Prediction of Electronic Properties of Radical-Containing Polymers at Coarse-Grained Resolutions

Riccardo Alessandri and Juan J. de Pablo*

Pritzker School of Molecular Engineering, University of Chicago, Chicago, Illinois 60637, United States

E-mail: depablo@uchicago.edu

Abstract

The properties of soft electronic materials depend on the coupling of electronic and conformational degrees of freedom over a wide range of spatiotemporal scales. Description of such properties requires multiscale approaches capable of, at the same time, accessing electronic properties and sampling the conformational space of soft materials. This could in principle be realized by connecting the coarse-grained (CG) methodologies required for adequate conformational sampling to conformationally-averaged electronic property distributions via backmapping to atomistic-resolution level models and repeated quantum-chemical calculations. Computational demands of such approaches, however, have hindered their application in high-throughput computer-aided soft materials discovery. Here, we present a method that, combining machine learning and CG techniques, can replace traditional backmapping-based approaches without sacrificing accuracy. We illustrate the method for an emerging class of soft electronic materials, namely non-conjugated, radical-containing polymers, promising materials for all-organic energy storage. Supervised machine learning models are trained to learn the dependence of electronic properties on polymer conformation at CG resolutions. We then parametrize CG models that retain electronic structure information, simulate CG condensed phases, and predict the electronic properties of such phases solely...
from the CG degrees of freedom. We validate our method by comparing it against a full backmapping-based approach, and find good agreement between both methods. This work demonstrates the potential of the proposed method to accelerate multiscale workflows, and provides a framework for the development of CG models that retain electronic structure information.

INTRODUCTION

Radical-containing polymers, also known as open-shell macromolecules, macromolecular radicals, or simply radical polymers, possess intriguing redox, optoelectronic, and magnetic characteristics that make them appealing for applications ranging from energy storage and optoelectronics to spintronics and memory storage.\textsuperscript{1–6} Non-conjugated, radical-containing polymers are organic polymers that have a non-conjugated backbone bearing pendant stable radical sites. They constitute a class of charge-carrying polymers that do not rely on $\pi$-conjugation to transport charges successfully. The rational design of radical polymers with enhanced characteristics could be greatly advanced by deriving relationships that connect their molecular structure, morphology, and electronic properties. These relationships are inherently multiscale, involving the coupling of electronic and conformational degrees of freedom over a wide range of spatiotemporal scales. New modeling approaches capable of describing such coupling are needed.

Recent work has shown that coarse-grained (CG) models can be used to probe polymeric material length- and timescales reaching the mesoscale.\textsuperscript{7–10} In contrast, explicit quantum-chemical calculations—usually using density functional theory (DFT)—, which are necessary to access electronic properties,\textsuperscript{11–13} are extraordinarily demanding and can only capture picosecond and Angstrom-level processes. To bridge these two scales, that is, to take into account large-scale morphological features generated via self-assembly processes (possibly as a function of processing conditions) when computing electronic properties, recent efforts have sought to introduce new multiscale modeling approaches.\textsuperscript{12,14} In such approaches, the
soft material morphology generated via CG simulations is backmapped to the atomistic resolution required for the quantum-chemical calculations, and such calculations are then performed on conformations drawn from the backmapped morphologies. These multiscale approaches have been primarily developed in the context of organic semiconductors, and provide a means to access the conformational dependence of electronic properties. However, considerable computational demands and workflow complexity of such have limited their applicability for high-throughput computer-aided materials discovery.

Computational studies of radical-containing polymers have been limited. Kemper et al. performed what appear to be the first and one of the few molecular dynamics investigations of radical polymers. They simulated the widely-used p-type radical polymer poly(2,2,6,6-tetramethylpiperidinyloxy-4-yl methacrylate) (PTMA), a polymethacrylate with pendant nitroxide radical TEMPO (2,2,6,6-tetramethylpiperidin-1-oxyl) groups, and provided early insights into some of PTMA’s molecular-scale dynamics and its coupling to electronic properties. More recent efforts that relied on coarser models have sought to predict the conductivity of radical polymers. Such models require as input molecular-level information about the specific redox-active unit in order to be predictive and to differentiate between the performance of different polymers. A different type of effort has focused on molecule-specific characteristics by exploring sets of different redox-active units with DFT-based methods. A limitation of these studies has been the fact that they do not account for condensed-phase conditions, which may affect the resulting electronic properties.

In this work we rely on machine learning (ML) to connect conformational properties to electronic structure properties in condensed phases. The use of ML techniques to study polymers has been limited by a variety of inherent challenges. These include the complexity of data representation in systems that are disordered, the dependence of properties on an ensemble of configurations rather than on a single one, and the dependence of processes on interactions that span multiple length- and timescales. To the best of our knowledge, previous work aimed at predicting the electronic properties of polymers using ML has focused exclu-
sively on polythiophene-based conjugated polymers. Jackson et al., in particular, introduced the concept of electronic coarse-graining (ECG), a ML-based methodology aimed at the prediction of electronic properties based on a molecule’s CG representation, and applied it mostly to prototypical conjugated polymers and single molecules in vacuum. However, no actual CG models—that require not only a CG mapping but also bonded and nonbonded interaction parameters—have been developed so far in the ECG framework; therefore, no validation against traditional backmapping-based approaches has been made. A comparison against backmapping-based approaches has been made by Simine et al., where a similar approach to ECG is used to infer the absorption spectra of a polythiophene system from CG degrees of freedom. However, significant discrepancies between the two approaches were reported and ascribed to the backmapping protocol. Building on the ECG work, in what follows we present an efficient ML-enabled method that can replace state-of-the-art but demanding backmapping-based multiscale approaches. We illustrate the method in the context of nonconjugated, radical-containing polymers, which represent an emerging class of conducting materials. We begin by (1) training supervised ML models to learn the dependence of several electronic properties on molecular conformations at CG resolution, and identifying CG mappings that retain electronic structure information. For this first part, we take inspiration from the ECG methodology. Next, we (2) parametrize CG models that retain electronic structure information using established coarse-graining techniques. Finally, we (3) use such CG models to simulate condensed phases and the trained ML models to predict electronic properties solely from the CG degrees of freedom. We validate our method by comparing its results to those obtained via a standard backmapping-based approach, and show that the proposed method offers a $10^6$ speedup with respect to backmapping-based approaches without loss of accuracy. By exploring different coarse-graining strategies, we are able to delineate general guidelines for development of CG models that retain electronic structure information, and outline a viable pathway to accelerate multiscale workflows aimed at predicting electronic properties while encompassing CG-level spatiotemporal scales.
RESULTS

Method overview. Figure 1 shows a schematic of the proposed method applied to non-conjugated, radical polymers. Only the main aspects of the method are outlined here, and readers are referred to the Materials and Methods section for additional details. To illustrate the method, we use the well-studied, p-type radical polymer PTMA, which bears pendant TEMPO units (Figure 1A). An all-atom (AA) model is used to generate condensed-phase conformations of PTMA using MD simulations. The electronic properties of the resulting configurations are computed via DFT calculations (Figure 1B). Given the nonconjugated backbone of this class of radical-containing polymers, the monomers can be treated as electronically independent (see, e.g., the singly-occupied molecular orbital (SOMO) distribution in Figure 1A). The data generated in this way—all-atom molecular conformations with associated electronic properties—are used as training data to develop supervised ML models. More specifically, we train feed-forward artificial neural networks (NNs) using the conformations (in the form of a reciprocal distance matrix) as input, and the electronic properties as labels. The all-atom molecular conformations can be mapped onto the corresponding CG configurations and NNs can therefore be trained at different CG levels of resolution (Figure 1B). We build two different datasets: one containing single monomer conformations and associated monomer conformation-dependent properties (SOMO energy level, spin density), and a second containing dimer conformations and associated dimer conformation-dependent properties (electronic couplings). After training, we generate two main outputs: (1) trained NNs for the different electronic properties (one for each different resolution) and (2) information on the CG mapping required for the CG model to retain sufficient information about specific electronic properties. The latter information, together with established coarse-graining techniques, are used to develop CG models for simulation of polymer condensed phases over larger spatial and temporal scales; the trained NNs allow one to retrieve electronic properties at the CG level with only negligible computational demands (Figure 1C), thereby permitting the investigation of electronic properties over large ensembles of CG morphologies.
Figure 1: Schematic representation of the proposed method for non-conjugated radical polymers. (A) PTMA radical polymer: its non-conjugated backbone localizes electronic structure features—such as the SOMO orbital represented—to the pendant radical groups. Hydrogen atoms are only shown on the radical group on which the SOMO orbital is localized. (B) Development: starting from an all-atom (AA) model, conformations are sampled via MD simulations. Quantum-chemical calculations are performed on those conformations to generate the training dataset; the dataset serves to train machine learning models—in this case NNs—at different levels of CG resolution. The information gained during training is used to inform the development of CG models that are built on reference AA data via structure-based coarse-graining techniques. (C) Application: the CG models permit study of structures over larger spatiotemporal scales; the trained machine learning models are used to predict electronic properties based on the conformations taken from the CG morphology. The renderings of the morphologies show only polymer backbones for clarity.

