Kernel Charge Equilibration: Efficient and Accurate Prediction of Molecular Dipole Moments with a Machine-Learning Enhanced Electron Density Model

Carsten G. Staacke,¹ Simon Wengert,¹ Christian Kunkel,¹ Gábor Csányi,² Karsten Reuter,¹ and Johannes T. Margraf¹, a)

¹) Fritz-Haber-Institut der Max-Planck-Gesellschaft, Faradayweg 4-6, D-14195 Berlin, Germany
²) Engineering Laboratory, University of Cambridge, Cambridge CB2 1PZ, United Kingdom

State-of-the-art machine learning (ML) interatomic potentials use local representations of atomic environments to ensure linear scaling and size-extensivity. This implies a neglect of long-range interactions, most prominently related to electrostatics. To overcome this limitation, we herein present a ML framework for predicting charge distributions and their interactions termed kernel Charge Equilibration (kQEq). This model is based on classical charge equilibration models like QEq, expanded with an environment dependent electronegativity. In contrast to previously reported neural network models with a similar concept, kQEq takes advantage of the linearity of both QEq and Kernel Ridge Regression to obtain a closed-form linear algebra expression for training the models. Furthermore, we avoid the ambiguity of charge partitioning schemes by using dipole moments as reference data. As a first application, we show that kQEq can be used to generate accurate and highly data-efficient models for molecular dipole moments.

a) Electronic mail: margraf@fhi.mpg.de
I. INTRODUCTION

Kernel and neural network (NN) based machine-learning (ML) methods have in recent years become established as an essential addition to the toolbox of computational chemistry.\textsuperscript{1–4} In particular, ML-based interatomic potentials have had great success in providing energies and forces with quantum mechanical accuracy at a fraction of the cost of first-principles calculations.\textsuperscript{1,5–10} To achieve size-extensivity and a linear computational scaling with system size, these ML potentials typically rely on a local representation of atomic environments and consequently assume that the energy can be decomposed into local atomic contributions.\textsuperscript{9,11} This simple idea has led to a strong focus of chemical ML research on developing sophisticated representations of local atomic environments and, relatedly, NN architectures that directly embed atoms in their neighborhood.\textsuperscript{11–18}

At the same time, it is clear that the assumption of locality does not hold for all systems to the same extent.\textsuperscript{19} Indeed, strongly polar or ionic systems display very long-ranged Coulomb interactions. Even for a fairly unpolar (\textit{e.g.} organic) system, the locality of the energy does not necessarily imply that other electronic properties are similarly local. In particular, electronic properties such as molecular orbital energies or dipole moments can break locality assumptions.\textsuperscript{20} Consequently, such properties tend to be more challenging to predict with purely data-driven ML methods.\textsuperscript{16,20} Beyond this methodological challenge, dipole moments are an intrinsically interesting target as they govern the asymptotic decay of interactions of neutral molecules and their absorption cross-sections in vibrational spectroscopy.

A promising route to overcome the limitations of local ML models is to include known physical interactions explicitly.\textsuperscript{19,21,22} For example, a description of long-range electrostatics can be obtained by learning atom-centered charge distributions (\textit{e.g.} atomic charges, dipoles or partitioned electron densities).\textsuperscript{19,23–25} A prominent recent example of this is the MuML dipole model of Veit \textit{et al.}, which uses atomic charges and atom-centered dipoles to predict molecular dipole moments.\textsuperscript{20} This idea takes advantage of the fact that the charge distributions around atoms can be predicted with reasonable accuracy from local environments, even if their interactions are long-ranged. While this solves some of the issues of local interatomic potentials, there are also significant downsides: Firstly, charge conservation of the overall system is generally not ensured, and secondly non-local charge transfer (\textit{e.g.} through conjugated \(\pi\)-systems) is not captured.\textsuperscript{26}
These issues can be fundamentally addressed by switching the target of the ML model: Instead of predicting the charge distribution directly, one can predict a charge dependent energy expression. The charge distribution is then obtained by minimizing this energy expression under the constraint that the charge is conserved. This idea is closely related to classical charge equilibration approaches like QEq. In this manner, charge conservation is rigorously ensured, the description of non-local charge transfer is enabled and a simple route to analytical derivatives is provided through a Hellmann-Feynman-like approach. The advantage of this approach, compared to directly predicting the charge density, can perhaps be understood in analogy to the choice of initial guess in density functional theory (DFT) calculations: While it is common practice to construct the initial guess from a superposition of atomic densities, it has been found that the superposition of atomic potentials yields a significantly improved starting point.

