ref. 33, which were obtained using a Green’s
function approach for treating phonon renormalization. The
latter calculations were carried using a different structural model
that does not correspond to a thermally accessible state, whence
one cannot expect quantitative agreement. There are further
methodological differences between the Green’s function
approach in ref. 33 and the present work. The latter takes into
account quantum statistic but only handles renormalization of
the second-order FCs, whereas the present approach is classical
but effectively accounts for higher-order terms via temperature-
dependent third-order FCs. 30

DISCUSSION

Regression techniques can reduce the computational effort
associated with FC extraction compared to enumeration techni-
cues by one or more orders of magnitude, in particular in the case
of higher-order expansions and/or large systems with low
symmetry. In this study, we have therefore assessed both
efficiency and efficacy of several regression methods in different
application scenarios. Specifically, we considered second-order
FCs and derived properties for large systems, third-order FCs, and
thermal conductivity as well as fourth and higher-order FCs and
their sampling for strongly anharmonic systems.

While the discussion below is explicitly based on the results
presented in the preceding sections, they are further supported by
our experience with various other materials including, e.g., metals,
oxides, carbides, and chalcogenides of varying dimensionality
including two and three-dimensional systems, interfaces, and
defects. Hence, we consider our conclusions to be applicable to
other materials and application areas to a reasonable extent.

Second-order FCs in large systems

For second-order FCs in large systems the regression approach
reduces the computational effort by more than one order of
magnitude compared to the direct approach (Fig. 4). OLS with
cutoff selection yields prediction errors that are on par with more
advanced regression techniques such as RFE-OLS, LASSO, or ARDR.
The latter are, however, at least one to two orders of magnitude
more demanding in terms of computer time, which can become a
concern for very large systems.

For OLS to work properly the linear system to be solved must be
overdetermined. The configurations used for regression can be
obtained by rattling the atomic positions. As a result, the
information density, i.e. the number of force components that are
sizeable, is high, which is usually not the case for enumerated
structures such as the ones used in the direct approach. As a
result, a much smaller number of configurations is required in
order to obtain a well conditioned sensing matrix.

We have furthermore found that inclusion of a few higher-order
FC terms (here third-order FCs with a short cutoff) accelerates
convergence of the lower-order FCs of interest with respect to
training set size. Here, the third-order terms allow extraction of the
"true" second-order expansion terms, which otherwise would have
to effectively account for anharmonicity in the PES.

Third-order FCs and thermal conductivity

The regression approach also drastically reduces the number of
reference calculations needed to recover the parameters of third-
order FC expansions. Here, care must be taken to verify not only
the convergence of the CV-RMSE scores with respect to the
references forces but to consider the actual property of interest, in
the present case the thermal conductivity.

As shown previously, OLS with cutoff selection provides a
viable route to obtaining well-converged thermal conductivity
values at a fraction of the computational cost of the direct
approach. Several generic feature selection methods provide,
however, viable alternatives that require adjusting a smaller
number of hyperparameters and are hence more easily extensible
to complex systems and higher-order expansions.

Here, RFE-OLS, ARDR, and ad-LASSO have been found to work
very reliably and efficiently, yielding both fast convergence and
a small number of features (sparse solutions). Standard LASSO
produces denser solutions, which leads to less predictive models
and has a detrimental impact when predicting the thermal
conductivity.

Higher-order FC models and anharmonic PESs

We also considered the construction of FC expansions beyond
third-order, which is usually impractical with enumeration
approaches. Specifically, we constructed fourth-order FC models
with up to more than 13,000 parameters for the inorganic
crystalline Ba8Ga16Ge30 using OLS with cutoff selection. Using the
final model effective FCs were generated and used to compute
the temperature dependent DOS and thermal conductivity,
yielding results in agreement with experiment as well as previous
computational work.

These results demonstrate that the "standard" and computa-
tionally relatively cheap OLS method can be used to capture
strongly anharmonic effects across a wide temperature range. It
can easily be extend to higher order anharmonicity and properties
other than the thermal conductivity, as shown by an eighth-order
model for a Ni surface, which allows one to model the
temperature dependence of the surface layer spacing (see
Supplementary Information).