Monomer conformation-dependent electronic properties. We first examine the performance of the NN models for prediction of electronic properties that depend on the conformation of a single PTMA monomer, using the SOMO energy level and the spin density as examples of the target properties to be learned. The SOMO energy level determines the position of the transport energy level and hence the charge transport type (p-type vs. n-type), while knowledge of the spin density distribution gives access to the degree of (de)localization
of the unpaired electron(s). The latter property contributes to the stability of the radical center. Both energy levels and spin densities are conformation-dependent; the extent to which they depend on conformation contributes to the degree of energetic disorder in the system.

Figure 2 shows that SOMO energies and spin densities can be predicted at CG resolution. Figure 2 shows correlation plots between the DFT reference SOMO energies and spin densities—computed at the B3LYP/6-311G(d,p) level of DFT (Materials and Methods)—and the same quantities as predicted by the NN model at AA, united-atom (UA), and CG resolutions. The performance of the NN on the AA configurations ($R^2 = 0.978$, $RMSE = 0.024$ eV for SOMO energies; $R^2 = 0.932$, $RMSE = 0.004$ for spin densities) represents the maximum achievable performance for a given molecular representation, given that all other resolutions involve a loss of information for the NN model to perform the regression task on. At UA resolution, the NN also achieves very good predictive performance ($R^2 = 0.928$, $RMSE = 0.044$ eV for SOMO energies; $R^2 = 0.903$, $RMSE = 0.005$ for spin densities), just below that obtained at AA resolution. At CG resolution, there is a noticeable drop of performance with respect to the UA and AA cases, but the NN still provides $R^2 = 0.793$, $RMSE = 0.074$ eV for SOMO energies and $R^2 = 0.744$, $RMSE = 0.009$ for spin densities. To which degree these increased errors would eventually impact the calculation of charge transport rates is difficult to estimate and will require a separate study. The CG mapping used here is “GBNO1” (see Figure 3B); a detailed investigation of the NN model accuracy for different CG mapping choices and resolutions is presented in the next section.

It is useful to investigate the minimum data requirements needed by the NN to learn monomer conformation-dependent electronic properties. Figure S1 shows that after $\approx 4,000$ datapoints, the performance of the NN starts to plateau at all resolutions. This represents a relatively small number of datapoints, and calculations at higher levels of theory could therefore be easily used to train such NNs. We also note that the required NNs are rather small—e.g., 4 hidden layers with 3 neurons each already maximize the performance at CG
resolutions—, and the same performance is achieved for a relatively wide range of hyperparameters (see Figure S8).

Impact of CG resolution and specific CG mapping. The CG level of resolution and the specific CG mapping have considerable influence on the ability of the NN to predict electronic properties. In this section we provide an analysis of resolution that is useful for identifying design principles for CG models that represent the best compromise between computational efficiency and accuracy.

To design CG mappings, we resort to two different strategies. First, we apply the Graph-Based Coarse-Graining (GBCG) algorithm of Webb et al. to arrive at a series of increasingly coarser mappings in a systematic and automated manner with minimal human interven-
tion. Second, we devise a mapping according to the Martini 3 building-block coarse-graining strategy. All the mappings are shown in Figure 3B; the atom-to-bead correspondence is given in detail in Figures S10 and S12. Additional details are provided in the Materials and Methods section and the Supplementary Material.

Figure 3A shows how the predictive performance of the NN model—as quantified by the R² coefficient—changes as a function of the resolution of the molecular representation for the prediction of the SOMO energy level. We see that GBCG mappings lead to poor predictive performance (R² < 0.2), even at the finest CG resolution level, GBCG1. The same is true for the Martini 3 CG mapping (CGM3, the resolution of which is between GBCG1 and GBCG2). Given the high predictive performance of the UA resolution, these results indicate that these “standard” CG mappings are missing key degrees of freedom. Notably, they do not include the nitroxide group explicitly. As can be seen from Figure 3A (black line), the explicit inclusion of the nitroxide group (i.e., the positions of the nitrogen and oxygen atoms are included in the distance matrix used as the NN input vector) improves considerably the performance across the different CG mappings, with R² remaining around 0.8 for GBCG2–GBCG4. Even for resolution GBCG5, which represents the PTMA monomer with only 1 site, when expanded to include the explicit description of the nitroxide group one can achieve a R² above 0.6. This result is expected, and very recently observed also in Ref. 35: the SOMO wavefunction is localized around the nitroxide group (see Figure 1A) and, consequently, the conformation of this group relative to the other atoms in the TEMPO unit governs the SOMO wavefunction coefficients and associated energy.

Further confirmation of the importance of the nitroxide group for prediction of SOMO energies comes from a complementary test, where we exclude either the nitrogen or the oxygen atom of this group from the UA and AA resolutions (Figure S2). Such exclusions lead to a dramatic decrease of predictive performance at the AA and UA levels, with R² values < 0.5 (Figure S2). Figure S2 shows that excluding the nitrogen from the UA and AA representations leads to R² values of ≈ 0.42 while excluding the oxygen from the same
representations degrades $R^2$ down to $\approx 0.1$, indicating that the oxygen position is more informative than that of the nitrogen for SOMO energy predictions. In the case of strongly localized radical groups such as the nitroxide in PTMA, the inclusion of the nitroxide atoms is essential for the NN model to be able to accurately infer the SOMO energy.

Figure 3: Dependence of the predictive performance of NN models on CG resolution. (A) $R^2$ score on the test set as a function of CG resolution for prediction of the SOMO energy level: all-atom (AA), united-atom (UA), and CG mappings obtained with the systematic graph-based coarse-graining approach (GBCGX, with $X=\{1..5\}$; blue line, hollow circles), and CG Martini 3 mapping (CGM3, blue triangle). The black data points (black line, hollow black circles for the GBCG-based mappings, black triangle for the CGM3-based mapping, CGNO) represent mappings where the nitrogen and oxygen atoms of the nitroxide group are described by an explicit CG site each (represented by semitransparent black circles in the renderings of Panel B). (B) Renderings of the AA, UA, and CG mappings and the underlying atomistic structure. The number of sites that each resolution entails is also reported. For the detailed atom-to-bead correspondence, see Figures S10 and S12. The NN’s hyperparameters are optimized for each resolution.

**Dimer conformation-dependent electronic properties.** We now examine the accuracy of NNs for the prediction of electronic properties that depend on the conformation of a dimer, using the electronic couplings as the target property to be learned. Electronic couplings are a key molecular-scale property that determines charge transport, the latter being in turn critical for many applications of radical-containing polymers.$^{1,2}$ Charge transport in these materials is generally believed$^{1,2}$ to occur via a hopping mechanism, where a charge jumps between a radical site and an adjacent ionized site—in the case of PTMA, and of p-type transport in general, a cation site. We approximate the electronic coupling
as the orbital overlap between the SOMO of the neutral radical and the lowest unoccupied molecular orbital (LUMO) of the cation, $\langle \phi_{SOMO} | \phi_{LUMO} \rangle$ (Materials and Methods), and we train NNs to learn its base 10 logarithm, $\log_{10}(\langle \phi_{SOMO} | \phi_{LUMO} \rangle)$.

As shown in Figure 4, the logarithm of the electronic coupling can be learned with good and comparable accuracy at AA, UA, GBCG1-GBCG2, and CGM3 resolutions. The predictive performances (on the test set)—as quantified by the $R^2$ coefficient—are around 0.8-0.81 at AA, UA, GBCG1, and CGM3 resolutions. The performance gap between AA and UA resolutions and the best CG resolutions are small, particularly when compared to the monomer conformation-dependent properties depicted in Figure 2. Electronic couplings, however, are significantly more complex—i.e., data-intensive and requiring larger NNs—and difficult to learn than energy levels or spin densities (compare Figure 4A to Figure S1), across all resolutions. In particular, if a training dataset of $\approx 4,000$ data points was sufficient to train good-performing NNs for the prediction of SOMO energies (Figure S1), at least $\approx 100,000$ data points are necessary for the analogous task on electronic couplings (Figure 4A). After $\approx 100,000$ data points, the predictive performance starts to plateau across the different CG resolutions. Moreover, NNs with 400 neurons per layer are required (see Table S1). This complexity—both in the dataset size required, and NN architecture—is consistent with previous reports. Overall, while more demanding, our results confirm that the task of predicting electronic couplings at CG resolutions can be accomplished. This is particularly important, given their role in determining charge transport.