So far only few examples of ML-based charge equilibration models have been reported, however. Most notably, Goedecker and co-workers applied a NN-based QEq model to ionic crystals. The corresponding models were trained to predict the energies and forces of reference DFT calculations, using the predicted partial charges merely as an intermediate quantity. More recently, Behler, Goedecker and co-workers combined this approach with local NN potentials for the description of organic molecules and MgO surfaces. Here, the charge equilibration models were trained on partial charges from reference DFT calculations. Finally, Xie, Persson and Small applied a more flexible charge-dependent NN to describe lithium hydride nanoparticles, using training data from constrained DFT calculations.

Herein, we present a new kernel-based approach to charge equilibration termed kQEq. These models are directly trained on molecular dipole moments and thus avoid the ambiguity associated with choosing population analysis or projection approaches required in other methods. A closed-form linear algebra expression for training kQEq models is derived and their accuracy is benchmarked on the prediction of molecular dipole moments. Finally, limitations and possible extensions are discussed.
II. THEORY

A. Charge Equilibration

Different conventional (i.e. non-ML) charge equilibration and electronegativity equalization methods have been proposed in the literature.\textsuperscript{27,33–40} In the derivation of the charge equilibration approach we largely follow the formalism of Goedecker and coworkers,\textsuperscript{26,29–31} which is in turn based on the QEq method of Rappé and Goddard.\textsuperscript{27,41} In this context, QEq can be understood as a kind of semi-empirical, orbital-free density functional theory, where the electron density $\rho(\mathbf{r})$ is expanded as:

$$\rho(\mathbf{r}) = \rho_0(\mathbf{r}) + \delta \rho(\mathbf{r}),$$

(1)

where, $\rho_0(\mathbf{r})$ is a reference density (here the superposition of isolated atom densities) and $\delta \rho(\mathbf{r})$ is a fluctuation term, which describes charge transfer and polarization in the interacting system. We expand $\delta \rho$ into a linear combination of normalized $1s$ Gaussians centered at the atomic positions $\mathbf{r}_A$ and of width $\alpha_A$

$$\delta \rho(\mathbf{r}) \approx \sum_{A=1}^{N} -q_A \frac{1}{\pi^{3/2} \alpha_A^3} \exp \left( \frac{|\mathbf{r} - \mathbf{r}_A|^2}{\alpha_A^2} \right),$$

(2)

where $N$ is the number of atoms and $q_A$ are the expansion coefficients. Note that we use the negative of the expansion coefficients $q_A$ here, so that these can directly be interpreted as atomic partial charges. With this approximation, the electron density is completely defined via the charges $q_A$ and it remains to find their optimal values.

To this end, a simple form of the charge-dependent electrostatic energy is assumed:

$$E[\rho] = E_0 + \sum_{A=1}^{N} \left( \chi_A q_A + \frac{1}{2} \eta_A q_A^2 \right) + \frac{1}{2} \int \int \frac{\delta \rho(\mathbf{r}) \delta \rho(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r} d\mathbf{r}'.$$  

(3)

Here, $E_0$ is a charge-independent reference energy, which we set to zero throughout. The second term (labeled 'Site-Energy') is the well-known second-order Taylor expansion of the atomic energy with respect to the charge, with the atomic electronegativity $\chi_A$ and the hardness $\eta_A$.\textsuperscript{42} It provides the energetic contribution incurred by adding or removing electron density from a given atom. The third term (labeled 'Coulomb-Integral') is the classical
Coulomb potential of the fluctuation density $\delta \rho$. This integral can be computed analytically, using the definition of $\delta \rho$ (see Eq. 2):

$$
\int \int \frac{\delta \rho(r) \delta \rho(r')}{|r - r'|} dr dr' = \sum_{A=1}^{N} \left( \frac{2\gamma_{AA}^2}{\sqrt{\pi}} q_A^2 + \sum_{B=1}^{N} q_A q_B \frac{\text{erf}(\gamma_{AB} r_{AB})}{r_{AB}} \right),
$$

(4)

with $\gamma_{AB} = \sqrt{(\alpha_A^2 + \alpha_B^2)^{-1}}$.

