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# Structure-Guided Machine Learning for Efficiency Prediction of Organic Photovoltaics Using Experimentally Informed Molecular Descriptors

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**ABSTRACT:** The efficiency of organic photovoltaics was estimated using a machine learning (ML) approach. We used the organic photovoltaics database built in-house by the Korea Research Institute of Chemical Technology. The dataset comprises reliable and representative experimental results for 1,010 ternary organic solar cells (D1:D2:A), obtained through repeated measurements. The data included 67 donors and 24 non-fullerene acceptors, device structures, donor/acceptor structures, donor-to-acceptor ratios, active-layer thicknesses, experimental conditions, and local symmetry.

We fragmented the donors and acceptors using a self-developed method. A dataset was created by generating descriptors of the fragmented molecules and used to train various ML algorithms, including random forest, XGBoost, LightGBM, support vector regression, and multilayer perceptron. Model performance was evaluated using the coefficient of determination ( $R^2$ ). XGBoost showed the highest  $R^2$  of 0.849. The contributions of key features were interpreted using SHAP analysis. This paper presents an ML framework that combines molecular fragmentation and data-driven modeling.

## 1. INTRODUCTION

Organic photovoltaics (OPVs) have attracted considerable attention as a promising renewable energy technology owing to their low manufacturing cost, mechanical flexibility, and continuous improvements in power conversion efficiency (PCE).

Since the first demonstration of OPV devices in 1986, significant progress has been achieved through advanc



es in active-layer materials and device architectures.<sup>1</sup> In particular, the introduction of donor–acceptor (D–A) bulk heterojunction (BHJ) structures enabled efficient exciton dissociation and charge transport, leading to substantial improvements in PCE.

Early efforts to enhance OPV performance relied heavily on the synthesis of novel donor and acceptor materials, as well as device-level optimization. Parallel to experimental developments, computational chemistry approaches based on density functional theory (DFT)<sup>2,3</sup> and semi-empirical methods (e.g., AM1<sup>4</sup>, PM6<sup>5</sup>, and PM7<sup>6</sup>) were employed to screen candidate materials and understand structure–property relationships. However, such approaches become computationally prohibitive when applied to large molecular libraries, limiting their applicability to high-throughput OPV material discovery.<sup>7–10</sup>

To overcome these limitations, data-driven approaches and machine learning (ML) techniques have increasingly been adopted to predict OPV performance. Early studies employed quantitative structure–property relationship (QSPR)<sup>11,12</sup> models, followed by more advanced ML frameworks using experimentally measured or DFT<sup>13</sup>-derived descriptors. While these approaches demonstrated encouraging predictive capabilities, they often suffered from limited dataset sizes, computational bias, or reliance on idealized molecular representations.

More recently, fragmentation-based molecular representations have been introduced to reduce molecular complexity and improve model scalability. Wu et al.<sup>14</sup> employed literature-derived fragmentation schemes to construct fingerprint-based ML models, whereas Kim et al.<sup>15</sup> utilized synthesis-oriented functional group indexing to achieve improved prediction accuracy. Despite these advances, most existing studies remain focused on relatively simple donor–acceptor systems and are primarily based on literature-reported data<sup>13,16–26</sup>, which are often biased toward high-efficiency devices and do not adequately reflect diverse experimental conditions encountered in practical laboratories.

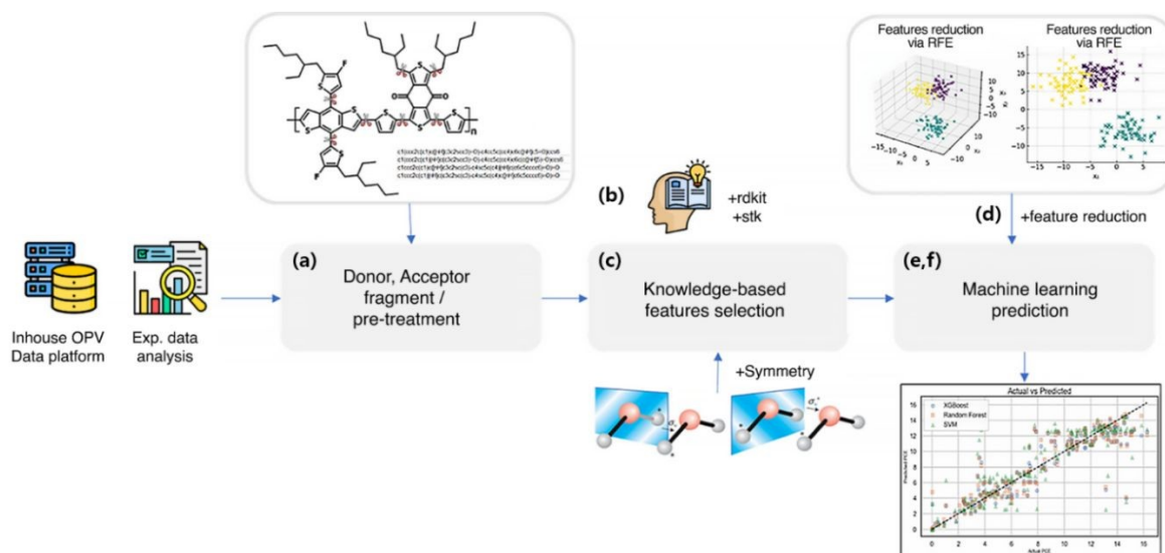
Machine-learning studies on ternary OPV systems have also emerged, with particular emphasis on D:A1:A2<sup>27</sup> configurations composed of small-molecule donors and acceptors. While these studies achieved meaningful success, their molecular representations are less suitable for polymer-rich systems, where repeating units, structural heterogeneity, and multiple donor components introduce additional complexity that is difficult to capture using conventional descriptors.