Another aspect that differentiates electronic couplings from the monomer-dependent electronic properties is the degree of chemical detail required by the CG mapping resolution. In particular, resolving the nitrooxide group is not critical in the case of electronic couplings: adding an explicit description of the nitrooxide group does not improve the results within the statistical uncertainty of our predictions (compare GBCG1-2 (solid) to GBNO1-2 (dashed) and CGM3 (solid) to CGNO (dashed) in Figure 4A). This observation can be understood in terms of the electronic coupling being dominated by the relative position between the two
monomers. The nitroxide groups do not contribute to the definition of this relative position, and hence their explicit representation does not improve predictions of electronic couplings significantly.

![Graph showing predictive performance of NN models for dimer conformation-dependent electronic properties at all-atom and coarse-grained resolutions.](image)

**Figure 4:** Predictive performance of NN models for dimer conformation-dependent electronic properties at all-atom and coarse-grained resolutions. (A) NN performance as a function of training dataset size for different resolutions: all-atom (AA), united-atom (UA), GBCG1-2, GBN01-2, and CG Martini (CGM3 and CGNO). See Figure 3B for a schematic representation of the mappings. (B) Two-dimensional histograms showing the best accuracy achieved by the NNs for the different resolutions. The $R^2$ scores and RMSE are indicated on each plot and represent the results on the test set. The NN’s hyperparameters are optimized for each resolution and dataset size.

**Electronic properties of condensed-phase CG simulations.** We now develop CG models that retain electronic structure information and apply the trained NNs to predict electronic properties from condensed-phase CG simulations. We validate the NN predictions by comparing them against the electronic properties obtained via the reference, state-of-the-art approach that requires backmapping and explicit quantum-chemical calculations.

To investigate what the most suitable coarse-graining strategies are when developing CG models that retain electronic structure information, we first build two CG models with the same mapping and bonded interactions but different nonbonded interactions. To realize this,
we choose the CGM3 mapping that, as we have seen, allows for good accuracy in predicting electronic couplings. At the same time, the CGM3 mapping allows us to test two sets of nonbonded interactions derived via two different CG philosophies, namely the building-block CG Martini 3 approach and the structure-based CG IBI method. For details on the CG models, see Materials and Methods and the Supplementary Material.

We first consider the results generated with the CGM3 IBI model (Figure 5A-C). The model is used to produce a morphology via CG MD, and we then use two methods to retrieve the electronic couplings of such a CG morphology: the reference approach that involves backmapping to atomistic resolution, and explicit DFT calculations (taken to be the ground truth; in red in Figure 5C) and the proposed ML-based method (in blue in Figure 5C). The agreement between the two electronic coupling distributions is excellent, with both the NN-predicted mean (μ) and standard deviation (σ) being around 1% of the corresponding reference values. We stress that the trained NN is able to predict electronic couplings of a CG simulation with an accuracy comparable to that of the reference approach but at a fraction of the computational cost (see next section for a discussion on the computational efficiency). Moreover, the NN was trained on CG conformations mapped from an AA MD simulation and had not seen conformations drawn from a CG simulation before. However, the structural accuracy of the CG model (Figure 5B) is such that the CG model spans a conformational space that is consistent with the underlying AA structure, and therefore the NN is able to make predictions on conformations drawn from previously unseen CG morphology realizations with excellent accuracy.

In contrast to the CGM3 IBI model, the CGM3 Martini 3 model leads to unsatisfactory electronic property predictions (Figure S4). In fact, not only does the electronic coupling distribution obtained with the Martini model exhibit a considerably larger standard deviation (+15%, Figure S4C) than the reference distribution, but that distribution also shows qualitative discrepancies with respect to the reference distribution (i.e., it is not a unimodal distribution). These discrepancies can be rationalized by looking at the structure of the Martini
CG morphology, for example by analyzing TEMPO-TEMPO and nitroxide-nitroxide radial distribution functions (RDFs) (Figure S4B). We see that the RDFs are qualitatively different, showing, for example, an extra peak at \( \approx 0.4 \) nm in the case of the nitroxide-nitroxide RDF. These discrepancies are indicative of molecular conformations occurring in the Martini CG morphology that have no AA counterpart, and which are hence non-physical. In contrast, structural accuracy is the parametrization target of structure-based coarse-graining techniques such as IBI and, therefore, as shown in the previous paragraph, the resulting CG models represent the underlying AA structure with high fidelity (Figure 5B). This structural accuracy appears to be an essential requirement when developing CG models that retain electronic structure information.

Having settled on IBI as the strategy of choice for developing CG models that retain electronic structure information, we now turn to the development of a CG model that is able to retain information on not only electronic couplings but also energy levels. Informed by the results of the previous sections, we know that an explicit description of the nitroxide group is required for energy level prediction and we therefore develop a corresponding CG model. We choose the GBNO2 mapping (Figure 5D), as it represents a good compromise between NN predictive accuracy—showing accurate electronic coupling (Figure 3) and energy level (Figure 4) predictions—and coarse-graining degree. We derive nonbonded interactions via IBI for all the GBCG2 beads (Supplementary Material). The nitrogen and oxygen atoms of the nitroxide group are instead described by two virtual dummy sites that do not interact via nonbonded interactions with any of the other CG sites. As such, the GBNO2 model is expected to allow for predictions of both electronic couplings and energy levels.

As done for CGM3-based CG models, we generate a morphology via CG MD using the GBNO2 model just described and predict this time both energy levels and electronic couplings using the trained NNs (Figure 5D-F). Again, we compare the NN predictions to the ground truth taken to be the energy levels and electronic couplings obtained with the reference, backmapping-based approach (Figure 5F-5G). The agreement between the two
Figure 5: Application and validation of the proposed ML-based method to PTMA, a prototypical radical polymer. (A) Schematic representation of the CG model (CGM3 mapping) and the underlying AA structure. (B) Representative AA vs. CG RDFs for the CGM3 model with nonbonded interactions obtained via IBI. (C) Electronic couplings computed with the reference backmapping-based approach (red) and the proposed ML-based method (blue) starting from a CG morphology generated via CG MD using the CGM3 model. (D) Schematic representation of the CG model (GBNO2 mapping) and the underlying AA structure; the nitroxide atoms are described by two virtual dummy sites (transparent gray circles; see text for details). (E) Representative AA vs. CG RDFs for the GBNO2 model. (F) Electronic couplings and (G) SOMO energies computed with the reference backmapping-based approach (red) and the proposed ML-based method (blue) starting from a CG morphology generated via CG MD using the GBNO2 model. Mean ($\mu$) and standard deviation ($\sigma$) of the electronic property distributions are reported in the legends.
electronic coupling distributions is also excellent in this case, with the NN-predicted $\mu$ being within 1% of the reference one and $\sigma$ matching the reference value. In contrast, the agreement between the two SOMO energy distributions is only satisfactory, with a very good agreement between the NN-predicted and reference $\mu$ (< 1%) but a NN-predicted $\sigma$ that is 22% lower than the reference. We note that in this case, we are looking at an electronic property that depends strongly on the intra-monomer degrees of freedom. The degeneracy of the CG representation, i.e., the fact that multiple AA conformations correspond to the same CG conformation, is likely responsible\textsuperscript{34} for the narrowing of the SOMO distribution predicted at CG resolution. As discussed later, a recently proposed\textsuperscript{34} Deep Kernel Learning approach may remedy this shortcoming and improve the prediction. Albeit with this limitation to keep in mind for the SOMO energy predictions, the GBNO2 model retains information on both energy levels and couplings, and can be used to explore large ensembles of morphologies at CG resolution and retrieve electronic structure information with negligible computational costs.

**Comparison to quantum chemistry approaches.** To highlight the importance of considering condensed-phase effects in soft materials when predicting electronic properties, including electronic couplings, we compare our results to some of the Boltzmann-averaged electronic couplings that have been used in recent, state-of-the-art quantum-chemical modeling studies\textsuperscript{13,24} aimed at understanding transport in radical polymers. In such studies, the electronic coupling for a given molecular species is evaluated by first generating gas-phase dimer structures either randomly\textsuperscript{24} or with more elaborate dimer surface sampling algorithms.\textsuperscript{13} For each of the generated dimer structures, the electronic coupling is computed with the level of theory of choice, along with either a total or binding energy for that structure. The energies, which are indicative of the relative stability of each structure in the gas phase, are then used to obtain a Boltzmann-averaged electronic coupling according to:

$$\langle V \rangle = \frac{\sum_{i=1}^{N} V_i \exp\left(-\frac{E_i}{k_B T}\right)}{\sum_{i=1}^{N} \exp\left(-\frac{E_i}{k_B T}\right)}$$

(1)
where $V_i$ is the electronic coupling of the $i$-th dimer, $E_i$ its binding energy (calculated as the single point energy difference between the dimer ($E_{i\text{dimer}}$) and the isolated species ($E_{i\text{radical}}$, $E_{i\text{cation}}$), according to: $E_i = E_{i\text{dimer}} - E_{i\text{radical}} - E_{i\text{cation}}$, $k_B$ the Boltzmann constant, $T$ the temperature (300 K), and $N$ the total number of dimers.