This allows rewriting Eq. 3 to:

$$
E[\rho] = E_0 + \sum_{A=1}^{N} \left[ \chi_A q_A + \frac{1}{2} \left( \eta_A + \frac{2\gamma_{AA}}{\sqrt{\pi}} \right) q_A^2 \right] + \frac{1}{2} \sum_{A,B} q_A q_B \frac{\text{erf}(\gamma_{AB} r_{AB})}{r_{AB}},
$$

(5)

which makes it explicit that $E[\rho]$ only depends on the charges $q_A$. We may therefore equivalently use the notation $E(q_1, \ldots, q_N)$. Note that this equation has the familiar form of the original QEq formulation, with the slight difference that the hardness parameter in QEq implicitly includes the electrostatic idempotential $\frac{2\gamma_{AA}}{\sqrt{\pi}}$, whereas here $\eta_A$ only describes the non-classical (e.g. exchange-correlation) contributions to the hardness.

Given the definitions of $\rho$ and $E[\rho]$, we now search for the density that minimizes the energy functional under the constraint that the total number of electrons is conserved. From the definition of $\delta \rho$, it can be seen that this is equivalent to the constraint that $\sum_A q_A = Q_{\text{tot}}$, with the total system charge $Q_{\text{tot}}$. This can be achieved by defining the Lagrangian

$$
\mathcal{L}(q_1, \ldots, q_N, Q_{\text{tot}}) = E(q_1, \ldots, q_N) + \lambda \left( \sum_{A=1}^{N} q_A - Q_{\text{tot}} \right).
$$

(6)

The constrained minimization of the charges can then be performed by setting up a linear system of equations so that:

$$
\frac{\partial \mathcal{L}(q_1, \ldots, q_N, \lambda)}{\partial q_A} = \sum_{B=1}^{N} H_{A,B} q_B + \lambda + \chi_A = 0
$$

(7)

$$
\frac{\partial \mathcal{L}(q_1, \ldots, q_N, \lambda)}{\partial \lambda} = \sum_{A=1}^{N} q_A - Q_{\text{tot}} = 0
$$

(8)

with the elements of the hardness matrix $H$ defined as:

$$
H_{A,B} = \begin{cases} 
\eta_A + \frac{2\gamma_{AA}}{\sqrt{\pi}} & \text{for } A = B, \\
\frac{\text{erf}(\gamma_{AB} r_{AB})}{r_{AB}} & \text{otherwise}.
\end{cases}
$$

(9)
In matrix notation, this linear system can be formulated as:

\[
\begin{pmatrix}
H_{1,1} & H_{1,2} & \cdots & H_{1,N} & 1 \\
H_{2,1} & H_{2,2} & \cdots & H_{2,N} & 1 \\
\vdots & \vdots & \ddots & \vdots & \vdots \\
H_{N,1} & H_{N,2} & \cdots & H_{N,N} & 1 \\
1 & 1 & \cdots & 1 & 0
\end{pmatrix}
\begin{pmatrix}
q_1 \\
q_2 \\
\vdots \\
q_N \\
\lambda
\end{pmatrix} =
\begin{pmatrix}
\chi_1 \\
\chi_2 \\
\vdots \\
\chi_N \\
-Q_{\text{tot}}
\end{pmatrix}
\tag{10}
\]

with the hardness matrix $\bar{H}$, the charge vector $\bar{q}$ and the electronegativity vector $\bar{c}$. Here, the bar-notation is used to indicate that these arrays are extended by one dimension due to the Lagrange multiplier. The corresponding $N$-dimensional sub-arrays are indicated by $H$, $q$ and $c$ in the following.

**B. Kernel Charge Equilibration**

The QEq approach as outlined above only requires three parameters, namely the electronegativity ($\chi_A$), the non-classical contribution to the hardness ($\eta_A$) and the atomic size ($\alpha_A$) for each species in the system. As a flipside of this elegant simplicity, the accuracy and transferability of the QEq method is limited, however. In the kernel Charge Equilibration (kQEq) method proposed herein, we follow the basic idea of Goedecker and coworkers to overcome this limitation.\(^{29}\) This is achieved by allowing the electronegativity of an atom to change as a function of its chemical environment. Importantly, taking advantage of the fact that both QEq and Kernel Ridge Regression (KRR) are formulated as linear problems, we obtain a closed-form expression for training these models.