To put the present results in context, we note that the benefits
of regression with regularization, typically discussed under the
heading of CS, have been particularly emphasized for systems
with many DOFs, which includes both the clathrate system (over 13,000 parameters) and the Ni surface (2000 parameters) considered here. While for other linear models, specifically cluster expansions, it can be difficult to generate that many reference data points, for FC expansions generating reference data is relatively easy and often simpler than tuning of model parameters and hyperparameters. For example, for the clathrate system, one structure provides 162 reference force components and hence already with less than a hundred such structures one obtains an overdetermined system.

One could argue that regularization would allow one to reduce the number of reference data points considerably, making techniques such as ad-LASSO, LASSO, or even ARDR competitive. In this regard, we point out that the RFE-OLS fits for the clathrate system show that a sensible model has about 4000–5000 parameters. Even with regularization, one should thus require approximately as many reference data points, which is often not a substantial enough reduction to warrant the substantially larger computational effort associated with regularization techniques.

Conclusions and outlook

Generic feature selection algorithms, in particular ARDR, RFE, and ad-LASSO can yield physically sound FC expansions at a fraction of the cost of enumeration approaches. This approach can be very powerful as demonstrated here for extracting third-order FCs and thermal conductivity. The application of (standard) LASSO approach is, however, not indicated due to its tendency to over-select, which leads to very slow convergence with training set size. For large unit cells with low symmetry and/or high-order expansions these techniques come, however, with a non-negligible cost that can be more than two orders of magnitude higher than that of OLS. The cost is still much smaller than those of DFT calculations but since the underlying problem is not as amenable to parallelization it can still become a factor to consider in practice. In such cases OLS with cutoff selection provides a viable route, with trivial parallelization over multiple cutoff parameter sets. The viability of the latter approach has been demonstrated here for both second-order FCs in large low-symmetry unit cells and high-order FCs in low-symmetry systems. Regression techniques are in principle very attractive for high-throughput schemes, since they require much less computational effort than enumeration approaches. For the regression approach to be viable one must, however, not only consider the computational effort but also the amount of human intervention required. In the future, it is therefore desirable to set up protocols that automatically construct and validate FC models. In this context, we have made the data related to the analysis of higher-order FC expansions publicly available\textsuperscript{64} to provide a standardized benchmark set for future work.

METHODS

Second-order FCs: Large low-symmetry systems

Calculations were carried out for supercells comprising $N \times N \times N$ conventional BCC cells with $N \in [4, \ldots, 10]$ that contain a single vacancy. Reference forces were computed using the embedded atom method (EAM) potential model TA1 from ref. 55. Using an empirical potential rather than DFT calculations in this example allows us to compute reference second-order FCs for very large configurations. For the PHONOPY calculations we used a displacement amplitude of 0.01 Å while for the HPHIVE calculations, we generated 30 structures for each $N$ by drawing random displacements from a normal distribution with a standard deviation of 0.01 Å.

Third-order FCs: Thermal conductivity

Reference forces were obtained from DFT calculations using the projector augmented wave method\textsuperscript{56,57} as implemented in VASP\textsuperscript{58,59} and an exchange-correlation functional\textsuperscript{60} based on the generalized gradient approximation. The Brillouin zone was sampled using only the Γ-point. The plane-wave energy cutoff was set to 245 eV, an additional support grid for fast-Fourier transformations was used during the force calculation, and the projection operators were evaluated in reciprocal space.

Fourth-order FCs: Strong anharmonicity

Reference forces were obtained for the 54-atom cells described below from DFT calculations using the projector augmented wave method\textsuperscript{56,57} as implemented in VASP\textsuperscript{56,57}. The exchange-correlation potential was represented using the vdW-DF-Cx method, which combines semi-local exchange with non-local correlation\textsuperscript{54,61,62}, as it has been previously shown to provide a good description of the vibrational modes of this system\textsuperscript{50}. The Brillouin zone was sampled using a Γ-centered $3 \times 3 \times 3$ k-point mesh and the plane-wave energy cutoff was set to 243 eV.

DATA AVAILABILITY

The data related to the analysis of third-order models in Si and fourth-order models in $\text{Ba}_2\text{Ga}_5\text{Ge}_{12}$ is publicly available in the form of a GitHub repository at https://gitlab.com/materials-modeling/hiphive-examples.

CODE AVAILABILITY

Code related to the analysis of third-order models in Si and fourth-order models in $\text{Ba}_2\text{Ga}_5\text{Ge}_{12}$ is publicly available in the form of a GitHub repository at https://gitlab.com/materials-modeling/hiphive-examples.

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COMPETING INTERESTS
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