For a comparison to the method introduced here, we must compute the Boltzmann-averaged electronic coupling at the same level of theory used in the present work. Hence, we randomly generate 10,000 dimer structures following the procedure by Li and Tabor (Supplementary Material for details) and compute the $\langle \phi_{\text{SOMO}} | \phi_{\text{LUMO}} \rangle$ orbital overlap for each structure. For the same dimer structure, we also compute the binding energy (at the $\omega$B97X-D/6-311G(d,p) level of DFT) and calculate the Boltzmann-averaged electronic coupling according to Eq. 1. The resulting coupling is shown with a vertical dashed line in Figure 5C and Figure 5F. We can see that the Boltzmann-averaged coupling overestimates the magnitude of the mean electronic coupling that arises in condensed phases. The discrepancy between the two is not surprising, given that the gas-phase sampling of the dimer structures neglects any solid-state packing preference dictated by (1) steric constraints (including the fact that the monomers are attached to a polymer backbone) and (2) (un)favorable intra- and inter-chain interactions. A second intrinsic limitation of computing Boltzmann-averaged electronic couplings is the lack of an estimate of the so-called structural disorder present in the system, i.e., the width of the electronic coupling distribution. This disorder may affect the charge transport landscape and may therefore impact the conductivity of the material. In contrast, the proposed method, whose accuracy depends on the structural accuracy of the CG model, gives access to not only (1) electronic couplings that account for the packing in the condensed phase, but also (2) the full distributions of electronic couplings, thereby allowing for quantification of the structural disorder in the system.
Having demonstrated the feasibility of designing and developing CG models that retain electronic structure information, and using such models as a more efficient substitute for current multiscale approaches that involve backmapping procedures, we conclude by briefly outlining a few opportunities for future extensions of the presented method.

The computational efficiency is a key advantage of the proposed method with respect to backmapping-based approaches, and is expected to enable innovative high-throughput investigations of electronic properties over ensembles of soft material morphologies (see below). The upfront cost of the dataset generation and ML model training is (considering both electronic couplings and SOMO energies): \( \approx 375 \cdot 10^3 \) CPU hours for training data generation and \( \approx 4.2 \cdot 10^3 \) CPU hours for training. Note that this estimation takes into account both the exploration of suitable mappings and the hyperparameter grid search performed for all the NNs. The training data generation dominates the upfront cost. Computing electronic properties with the backmapping-based approach requires a total of \( \approx 27 \cdot 10^3 \) CPU hours for a single snapshot (\( \approx 2.5 \cdot 10^3 \) hours for backmapping and relaxation, and \( \approx 24.5 \cdot 10^3 \) hours for the quantum-chemical calculations). Instead, the NN predictions take \( \approx 1.5 \cdot 10^{-1} \) CPU hours per snapshot. See Table S4 for a detailed breakdown. Therefore, if electronic properties are predicted for 13 CG frames, we break even with the upfront training cost (see Figure S25). For any further frame, the proposed method will provide a speedup of \( 10^6 \) times. Further efficiency gains may be realized if more generic ML models, i.e., models trained on a larger fraction of the (CG) chemical space, will be developed—see also discussion below.

The fact that neither the GBCG nor the Martini CG mappings describe the nitroxide group explicitly makes these mappings unable to predict the SOMO energy level. Although it is unsurprising that such fine detail is being averaged out by the CG mappings, this result highlights the limitation of current methods to devise CG mappings that retain specific electronic structure information (e.g., SOMO energy). While less strongly localized radical units might be less affected by this—i.e., we speculate that if a radical is more delocal-
ized, a coarser representation may still be able to capture the conformational-dependence of the SOMO energy—a systematic way to identify mappings that preserve specific electronic properties would be greatly beneficial for automating the CG mapping step of the proposed method.

We have shown that a structurally-accurate CG model is essential for the proposed method. Hence, in general, structure-based coarse-graining techniques—such as IBI—should be used when deriving interactions for CG models that retain electronic structure information. However, other kinds of CG models such as chemically-specific, building-block CG force fields (e.g., Martini) that do not use the structure as a target for parametrization, may still be valuable. First, while structural inaccuracies cannot be excluded for Martini-like CG approaches, they are also not necessarily present in every CG model developed within such frameworks. In fact, the latest Martini force field update (version 3) has taken structural accuracy more into account via bonded parameter optimization that targets the AA molecular volume and shape and a wider range of systematically pre-calibrated nonbonded interactions; for example, structural accuracy is expected to improve for conjugated structures.

Hence, more generally, regardless of how the CG model potentials are obtained, as long as structural accuracy is preserved—which can be inspected, e.g., by comparing AA and CG RDFs—we expect a CG model to be suitable for the proposed method. Second, given the computational efficiency of Lennard-Jones based models such as Martini as compared to models that use (IBI-derived) custom nonbonded potentials (Table S5), and the transferability of CG models developed within building-block CG frameworks, it is still valuable to investigate how such CG strategies can contribute to the development of CG models that retain electronic structure information.

With respect to the ML model and featurization schemes used here, there is ample room for improvement. Here we used NNs due to their simplicity of implementation, and application flexibility. However, recently Gaussian process regression has been combined with NNs to realize a Deep Kernel Learning (DKL) approach to ECG. The approach al-
allows to incorporate the distributional nature of the electronic property predictions resulting from the CG mapping degeneracy, which, as we have seen in the present work, may be needed for electronic properties that strongly depends on detailed intramolecular features (Figure 5G). DKL moreover provides prediction uncertainties and it is hence suitable for active learning strategies. Regarding the input featurization, the (reciprocal) distance matrices used here constitute one of the simplest molecular representations that guarantees rotational and translation invariance. However, a host of more elaborated featurization schemes, such as representations based on symmetry functions or graph convolutions, are available. In particular, input representations containing chemical identity and possibly other physical descriptors are expected to play a key role in order to explore the question of whether ML models able to predict electronic properties at CG resolution and that are transferable across CG chemical space can be trained. Finally, applications to conjugated polymers, in contrast to the nonconjugated polymers studied here, need to consider that electronic states can delocalize along the polymer backbone. To tackle this problem, long short-term memory networks combined with a ∆-ML approach have been shown to be a promising strategy to achieve the molecular-weight transferability necessary to consider conjugation along polymer backbones.

The proposed method opens up interesting avenues for applications that require efficient modeling of radical-containing polymers, and more generally, in soft electronic materials. For example, electronic properties could be studied as a function of morphology processing conditions in a high-throughput fashion. In particular, charge transport networks and fluctuations of electronic couplings over CG spatiotemporal scales can be studied by the proposed approach. Electronic couplings thus gathered could be fed into, for example, kinetic Monte Carlo simulations aimed at computing conductivities. As discussed before, such high-throughput studies are made possible by the $10^6$ increase in efficiency of the proposed method with respect to current backmapping-based approaches.

In conclusion, an efficient ML-enabled method has been presented that connects elec-
tronic properties of soft materials with the CG spatiotemporal scales required to sample their phase space. The method is able to replace current backmapping-based multiscale approaches and directly connect electronic structure information to CG degrees of freedom. The method has been demonstrated for nonconjugated, radical-containing polymers but it is suitable for any soft material with electronic properties. Overall, the findings reported underscore the potential of the proposed strategy to propel the investigation of soft materials’ electronic properties that depend on conformational degrees of freedom over a wide range of spatiotemporal scales, a critical step towards the bottom-up design of soft electronic materials.

MATERIALS AND METHODS

AA models. Initial parameters were obtained for a OPLS-AA/CM1A force field from the LigParGen server. QUBEKit was subsequently used to derive parameters for bond and angle potentials from DFT calculations (B3LYP/6-311++G(d,p)) that are tailored to PTMA. Charges were obtained with Gaussian via the CHELPG method. The resulting force field led to a density for methyl methacrylate and TEMPO—molecular fragments that make up PTMA and for which experimental reference densities are available—in good agreement (within 2.5%) with experiments (Table S2).