The environment-dependent electronegativity in kQEq is defined via a kernel regression Ansatz as:

\[
\chi_A(p_A) = \sum_{B=1}^{N_{\text{train}}} w_B k(p_A, p_B),
\tag{11}
\]

where $k$ is a kernel function, $p_A$ is a representation vector that encodes the chemical environment of atom $A$, $w_B$ is a regression coefficient and $N_{\text{train}}$ is the number of atoms in the training set. We use the SOAP kernel and representation vector, which are widely used in the construction of interatomic potentials and as descriptors of local environments.\(^{18}\) We refer to the original literature and a recent review for a detailed account of the corresponding
theory and implementation.\textsuperscript{1,11,43} In general, the kernel function measures the similarity of chemical environments and is defined as:

\[ k(p, p') = (p \cdot p')^2. \]  

(12)

To derive an expression for the regression coefficients \( w \), we begin by noting that the prediction of charges with QEq can be expressed as a matrix multiplication of the matrix \( A \) with the vector of electronegativities \( c \):

\[ q = -Ac \]  

(13)

where \( A \) is the \( N \)-dimensional submatrix of \( \overline{H}^{-1} \). Using Eq. 11, \( c \) can also be written in terms of a matrix-vector multiplication:

\[ c = Kw \]  

(14)

so that

\[ q = -AKw. \]  

(15)

Here, \( K \) is the kernel matrix (quantifying the similarity between the atoms in the system of interest and the atoms in the training set) and \( w \) is the vector of regression coefficients. There are in principle several options for defining the 'optimal' regression coefficients. One could, e.g., fit them to partial charges obtained from some partitioning of the DFT density. However, the choice of a partial charge model is necessarily somewhat arbitrary and does not guarantee an accurate description of electrostatic interactions. We therefore instead use molecular dipole moments \( \mathbf{\mu} \) as a reference, which are unambiguously defined physical observables for finite systems and define the leading order term of molecular interactions in the long-range limit.

In a first-principles calculation, the dipole moment is calculated as:

\[ \mathbf{\mu} = \int r n(r) dr, \]  

(16)

where \( n(r) \) is the total charge density (obtained from the sum of the electron and nuclear charge distributions). For a charge equilibration model like kQEq, this simplifies to

\[ \mathbf{\mu} = \sum_{A=1}^{N} r_A q_A = \mathbf{Rq}. \]  

(17)
using the $3 \times N$ matrix $\mathbf{R}$ with columns $\mathbf{r}_A$. Note that for an unambiguous definition of $\mathbf{\mu}$ for charged systems, center-of-mass shifted coordinates are used throughout.

Combining Eqs. 17 and 15, we obtain:

$$
\mathbf{\mu} = -\mathbf{R}(\mathbf{A}\mathbf{Kw}),
$$

(18)

for the kQEq dipole moment. To determine the regression weights $w$, we then set up a regularized least-squares problem with the loss-function:

$$
L = ||(\mathbf{\mu} - \mathbf{\mu}_{\text{ref}})||^2 + \sigma \mathbf{w}^T \mathbf{Kw} = || -\mathbf{R}(\mathbf{A}\mathbf{Kw}) - \mathbf{\mu}_{\text{ref}}||^2 + \sigma \mathbf{w}^T \mathbf{Kw},
$$

(19)

where $\sigma$ is a regularization hyperparameter and the term $\mathbf{w}^T \mathbf{Kw}$ comes from the use of Tikhonov regularization in a kernel regression model. Note that we use the simplest form of regularization, with a single parameter $\sigma$ to control for overfitting. In principle different regularization strengths could be used for each training point (e.g. proportional to the dipole magnitude).

In a final step, we set $\frac{dL}{dw} = 0$ to minimize $L$ and solve for $w$, to obtain:

$$
\mathbf{w} = -(\mathbf{A}^T \mathbf{R}^T \mathbf{R} \mathbf{A} \mathbf{K} + \sigma \mathbf{1})^{-1} \mathbf{A}^T \mathbf{R}^T \mathbf{\mu}_{\text{ref}}
$$

(20)

The above equations are formulated for a single kQEq problem (i.e. a single molecule or system). In practice we train on multiple systems at once. This can still be achieved with a single linear algebra equation by using blocked matrices for $\mathbf{A}$ and $\mathbf{R}$, and by concatenating the dipole vector elements of all training systems into a single vector.