Datasets. Two datasets were built: one with single PTMA monomer conformations and associated monomer conformation-dependent electronic properties (namely, SOMO energy level and spin density) and a second one with dimer conformations and associated dimer conformation-dependent electronic properties (electronic couplings). We refer to these as “monomer dataset” and “dimer dataset”, respectively. The following protocol was used to ensure good conformational sampling for the training data generation. We ran a condensed-phase AA MD simulation at 680 K (i.e., \( \approx 1.3 \cdot T_g \); see Figure S9 for the estimation of the glass transition temperature, \( T_g \), of the AA model) and, after 30 ns of equilibration, we gathered snapshots every 10 ns. Each snapshot was relaxed at 300K for at least 20 ns. The
relaxed snapshots were then used to extract the conformations for which quantum-chemical calculations were performed. Two different procedures were used to extract conformations for the two datasets, both leveraging the MDAnalysis python library\textsuperscript{47,48} and Open Babel.\textsuperscript{49} For the monomer dataset, 4 snapshots containing 3,000 conformations each were used. For each conformation, coordinates were extracted up to the first carbon of the backbone ("C01" of Figure S10) and the carbon atom capped with 3 hydrogen atoms. For each such conformation, a reciprocal distance matrices at the different resolutions were computed and stored and an input file for a single point calculation (B3LYP/6-311++G(d,p)) was written. The $(4 \cdot 3,000)$ 12,000 conformations were filtered to increase the diversity of the dataset by removing all conformations that had a RMSD of less than 0.8 Å with any other conformation. Gaussian\textsuperscript{46} was used to obtain the SOMO energy and Mulliken spin density (B3LYP/6-311++G(d,p)). A final dataset of 10,778 datapoints was obtained. For the dimer dataset, 24 snapshots containing about 9,200 dimer conformations each were used. A dimer was selected if the distance between the respective nitrogen atoms was within a cutoff of 10 Å. For each selected dimer, the DFT-optimized geometry of the monomer was aligned (by minimizing the RMSD\textsuperscript{50,51}) to each MD monomer structure. In this way, intramonomer vibrations were neglected. Intermonomer vibrations are much faster than the intermonomer motions and affect negligibly intermonomer couplings.\textsuperscript{52} The electronic couplings were approximated as the orbital overlap\textsuperscript{52} between the SOMO of the neutral radical and the LUMO of the cation, $\langle \phi_{\text{SOMO}} | \phi_{\text{LUMO}} \rangle$. For each dimer, Gaussian\textsuperscript{46} was used to compute (1) the orbitals of monomer $i$ and (2) the orbitals of monomer $j$ of the of $i$-$j$ dimer both by performing a single point calculation using as starting point the converged orbitals of the DFT-optimized geometry; (3) the overlap matrix by performing a calculation that produces only the overlap matrix in the MO basis set of the dimer; finally, Multiwfn\textsuperscript{53} was used to evaluate $\langle \phi_{\text{SOMO}} | \phi_{\text{LUMO}} \rangle$ by providing the Gaussian checkpoint files from the three calculations described above. Note that the average overlap between the $\langle \phi_{\text{SOMO},i} | \phi_{\text{LUMO},j} \rangle$ and $\langle \phi_{\text{LUMO},i} | \phi_{\text{SOMO},j} \rangle$ values was taken, as the two values are in general different (although,
at least for the present system, they are very similar). A final dataset of 221,406 datapoints was obtained.

**Coarse-graining.** GBCG mappings were generated using the spectral grouping variant of the GBCG algorithm,\(^{37}\) as implemented at [https://github.com/xmwebb/GBCG](https://github.com/xmwebb/GBCG). The Martini CG mapping was devised following the Martini 3 guidelines.\(^{10,40}\) Bonded parameters for the CG models were obtained by fitting standard bond, angle, and dihedral potentials to reference distributions obtained from mapped atomistic simulation (Supplementary Material). Nonbonded parameters for the structure-based CG models were derived by iterative Boltzmann inversion (IBI)\(^{36}\) as implemented in the VOTCA package.\(^{54}\) 15 and 10 pair interactions were parametrized for the CGM3 and GBNO2 CG models, respectively. For more details, see the Supplementary Material. Nonbonded parameters for the Martini model—that is, Martini bead types—were assigned based on Martini 3 guidelines\(^{10,40}\) (and validated by computing octanol/water free energies of transfer, see Table S3) and available models for similar polymers\(^{55}\) (Supplementary Material).

**Backmapping.** Backmapping was performed with Backward,\(^{56}\) which relaxes the AA structure with the atomistic force field after an initial geometrical reconstruction of the AA structure based on the CG particle positions. The geometrical reconstruction is specified via CG-to-AA mapping files.

**Molecular dynamics simulations.** Starting configurations for all polymers simulations were set up with Polyply,\(^{55}\) and MD simulations run with Gromacs versions 2021.x or more recent.\(^{57}\) AA simulations used a time step of 1 fs and the Verlet scheme with a nonbonded cutoff of 1.1 nm, dispersion correction, and the particle mesh Ewald (PME) method for long-range electrostatic interactions. Bonds involving hydrogen atoms were constrained. Temperature and pressure were controlled by a Nosé-Hoover thermostat (coupling parameter, \(\tau_T\), of 1.0 ps) and a Parrinello-Rahman barostat (coupling parameter, \(\tau_P\) of 5.0 ps), respectively. The Berendsen barostat was used for equilibration purposes (\(\tau_P = 0.5\) ps). For the Martini CG simulations, the Verlet scheme with a straight nonbonded cutoff of 1.1
Temperature and pressure were controlled by a velocity-rescaling thermostat ($\tau_T = 1.0$ ps) and a Parrinello-Rahman barostat ($\tau_P = 12$ ps), respectively. For the IBI-based CG simulations, the group cutoff scheme was used in order to use tabulated potentials, with a nonbonded cutoff of 1.5 nm. Note that CG simulations with IBI-derived nonbonded potentials were run with Gromacs version 2019.5 because tabulated potentials are not available in more recent Gromacs versions.

**ML model details.** For each monomer or dimer conformation, a reciprocal distance matrix $D$ between all particles of a particular resolution was computed. Its elements are $D_{kl}^{ij} = |r_k^{(i)} - r_l^{(j)}|^{-1}$ where $r$ is the position vector, $i$ and $j$ are the monomer indices, and $k$ and $l$ are the atom indices. For the monomer dataset, $j = i$ and hence $D_{kl}^{ii} = |r_k^{(i)} - r_l^{(i)}|^{-1}$. Each matrix was flattened and the resulting one-dimensional vector (of dimension $N^2$) used as the input feature for the NN. The electronic properties (SOMO energy level, spin density, electronic coupling) are the labels associated to the input vectors for the supervised ML regression task. The SOMO energy and spin density (only the values on the nitrooxide nitrogen and oxygen atoms were considered, given that all other atoms have a spin density < 0.07; hence, two values were passed to the NN) were used as obtained from the DFT calculations. The base 10 logarithm of $\langle \phi_{SOMO} | \phi_{LUMO} \rangle$ was used in the case of the couplings. A fully-connected, feed-forward NN with a $M$-dimensional input layer followed by 4 batch-normalized hidden layers with the same number of neurons was used so as to have a NN flexible enough. $M$ is the dimension of the (flattened) input vector and it therefore depends on the molecular resolution ($M = N^2$). Hyperparameters that were optimized include: the number of neurons in the hidden layers, the batch size, and the number of training epochs. The default learning rate of the NAdam optimizer (0.001) was used for training. Visualizations of the hyperparameter grid search for SOMO energy prediction for some of the resolutions can be found in Figure S5-S8. Standard scaling was applied to the input and output features. 10% of the datasets (1,077 datapoints for the monomer dataset; 22,140 for the dimer dataset) was held out and used as the test set. The remaining datapoints were...
used to build training and validation sets by using 5-fold cross-validation. Hyperparameters were optimized by grid search based on the 5-fold cross-validated performance. For the best performing NN models for each property and molecular resolution, see Table S1. The final model performance was measured by applying the best model chosen based on the 5-fold cross-validation to the held-out test set. When predicting electronic couplings, as done when generating the dimer dataset: (1) couplings were inferred for dimers for which the distance between the nitrogen virtual sites (or the beads representing the nitroxide group in the case of the CGM3 mapping) was within a 10 Å cutoff; (2) the CG-mapped DFT-optimized structure were aligned to each monomer of the CG MD simulation snapshot structure and electronic couplings were inferred based on the reciprocal distance matrix computed for the dimer conformation after alignment. All the ML methods were implemented using the Keras and scikit-learn libraries.

**Data availability.** Data and code to reproduce the findings of this work are available at https://github.com/ricalessandri/ECG-RedoxPoly-Cathode. The developed AA and CG polymer models are also implemented in the Polyply library (https://github.com/marrink-lab/polyply_1.0). The TEMPO and PTMA monomer Martini models are also available on the Martini 3 small molecule library (https://github.com/ricalessandri/Martini3-small-molecules).

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Supporting Information Available

Additional results: NN performance vs. dataset size for SOMO energy prediction, extended Figure 3, extended Figure 4, extended Figure 5 (CGM3 Martini model); ML model hyperparameters; density and $T_g$ data of the AA model; details on the development of the CG models: mappings, bonded parameters, IBI nonbonded parameters, Martini nonbonded parameters; random dimer structure generation; computational performance data.

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Supporting Information:

Prediction of Electronic Properties of Radical-Containing Polymers at Coarse-Grained Resolutions

Riccardo Alessandri and Juan J. de Pablo

Pritzker School of Molecular Engineering, University of Chicago, Chicago, Illinois 60637, United States

E-mail: depablo@uchicago.edu
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## 1 Additional results

Figure S1: **NN performance vs. training dataset size for the prediction of SOMO energies.** NN performance as a function of training dataset size for different resolutions: all-atom (AA), united-atom (UA), GBNO1, GBNO2. Solid lines are results on the test set, while dash-dotted lines are results the fivefold cross-validated results.