C. Hyperparameters

Up to now, an environment-dependent description of the atomic electronegativity $\chi_A$ is defined, which can be learned from data. It remains to specify the non-classical contribution to the atomic hardness $\eta_A$ and the atomic radius $\alpha_A$ for each element. Herein, we choose these by very simple heuristics: $\alpha_A$ is set to be proportional to the original QEeq radius of the element in question. These radii are tabulated for all elements up to Lawrencium ($Z=103$). Empirically, we found that scaling these radii with single global scaling parameter $s_{\text{at}} = 0.75$ yields satisfactory results. Similarly, the non-classical hardness parameter $\eta_A$ is set to zero throughout, as we found that this yields robust models while keeping the empiricism of the
method as low as possible. These choices are quite simplistic and further optimization would certainly be possible. As shown below, already these simple defaults provide highly accurate results for the investigated systems though.

The main hyperparameters to be considered for SOAP are the cutoff radius $r_{\text{cut}}$ within which the neighborhood is expanded and the broadness of the Gaussians used to smear out the atomic positions ($\sigma_{\text{atom}}$). The choice of these lengthscales governs the range in which the environment of an atom affects its electronegativity and how sensitive it is to changes of the atomic positions. In the following, we keep the ratio between these parameters constant ($\sigma_{\text{atom}} = \frac{r_{\text{cut}}}{8}$) so that for larger cutoffs, the atomic positions are smeared out more strongly. The idea of keeping this ratio fixed is based on the fact that the expressiveness of a given atom-centered basis set is limited by the number of basis functions, meaning that it can either provide a high-resolution picture that is short-ranged or a lower-resolution picture that is longer-ranged. Alternatively, one could increase the number of basis functions for larger cutoff radii instead, but this would lead to a significantly increased computational cost. The particular constant of proportionality we use was found to work well empirically. Note that the Dscribe implementation of SOAP is used.\(^{43}\) Full details are provided in the supplementary information. The influence of the cutoff parameter is discussed below.

D. Error Metrics

To quantify the performance of the kQEq models for predicting molecular dipole moments, we use two complementary metrics. On one hand, we use the mean absolute error (MAE) of predicted absolute dipole moments. This is a common measure of accuracy, which allows direct comparison with previous models. Additionally, we use the root mean squared regularized relative error (RRMSE) as used by Hait and Head-Gordon in ref. [44]. This metric is defined as $\frac{|\mu - \mu_{\text{ref}}|}{\max (|\mu_{\text{ref}}|, 1 \text{ D})} \times 100\%$, with an arbitrary threshold of 1 Debye that discriminates between small and large dipole moments. In this way, a seamless transition from absolute error (for small dipoles) to relative error (for large dipoles) is achieved, which is necessary since the pure relative error is otherwise severely distorted towards systems with small dipoles.
III. RESULTS

A. Molecular Dipole Moments

As a first benchmark we trained kQEq models for predicting dipole moments of organic molecules. As reference data, the dipole moments of 7500 random molecules from the QM9 database were calculated at the PBE0/def2-TZVP level, using ORCA (data provided in the SI).\textsuperscript{45–47} This set spans a wide range of small to medium sized molecules containing the elements C, H, N, O and F. From these structures, we randomly selected a validation set (used to optimize the regularisation parameter $\sigma$) and a test set of 1000 molecules each. The training sets used below were drawn from the remaining 4000 molecules. Figure 1 depicts the learning curves of different kQEq models, using a range of SOAP cutoffs. For comparison, we also fitted a conventional QEeq model to the same data.

All kQEq models clearly outperform the conventional QEeq approach, underscoring the benefit of the additional flexibility obtained by using environment dependent electronegativities. Furthermore, it can be seen that the kQEq models improve continuously when given more data, whereas the MAE of the conventional approach quickly saturates. The model with the smallest SOAP cutoff used here (1.7 Å) shows the best performance for small training sets but stops improving when training on larger sets. Meanwhile, the larger cutoffs we tested (2.6 Å and 3.5 Å) continuously improve and reach an excellent accuracy of 0.15 D (compared to an intrinsic standard deviation of ca. 2 D).

Following the experience with interatomic potentials based on SOAP, we further tested a 1.7/3.5 Å ”double SOAP” representation.\textsuperscript{7,48} This combines a short-ranged/high-resolution kernel with a longer-range/low-resolution one. The corresponding model reaches an even better accuracy and displays the most robust performance across different training set sizes. The latter is particularly evident when considering the RRMSE, which shows that this model consistently improves the relative error when increasing the training set, whereas some of the other models improve the performance on total dipole moments at the expense of the relative error (\textit{i.e.} by describing small dipole moments less accurately).

Overall, these results show that the physical description of long-range contributions in kQEq allows the use of rather small cutoffs for the ML part, effectively focusing on the nearest neighbors. This is beneficial both in terms of transferability of the models and the
FIG. 1. Learning curves for QEq and kQEq models trained on PBE0 dipole moments for the QM9 dataset. Top: Mean absolute errors (MAE) of four different kQEq models compared to QEq. Three different randomized training sets are used and averaged for each model. Bottom: Regularized relative root mean square error (RRMSE) of the same models.

cost of computing the representation. Notably, the philosophically similar BpopNN model of Xie, Persson and Smalls uses a much larger cutoff radius of 13.2 Å.\textsuperscript{32} Meanwhile, the MuML dipole model of Veit \textit{et al.} (which is also based on SOAP) uses a cutoff of 5 Å.\textsuperscript{20} This indicates that kQEq does a good job of partitioning contributions into long-ranged physical terms and a short-ranged ML model.

To put this performance into perspective, we compare these models to two recent kernel ML models that are specifically tailored to predicting dipole moments, namely the operator ML approach of Christensen, Faber and Lilienfeld and the aforementioned MuML model of
The former uses a modified variant of the Faber-Christensen-Huang-Lilienfeld (FCHL) representation (FCHL⋆), which can incorporate the response of the ML model to applied electric fields and thus provides a physically rigorous and equivariant route to predicting dipole moments. Meanwhile, the latter uses a decomposition of the total molecular dipole into atomic monopole and dipole contributions, using the equivariant λ-SOAP approach.50 For reference we also include the learning curve of a naive FCHL model, which simply predicts the total dipole moment as a scalar (taken from ref. [49]).

As already discussed in reference [49], the FCHL⋆ model is a significant improvement over the scalar approach. It also significantly outperforms conventional QEq for all but the smallest training sets. Meanwhile the MuML and kQEq models display remarkably similar learning curves and represent a further improvement over FCHL⋆. Note that for consistency the QEq and kQEq models in Fig. 1 were trained on the same reference data (calculated at the B3LYP level) as the MuML model from ref. [20].

It is also instructive to consider the accuracy of the reference methods themselves. As mentioned above, Hait and Head-Gordon used the RRMSE to benchmark density functional methods against high-level Coupled Cluster references.44 According to this metric, the best
kQEq model reaches an accuracy of 8.1% on the dipole moments of QM9. This can be compared with the reported errors of popular hybrid functionals like PBE0 (5.2%) and B3LYP (7%). However, it should be noted that the benchmark in ref. [44] focuses on very small molecules and includes spin-polarized systems, so that this comparison should not be overinterpreted. Nonetheless, it indicates that kQEq models approach hybrid DFT accuracy at a much reduced cost.

B. Charge Analysis

Having established the high accuracy of the kQEq predicted dipole moments, we next turn to the predicted charges themselves. It is well known that the electron density cannot be unambiguously partitioned into atomic partial charges. Consequently, there is no way to objectively measure the quality of such partitionings. Indeed, there is a fundamental tension between describing the local charge density around each atom versus global properties such as dipole moments or electrostatic potentials, when approximating a continuous charge density with atom-centered spherically symmetric charges. Nonetheless, it is worth considering whether the predicted charges are reasonably intuitive and how they compare to standard population analysis schemes like the one of Mulliken or restricted electrostatic potential fits like the ChelpG scheme. Here, a double SOAP model trained on 2000 training configurations (at the PBE0 level) is used to provide the kQEq charges (see Fig. 1).

Figure 3 depicts the correlation of Mulliken and kQEq charges for all atoms in the test set. While there is a moderate overall correlation between the charges, there are also clear differences. In particular, we see two general trends. First, kQEq charges appear in a broader range (-1 to 2.5) while Mulliken charges lie between -0.5 and 0.5. Second, kQEq charges display a much more pronounced clustering into functional groups. This is particularly evident for oxygen. While the Mulliken analysis assigns similar charges to ether and carbonyl oxygen atoms, kQEq predicts the carbonyl groups to be significantly more polar. Similarly, the polarity of nitrile and fluoride functional groups is much higher in kQEq.

A similar comparison is shown for ChelpG charges in Fig. 4. Compared to the Mulliken charges, a somewhat better correlation with kQEq is observed and the charges fall into the same range. In particular, the large differences for polar functional groups (e.g. nitrile or carbonyl functional groups) are not observed here. Nonetheless, large deviations can be seen
in other cases, particularly for Fluoride functional groups.