Figure S2: **Effect of excluding either N or O from the molecular representation on the predictive performance of NN models for SOMO energy prediction.** $R^2$ coefficients obtained for AA and UA representations without the nitrogen ("only O", orange crosses) or without the oxygen ("only N", orange crosses). For comparison, data reproduced from Figure ??A of the main text are also shown.
Figure S3: **Predictive performance for electronic couplings: GBCG mappings.** (A) NN performance as a function of training dataset size for different resolutions: all-atom (AA), GBCG1-3, and GBNO1-3. See Figure ??B for a schematic representation of the mappings. The $R^2$ scores is for the test set.

Figure S4: **Application of the method using the Martini 3 CG model.** (A) Schematic representation of the CG model (CGM3 mapping) and the underlying AA structure. (B) Representative AA vs. CG RDFs for the CGM3 model with Martini 3 nonbonded interactions. (C) Electronic couplings computed with the reference backmapping-based approach (red) and the proposed ML-based method (blue) starting from a CG morphology generated via CG MD. Mean ($\mu$) and standard deviation ($\sigma$) of the electronic property distributions are reported in the legends.
# Machine learning models

Table S1: **Hyperparameters of the highest-performing NN models.** For the different properties and molecular resolutions, the hyperparameters—optimized via grid search—for the highest-performing NN models (as judged from the fivefold cross-validated $R^2$ score) are reported.

| property, resolution | learning rate | batch size | no. epochs | no. neurons / layer |
|-----------------------|----------------|------------|-------------|---------------------|
| couplings, AA         | 0.002          | 1024       | 3000        | 400                 |
| couplings, UA         | 0.002          | 1024       | 2000        | 400                 |
| couplings, GBCG1      | 0.004          | 1024       | 2000        | 400                 |
| couplings, GBNO1      | 0.001          | 256        | 2000        | 400                 |
| couplings, GBCG2      | 0.004          | 1024       | 2000        | 400                 |
| couplings, GBNO2      | 0.001          | 256        | 2000        | 400                 |
| couplings, GBCG3      | 0.001          | 256        | 2000        | 400                 |
| couplings, GBNO3      | 0.001          | 256        | 2000        | 400                 |
| couplings, GBCG4      | 0.001          | 256        | 2000        | 400                 |
| couplings, GBNO4      | 0.001          | 256        | 2000        | 400                 |
| couplings, GBCG5      | 0.001          | 256        | 2000        | 400                 |
| couplings, GBNO5      | 0.001          | 256        | 2000        | 400                 |
| couplings, CGM3       | 0.001          | 256        | 2000        | 400                 |
| couplings, CGNO       | 0.001          | 256        | 2000        | 400                 |
| SOMOs, AA             | 0.001          | 128        | 1000        | 3                   |
| SOMOs, UA             | 0.001          | 128        | 2000        | 3                   |
| SOMOs, GBNO1          | 0.001          | 256        | 2000        | 3                   |
| SOMOs, GBNO2          | 0.001          | 256        | 1000        | 3                   |
| SOMOs, GBNO3          | 0.001          | 128        | 1000        | 12                  |
| SOMOs, GBNO4          | 0.001          | 128        | 0100        | 12                  |
| SOMOs, GBNO5          | 0.001          | 256        | 2000        | 12                  |
| SOMOs, CGNO           | 0.001          | 256        | 2000        | 3                   |
Figure S5: **Hyperparameter grid search for SOMO energy prediction: AA resolution.** Hyperparameters tested via grid search are reported on the heatmap axes, that are colored according to the corresponding fivefold cross-validated $R^2$ obtained after training.

Figure S6: **Hyperparameter grid search for SOMO energy prediction: UA resolution.** Hyperparameters tested via grid search are reported on the heatmap axes, that are colored according to the corresponding 5-fold cross-validated $R^2$ obtained after training.
Figure S7: Hyperparameter grid search for SOMO energy prediction: GBCG1 resolution. Hyperparameters tested via grid search are reported on the heatmap axes, that are colored according to the corresponding fivefold cross-validated $R^2$ obtained after training.

Figure S8: Hyperparameter grid search for SOMO energy prediction: GBNO1 resolution. Hyperparameters tested via grid search are reported on the heatmap axes, that are colored according to the corresponding fivefold cross-validated $R^2$ obtained after training.
3 All-atom model

Table S2: **Mass densities: AA model vs. experiments.** Mass densities for the fragments constituting PTMA, namely methyl methacrylate (MMA) and TEMPO. Densities obtained via AA simulations are reported along with the reference experimental values. All values in g cm\(^{-3}\). Also reported are the error percentages with respect to the experimental values.

|          | exp.  | AA (err %)     |
|----------|-------|----------------|
| TEMPO (crystal) | 1.077\(^{\text{S1}}\) | 1.071±0.002 (−0.5%) |
| MMA      | 0.94\(^{\text{S2}}\)   | 0.963±0.003 (+2.4%) |

Figure S9: **Glass transition temperature, \(T_g\), estimation.** We estimate \(T_g\) of the AA model by cooling down a morphology from 800 K and equilibrating it at temperatures spaced every 20 K. The simulations were equilibrated at the target temperature for 30 ns and the density was extracted as the average density over the subsequent 20 ns of MD simulation. The data gathered in this way in the range 100-800 K are shown on the left-hand side plot. Two independent linear fits of the density-temperature data have been performed to individuate the glass and melt regions. To choose the best fits, the following procedure was employed: starting from the 4th datapoint from the lowest temperature and from the highest temperature for the glass and melt fits, respectively, we scanned through all combinations of fits by progressively increasing the number of datapoints to be included in the fits; the best fits were selected by choosing the ones that maximize the sum of the \(R^2\) coefficients of the linear fits (right-hand side plot). \(T_g\) was estimated to be at the point where the two fits cross, i.e., 511 K.
4 Coarse-grained models

Mappings. The CGM3 and GBNO2 mappings, for which CG model parameters have been derived, are given in Figure S10. GBCG mappings are given in Figure S11 (schematic representation) and Figure S12 (atom-to-bead correspondence). GBCG mappings that include an explicit nitroxide group description are given in Figure S13 (atom-to-bead correspondence).

Figure S10: Atom labels and mappings for PTMA: CGM3 and GBNO2. (left) Atom labels. (right) The atom-to-bead correspondence is given for CGM3 and GBNO2 mappings, along with the Martini 3 (M3) bead types $^{33}$ and the names for the IBI bead types.

| M3 bead type | IBI bead type | Atoms (CGM3 mapping) |
|--------------|---------------|----------------------|
| SC3 (PMMA)   | VNL           | C01 C17 H38 H39 C02 H18 H19 H20 |
| SN4a (PMMA)  | EST           | C03 O04 O05          |
| SC3 (TEMPO)  | C1            | C06 H21 C07 H22 H23 C11 H24 H25 |
| SC3 (TEMPO)  | C2            | C40 C12 H26 H27 H28 C13 H30 H31 |
| SC3 (TEMPO)  | C2            | C08 C15 H32 H33 H34 C16 H34 H35 H36 |
| TP5 (TEMPO)  | N4            | N09 O14              |

| IBI bead type | Atoms (GBNO2 mapping) |
|---------------|-----------------------|
| VNL           | C03 C01 C02 O04 C17 H18 H19 H20 H38 H39 |
| EST           | C06 O05 H21           |
| C1            | C08 C07 C15 C16 H22 H23 H32 H33 H34 H35 H36 H37 |
| C1            | C10 C11 C12 C13 H24 H25 H26 H27 H28 H29 H30 H31 |
| NO            | N09 O14               |
| virtual site  | N09                   |
| virtual site  | O14                   |

Cii (Cjj) indicates the connection with the previous (next) monomer.

Figure S11: Graph-Based CG mappings for a monomer of PTMA. From (left) all-atom (AA) resolution to (right) a 1-site CG model. Figure S12 details the atom-to-bead mapping. This Figure is a larger version of Figure ??B of the main text.
Figure S12: **Graph-Based CG mappings.** Beads are separated by commas. Atom labels correspond to the ones of Figure S10.