Two illustrative examples of these differences are shown for 1-(2-Methylcyclopropyl)ethanone and 2-Fluoropyrazine in Fig. 5 (both of which are not part of the kQEq training set). At first glance, the partitionings are qualitatively similar, meaning that the signs of most charges match. However, kQEq and ChelpG predict significantly larger absolute charges than Mulliken. In the case of 1-(2-Methylcyclopropyl)-ethanone, this is particularly evident for the carbonyl group, which is much more strongly polarized. Importantly, these differences have strong consequences for the overall description of the molecular electrostatics. Indeed, the dipole moment calculated from the Mulliken charges points in the opposite direction of the actual dipole. In contrast, the ChelpG dipole is well aligned with the reference, but too small in magnitude. Finally, the kQEq charges provide an accurate prediction of both the absolute dipole moment and its direction.
FIG. 4. Correlation between ChelpG and kQEq partial charges for the full test set (green) and individual elements: Oxygen (red), Carbon (grey), Hydrogen (turquoise), Nitrogen (blue) and Fluorine (purple).

The case of 2-Fluoropyrazine is particularly interesting because it illustrates the observed discrepancy between the charges assigned to C-F groups. Here, kQEq predicts the largest absolute charges on the corresponding C and F atoms. Interestingly, the Mulliken and ChelpG charges nonetheless overestimate the molecular dipole moment, while the kQEq prediction provides an excellent fit. This is because the large positive charge on the carbon atom stems not only from charge transfer to Fluorine, but also to the adjacent Carbon and Nitrogen atoms, which partially counter-balance the polarity of the C-F group.

The generally poor correlation between the charges obtained with different schemes raises some questions about the different roles and interpretations partial charges can have. On one hand, they can reflect a local partitioning of the electron density, as in the case of Mulliken, Hirshfeld or Bader charges. On the other hand, they can reflect the electrostatic potential on the surface of a molecules, as in the case of ChelpG and related schemes. While the latter
is arguably less arbitrary (as it is directly tied to a physical observable) it has well-known issues with assigning meaningful charges to atoms that are "buried" in bulky molecules.

The kQEq model proposed herein does not neatly fit into these categories. Firstly, it is not a charge partitioning scheme but an ML model. Mulliken and ChelpG charges can only be obtained by running a full DFT calculation, whereas the kQEq prediction is much cheaper. Secondly, while kQEq models are trained to reproduce molecular dipole moments, the charges themselves are obtained by minimizing the electrostatic energy expression in Eq. 5. Consequently, the Coulomb interactions between partial charges provide an important physical constraint on how the charge distribution is approximated.

C. Extrapolating Beyond Small Molecules

A well-known drawback of the conventional QEeq approach is that it suffers from an delocalization error akin to that observed for local density functionals. This is e.g. evident
in the fact that QEq models incorrectly dissociate molecules into partially charged atoms, since electronegativity differences between isolated atoms lead to spurious charge transfer.\textsuperscript{36} While kQEq could in principle cure this particular pathology (by assigning the same electronegativity to all isolated atoms), the more general delocalization tendencies of QEq will likely be inherited by kQEq to an extent. In this section we explore this issue by testing the performance of kQEq for the organic polymer chains dataset introduced by Veit \textit{et al.}\textsuperscript{20} (see Fig. 6).

This dataset consists of two types of structures. On one hand, glycine polypeptides in the $\alpha$-helix and $\beta$-strand configurations are considered. On the other hand, polynoic amino acids and $n$-amino carboxylic acids are included, which consist of a carboxylic acid and an amine group separated by a conjugated double bond or alkane spacer, respectively. Each of these systems shows characteristic changes of the total dipole moment as the polymer length increases. For the polypeptides, each additional amide bond is itself polar, so that the total dipole increases approximately linearly with the system size. However, the precise behaviour depends on the spatial orientation of these bond dipoles and their interactions, so that the $\alpha$-helix and $\beta$-strand configurations show different scalings. The polynoic amino acid chains also display a linearly increasing dipole moment. In this case this is not due to the addition of polar bonds, however, but due to the polarization of the delocalized $\pi$-electrons in the spacer. Finally, the dipole of the $n$-amino carboxylic acids remains constant upon increasing the chain length, since no polar bonds or delocalized electrons are present.