"GBCG1": [C01 C02 C17 H18 H19 H20 H38 H39, C03 O04, C06 O05 H21, C07 H22 H23, C08 C15 C16 H32 H33 H34 H35 H36 H37, N09 O14, C10 C12 C13 H26 H27 H28 H29 H30 H31, C11 H24 H25]

"GBCG2": [C03 C01 C02 004 C17 H18 H19 H20 H38 H39, C06 O05 H21, C08 C07 C15 C16 H22 H23 H32 H33 H34 H35 H36 H37, N09 O14, C10 C11 C12 C13 H24 H25 H26 H27 H28 H29 H30 H31]

"GBCG3": [C06 C01 C02 C03 004 C17 H18 H19 H20 H21 H38 H39, C08 C07 O09 O14 C15 C16 H22 H23 H32 H33 H34 H35 H36 H37, C10 C11 C12 C13 H24 H25 H26 H27 H28 H29 H30 H31]

"GBCG4": [C06 C01 C02 C03 004 C17 H18 H19 H20 H21 H38 H39, C08 C07 O09 O14 C15 C16 H22 H23 H24 H25 H26 H27 H28 H29 H30 H31 H32 H33 H34 H35 H36 H37]

"GBCG5": [C08 C01 C02 C03 004 005 C06 C07 N09 C10 C11 C12 C13 O14 C15 C16 C17 H18 H19 H20 H21 H22 H23 H24 H25 H26 H27 H28 H29 H30 H31 H32 H33 H34 H35 H36 H37]

Figure S13: **Graph-Based CG mappings with explicit nitroxide group description.** Beads are separated by commas. Atom labels correspond to the ones of Figure S10.

"GBNO1": [C01 C02 C17 H18 H19 H20 H38 H39, C03 O04, C06 O05 H21, C07 H22 H23, C08 C15 C16 H32 H33 H34 H35 H36 H37, N09 O14, C10 C12 C13 H26 H27 H28 H29 H30 H31, C11 H24 H25]

"GBNO2": [C03 C01 C02 004 C17 H18 H19 H20 H38 H39, C06 O05 H21, C08 C07 C15 C16 H22 H23 H32 H33 H34 H35 H36 H37, N09 O14, C10 C11 C12 C13 H24 H25 H26 H27 H28 H29 H30 H31]

"GBNO3": [C06 C01 C02 C03 004 005 C17 H18 H19 H20 H21 H38 H39, N09 O14, C08 C07 C15 C16 H22 H23 H32 H33 H34 H35 H36 H37, C10 C11 C12 C13 H24 H25 H26 H27 H28 H29 H30 H31]

"GBNO4": [C06 C01 C02 C03 004 005 C17 H18 H19 H20 H21 H38 H39, N09, O14, C08 C07 C10 C11 C12 C13 C15 C16 H22 H23 H24 H25 H26 H27 H28 H29 H30 H31 H32 H33 H34 H36 H37]

"GBNO5": [N09, O14, C08 C01 C02 C03 004 005 C06 C07 C10 C11 C12 C13 C15 C16 C17 H18 H19 H20 H21 H22 H23 H24 H25 H26 H27 H28 H29 H30 H31 H32 H33 H34 H35 H36 H37 H38 H39]
**Bonded parameters.** Bonded parameters were obtained by extracting bond, angle, and torsion distributions from mapped atomistic simulations and fitting them to standard harmonic bond, angle, and dihedral (i.e., improper dihedral) potentials. CG sites are mapped to the center of geometry (COG) of the group of atoms they represent (see Mappings.). The resulting CG parameters are given in Figure S15 and the distributions are shown in Figure S14. Similarly, CG parameters and distributions are shown in Figure S18 and Figure S19, respectively, for the GBNO2 mapping. The two virtual dummy sites used to describe the nitrogen and oxygen atoms of the nitroxide group are “virtual” sites because their displacement is not calculated by the integrator algorithm, but their position is instead recomputed from the new positions of the constructing CG particles (i.e., a number of CG sites that completely define the virtual sites’ position) after each integration step. They are “dummy” sites because they do not interact via nonbonded interactions with any of the other CG sites. See Figure S18 for their definition.

The bonded parameters for the PMMA-like backbone were taken from the Martini 3 PMMA model\(^4\) (and are shown in Figure S20) in the case of the CGM3 mapping. For the GBNO2 mapping, new parameters were obtained and are shown in Figure S21 (and corresponding distributions in Figure S22).

**Nonbonded parameters: Iterative Boltzmann inversion.** For the IBI procedure, reference atomistic RDFs are needed. Bonded parameters are taken from the previous step, and only nonbonded pair interactions are optimized using IBI as implemented in VOTCA.\(^5\) The same nonbonded exclusions used in the calculations of the reference atomistic RDFs should be used in the IBI iteration steps and hence in the final CG model. Typically, one excludes intramolecular pairs of CG beads which interact via bonded interactions. In the case of PTMA, we exclude all the intramonomer bead interactions and the 1-2 and 1-3 interactions between all the other beads. In GROMACS, the former is achieved by explicitly excluding the intramonomer pairs via a [exclusions] section, and the latter by setting nrexcl = 2, as shown in the topology file in Figure S15 and Figure S18 (the files are Polyply block files.
in .ff format; they use GROMACS-like syntax to define the repeating unit of a polymer needed by Polyply\textsuperscript{S4}).

A comparison between the AA reference RDFs and the RDFs obtained with the final IBI-derived CG potentials for PTMA is shown in Figure S16 (CGM3) and Figure S17 (GBNO2).

Figure S14: Bond, angle, and torsion distributions for a PTMA monomer (CGM3 mapping). Atomistic reference distributions are in blue, CG distributions are in red. Compare header to labels in Figure S15 for information on the particular degree of freedom shown.
Figure S15: **Topology of the PTMA CG model (CGM3 mapping).** Shown is the Polyply block .ff file (that uses GROMACS syntax). The exclusion settings apply to the CG IBI model. In the CG M3 model, the \[exclusions\] are omitted and \texttt{nrexcl = 1} (the Martini default). Units for the bond, angle, and dihedral parameters are indicated in the respective sections.

```plaintext
[moleculetype]
; name nrexcl
PTMA 2

[atoms]
; id type resnr residue atom cgnr charge mass
 1 VNL 1 PTMA VNL 1 0.0 54
 2 EST 1 PTMA EST 2 0.0 54
 3 C1 1 PTMA C1 3 0.0 54
 4 C2 1 PTMA C2 4 0.0 54
 5 C2 1 PTMA C3 5 0.0 54
 6 N4 1 PTMA N4 6 0.0 36

[bonds]
; i j funct length force_k
; harmonic (nm) (kJ mol\(^{-1}\) nm\(^{-2}\))
 1 2 1 0.262 15000 ; b_1
 2 3 1 0.253 15000 ; b_2
 3 4 1 0.326 25000 ; b_3
 3 5 1 0.326 25000 ; b_4

[constraints]
; i j funct length
; (nm)
 3 6 1 0.310 ; b_5
 4 6 1 0.238 ; b_6
 5 6 1 0.238 ; b_7

[angles]
; i j k funct ref.angle force_k
; harmonic (deg) (kJ mol\(^{-1}\) rad\(^{-2}\))
 1 2 3 2 132.9 70 ; ang_1
 2 3 4 2 129.0 60 ; ang_2
 2 3 5 2 121.9 95 ; ang_3

[dihedrals]
; i j k l funct ref.angle force_k
; harmonic (deg) (kJ mol\(^{-1}\) rad\(^{-2}\))
 3 4 5 6 2 140.00 20 ; dih_1
 2 4 5 6 2 150.00 80 ; dih_2

[exclusions]
 1 2 3 4 5 6
 2 3 4 5 6
 3 4 5 6
 4 5 6
 5 6
```
Figure S16: Pair radial distribution functions (RDFs) between CG bead types for the CGM3 mapping. In each panel, results using the CGM3 representation are obtained from mapped all-atom (AA) simulations (black) and CG simulations with pair potentials computed from iterative Boltzmann inversion (IBI) (red).
Figure S17: Pair radial distribution functions (RDFs) between CG bead types for the GBNO2 mapping. In each panel, results using the GBNO2 representation are obtained from mapped all-atom (AA) simulations (black) and CG simulations with pair potentials computed from iterative Boltzmann inversion (IBI) (red).
Figure S18: Topology of the PTMA CG model (GBNO2 mapping). Shown is the Polyply block .ff file (that uses GROMACS syntax). Units for the bond, angle, and dihedral parameters are indicated in the respective sections.
Figure S19: Bond, angle, and torsion distributions for a PTMA monomer (GBNO2 mapping). Atomistic reference distributions are in blue, CG distributions are in red. Compare header to labels in Figure S18 for information on the particular degree of freedom shown. Degrees of freedom $b_7 - b_8$, $ang_3 - ang_8$, and $dih_1$ and $dih_4$ (which have no explicit potential in the topology, see Figure S18) correspond to those that are either kept in place by the virtual sites or by the other potentials.
Figure S20: **Topology of the PMMA-like backbone model (CGM3 mapping).** Shown is the Polyplinker link .ff file (that uses GROMACS syntax). Units for the bond, angle, and dihedral parameters are indicated in the respective sections. From Ref. S4.

```plaintext
[ link ]
resname "PTMA"
[ bonds ]
; i  j  funct  length force_k
; harmonic  (nm) (kJ mol^-1 nm^-2)
VNL +VNL  1  0.315  4000  {"group": "vinyl backbone"}

[ link ]
resname "PTMA"
[ angles ]
; i  j  k  funct ref.angle force_k
; harmonic  (deg) (kJ mol^-1 rad^-2)
VNL +VNL ++VNL  2  115  35  {"group": "vinyl backbone"}

[ link ]
resname "PTMA"
[ angles ]
; i  j  k  funct ref.angle force_k
; harmonic  (deg) (kJ mol^-1 rad^-2)
EST  VNL +VNL  2  70  20
```