We again use a ”double SOAP” kQEq model trained on 2000 randomly drawn QM9 molecules for comparison. This model has a mixed performance for this test. For polynoic and $n$-amino carboxylic acids, the performance is very good. This is both in terms of the qualitative features (linear increase in dipole moments, vs. quick saturation) and the quantitative agreement. Furthermore, kQEq is a strong improvement over conventional QEq here. In contrast, the performance for the polypeptides is less satisfactory. While the linear trend is correctly captured, the magnitude of the dipoles is significantly underestimated, in particular for the $\beta$-strand. This behaviour is analogous to the underestimation of dipoles typically observed with local functionals, due to the delocalization error.\textsuperscript{44} While this shows that kQEq does not automatically cure all pathologies of QEq, a clear advantage of a ML approach is that it can be improved with more data. Indeed, by including the longest chains of each type explicitly in the training set, a retrained kQEq model can be generated that
FIG. 6. Dipole moment predictions for α-helix and β-strand polyglycine, polyenoic amino acids, and \(n\)-amino carboxylic acids.\(^{20}\) The kQEq and QEq models are trained on 2000 molecules from QM9. The retrained kQEq model additionally includes the largest polymer in each series, highlighted by a square symbol.

captures these trends more accurately, see Fig. 6. Potentially, an improved extrapolation behaviour could be obtained by using a more flexible expression for the site-energy in Eq. 3, effectively tackling the problem at its root.

Figure 6 also shows the best MuML model from Ref. [20] for comparison. This model performs quite well overall, in particular for the polypeptide systems, with slightly worse performance for the carboxylic acids. In this context, it is worth discussing the fundamental differences between MuML and kQEq. The former uses a fully local decomposition of the overall dipole moment. This works very well for situations where the charge distribution of a large system is essentially just a sum of smaller fragments (as for the polypeptides). In contrast, long-range charge transfer and polarization effects cannot be described by such a model, which is reflected in the underestimated dipole moments of longer polyenoic amino acids.
acid chains. QEq and kQEq are in principle able to describe such non-local effects. The QEq curve for n-amino carboxylic acid reveals that this is not necessarily an advantage however, as large unphysical charge transfer is predicted in this case. In other words, a purely local model will generally lead to more systematic and less dramatic failures than a poor non-local one. Fortunately, kQEq and related methods now provide a framework for sophisticated non-local charge equilibration models.

More generally, it should be noted that kQEq is not primarily intended as a stand-alone molecular dipole model. Since it is based on an energy functional, it can be used to model long-range electrostatic interactions in combination with local interatomic potentials. Indeed, it remains an open question whether dipole moments alone provide enough information for this purpose. Fortunately, the current approach can easily be expanded to also include higher moments, electrostatic potentials or reference partial charges. This will be explored in future work.

IV. CONCLUSION

In this work, we introduced kQEq, a kernel-based approach to charge equilibration in molecules. In contrast to conventional charge equilibration methods like QEq, a data-driven, environment-dependent description of atomic electronegativities is introduced. kQEq models trained on molecular dipole moments display excellent performance, en par with or better than state-of-the-art kernel models, specifically tuned to predicting dipole moments.\textsuperscript{20,49}

The formalism presented herein opens the door towards physics-based kernel ML models for predicting atomic charges, to be used in combination with reactive interatomic potentials. Importantly, the presented approach is quite general and can be extended to other fit targets (\textit{e.g.} quadrupole moments and electrostatic potentials) and to more flexible density representations (\textit{e.g.} using atom centered dipoles in addition to partial charges).

While this work focuses on molecular systems, the application to inorganic materials is also envisioned. For finite systems (\textit{e.g.} nanoparticles) this is in principle straightforward. The use of other fit targets may become important for larger systems, however, as the dipole moment alone might contain too little information in this case. For periodic systems multipole moments are in general ill-defined. Here, the current approach will have to be extended to reproduce other electronic properties.
It should also be noted that QEq-based frameworks are likely not equally well suited for different kinds of materials. In principle, one would expect the best performance for metallic systems, where charges are strongly delocalized and mobile. In contrast, polar insulators or interfaces may be less well described due to the intrinsic delocalization error of charge equilibration models. More general site-energy expressions could be developed to overcome these tendencies. Ideally, such developments will ultimately converge with recent developments in ML-based DFT, yielding a new generation of orbital-free density functionals.\textsuperscript{53–56} Work in this direction is ongoing in our groups.

DATA AND CODE AVAILABILITY

All data and a reference implementation of the kQEq method is available at https://gitlab.com/jmargraf/kqeq.

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