Figure S21: **Topology of the PMMA-like backbone model (CGM3 mapping).** Shown is the Polyplinker link .ff file (that uses GROMACS syntax). Units for the bond, angle, and dihedral parameters are indicated in the respective sections.

```plaintext
[ link ]
resname "PTMA"
[ bonds ]
; i  j  funct  length force_k
; harmonic  (nm) (kJ mol^-1 nm^-2)
VNL +VNL  1  0.305  12000  {"group": "vinyl backbone"}; b_1

[ link ]
resname "PTMA"
[ angles ]
; i  j  k  funct ref.angle force_k
; harmonic  (deg) (kJ mol^-1 rad^-2)
VNL +VNL ++VNL  2  126  40  {"group": "vinyl backbone"}; ang_1

[ link ]
resname "PTMA"
[ angles ]
; i  j  k  funct ref.angle force_k
; harmonic  (deg) (kJ mol^-1 rad^-2)
EST  VNL +VNL  2  90  20 ; ang_2
```
Figure S22: **Bond, angle, and torsion distributions for a PMMA-like backbone (GBNO2 mapping).** Atomistic reference distributions are in blue, CG distributions are in red. Compare header to labels in Figure S21 for information on the particular degree of freedom shown.

**Nonbonded interactions: Martini 3.** Bead types for the backbone beads are taken from the Martini 3\(^3\) PMMA model (available in the Polyply library)\(^4\) and are SC3 and SN4a (see also Figure S23). Martini 3 bead types for the 2,2,6,6-Tetramethyl-1-piperidineolate (TEMPO) unit have been chosen following recommendations based on the Martini 3 small molecule dataset,\(^6\) when available. When not, such as in the case of the nitroxide group, the bead type was chosen by computing the free energies of transfer and comparing to experimental data (see Table S3). Accordingly, we found that a TP5 bead gives a $\Delta G$ in fair agreement with the available experimental logP of 1.85.\(^7\)

We moreover analyzed the shape and molecular volume of the Martini CG model for TEMPO with respect to its corresponding AA model by computing solvent accessible surface areas (SASAs) and Connolly surfaces, according to the Martini 3 protocol described elsewhere.\(^3,6\) Results are shown in Figure S24 and show that although the SASA average values are in good agreement (within 2%), the Connolly surface comparison shows mismatches between the AA and CG models. In particular, the lower thickness of the CG Connolly surface is likely responsible for the RDF discrepancies that can be observed in Figure S4: namely, as a consequence of the too low thickness of the Martini 3 model, the TEMPO structures can come closer in space, likely giving rise to the extra peak at low distances in the nitroxide-nitroxide RDF (Figure S4B).
Figure S23: **Martini 3 bead types for the PTMA CG model.** Shown is the `[atoms]` section of the Polyply Polyply block `.ff` file (that uses GROMACS syntax). Martini 3 bead types are shown in the second column.

```plaintext
[ moleculetype ]
; name nrexcl.
PTMA  1

[ atoms ]
; id  type  resnr  residu  atom  cgnr  charge  mass
1   SC3  1       PTMA  VNL  1    0.0   54  ; Martini 3 PMMA model
2   SN4a 1       PTMA  EST  2    0.0   54  ; Martini 3 PMMA model
3   SC3  1       PTMA  C1  3  0.0   54  
4   SC3  1       PTMA  C2  4  0.0   54  
5   SC3  1       PTMA  C3  5  0.0   54  
6   TP5  1       PTMA  N4  6  0.0   36  
```

Table S3: **Transfer free energies for TEMPO.** The free energy relative to the transfer of the solute molecule from solvent S\(_1\) to S\(_2\) (\(\Delta G_{S_1 \rightarrow S_2}\)) obtained from experiments. These are shown along with the experimentally determined partitioning coefficient logP representing the same transfer (logP\(_{S_1 \rightarrow S_2}\)) Solvents are n-octanol (OCO) and water (W). All the free energies are in kJ mol\(^{-1}\). Statistical uncertainty for the computed \(\Delta G\) is below 0.3 kJ mol\(^{-1}\) in all cases. Experimental data from Refs. S7.

|          | \(\Delta G_{OCO \rightarrow W}\) | (logP\(_{OCO \rightarrow W}\)) exp | CG | AA |
|----------|---------------------|-------------------------------|----|----|
| TEMPO    | (SC3)\(_3\)-TP5     | (1.85)                        | 10.6 | 13.6 | —  |

Figure S24: **Solvent accessible surface areas and Connolly surfaces for TEMPO:** **AA vs. CG models.** The CG surface is depicted in blue while the AA surface in gray. Mean and standard deviation of the SASA distributions are as follows: \(\text{SASA}_{\text{AA}} = 4.06 \pm 0.01 \text{ nm}^2\) \(\text{SASA}_{\text{CG}} = 3.98 \pm 0.03 \text{ nm}^2\) (the CG SASA is 1.9% lower than the AA one).
5 Random dimer structure generation for Boltzmann-weighting

To randomly generate the dimer configurations to be Boltzmann-weighted, we use the algorithm by Li and Tabor and available at https://github.com/Tabor-Research-Group/redox_mol_screening. Briefly, the algorithm places two DFT-optimized monomers at an overlapping position at the origin. From this initial configuration, one of the two monomers is randomly translated and rotated. To avoid exploring conformational space where the coupling is zero, the maximum translation allowed was such that the closest heavy atom distance between two monomers was within 1.5 and 3.0 Å. The raw and Boltzmann-weighted overlaps are shown in Figure S25.

Figure S25: Coupling distributions obtained via the gas-phase random dimer structure generation. The grey distribution represents the histogram of the un-weighted overlaps; the red distribution represents the histogram of the Boltzmann-weighted overlaps, that is, the $y$ axis represents the Boltzmann factor $\exp(-\frac{E_i}{k_B T}) / \sum_{i=1}^{N} \exp(-\frac{E_i}{k_B T})$; the dark red is the Guassian fit of the Boltzmann-weighted distribution. The Boltzmann-averaged (Eq. ?? of the main text) is also reported. The disorder, $\sigma$, obtained from the Boltzmann-weighted distribution ($\sigma \approx 0.11$) underestimates greatly the disorder found in the condensed phase simulations ($\sigma \approx 0.85$) shown in the main text.
6 Computational performance

Table S4: Benchmark of methods: backmapping-based approach vs. proposed ML-based method. For the training benchmarks, “datapoints” refers to the number of different training runs that were executed (different hyperparameter combinations, different mappings).

|                             | CPU hours/datapoint | datapoints | total CPU hours |
|-----------------------------|---------------------|------------|-----------------|
| dataset generation (SOMOs)  | ≈0.91               | 10,778     | ≈9,808          |
| dataset generation (couplings) | ≈1.65             | 221,406    | ≈365,320        |
| training (SOMOs)            | ≈1.06               | 1,152      | ≈1,221          |
| training (couplings)        | ≈13.86              | 216        | ≈2,994          |
| prediction single snapshot (SOMOs) | ≈0.000009         | 3,000      | ≈0.03           |
| prediction single snapshot (couplings) | ≈0.000009       | 13,164     | ≈0.12           |
| Total                       |                     |            | ≈379,343        |
| backmapping (incl. relaxation) | N/A                | N/A        | ≈2,600          |
| QC calculations single snapshot (SOMOs) | ≈0.91             | 3,000      | ≈2,730          |
| QC calculations single snapshot (overlaps) | ≈1.65            | 13,164     | ≈21,721         |
| Total                       |                     |            | ≈27,051         |

Figure S26: Computational cost of the reference method vs. the proposed method. The estimated computational cost (in CPU hours) of the backmapping-based reference multiscale approach (blue, “reference”) is plotted along with the estimated computational cost (in CPU hours) of the proposed ML-based method (black, “this work”) as a function of the number of CG snapshots for which the electronic properties are predicted.
Table S5: **Benchmark of atomistic (AA) and coarse-grained (CG) models computational performance.** The performance is reported and compared also between the CG models parametrized using Martini 3 (M3) and IBI. The performance refers to MD simulations carried out on 10 CPUs and with GROMACS version 2019.5.

|                     | Performance (ns/day) | speed-up (AA vs. CG) | speed-up (M3 vs. IBI) |
|---------------------|---------------------|----------------------|-----------------------|
| AA                  | ≈17                 | 1                    | N/A                   |
| CG IBI              | ≈230                | ≈13.5                | 1                     |
| CG M3               | ≈3900               | ≈230                 | ≈17                   |

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