In this work, we focus on ternary OPV systems of the D1:D2:A type, in which two polymeric donors are blended with a single acceptor. To the best of our knowledge, this study represents the first systematic ML investigation of ternary OPV devices involving multiple polymer donors. By leveraging a curated experimental database that includes both high- and low-efficiency devices, we mitigate literature bias and construct a more balanced representation of real experimental conditions.

To accurately describe such complex systems, we propose a fragment-based molecular representation based on chemically meaningful fragmentation. Fragment-level physicochemical descriptors are generated using RDKit and Stk, and local symmetry information is incorporated as a physically motivated proxy for molecular packing tendencies without explicitly modeling solid-state morphology. Redundant and irrelevant features are further removed through REF-based feature selection, enabling the construction of robust and transferable machine-learning models for ternary OPV systems.



Using this representation, we develop machine-learning models to predict power conversion efficiency and key device parameters. Interpretability analyses provide insights into structure–process–property relationships, highlighting the contributions of specific molecular fragments and device parameters to device performance (**Figure 1**).



**Figure 1.** DL-assisted design framework of OPV materials: (a) polymer fragmentation and SMILES conversion (b) descriptor generation using RDKit and Stk, (c) data pre-treatment, (d) feature operation/processing, (e) machine learning, and (f) performance prediction.

## 2. RESULTS AND DISCUSSION

### 2.1. Collection of experimental OPV data

The experimental data used in this study were collected over the past five years by the Photoenergy Research Center at the Korea Research Institute of Chemical Technology. In 2018, our Digital Chemistry Research team established an in-house OPV data platform to systematically accumulate, analyze, and visualize experimental data without reliance on external online tools (**Figure S1**).

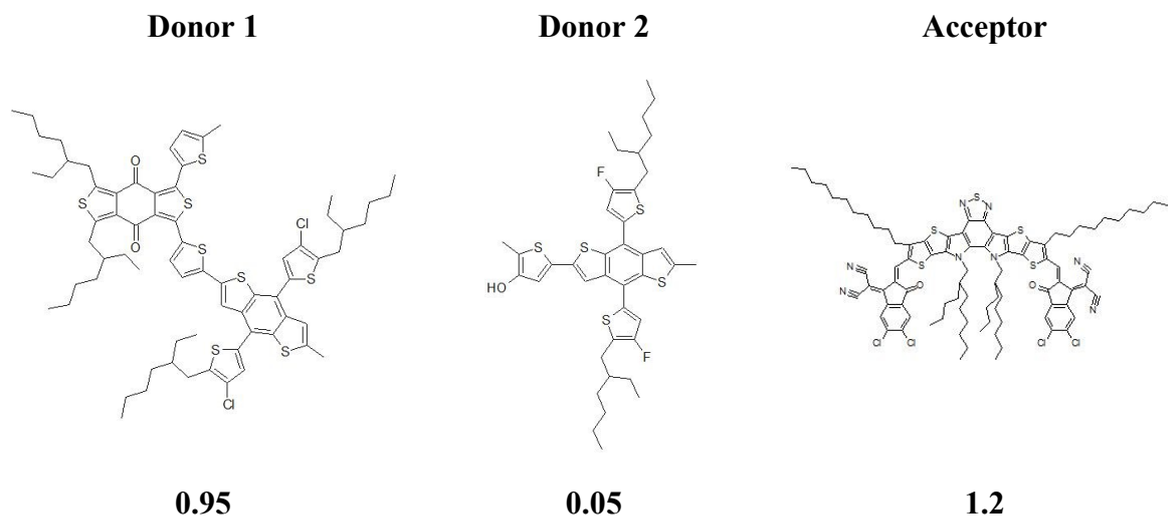
The dataset consists of two main categories: materials and devices. The materials category contains chemical and structural information on donor and acceptor compounds, while the device category records OPV device architectures and electrode-related experimental results. ChemAxon's Marvin JS<sup>28</sup> and JChem Microservices were used to convert compounds into MolV2000 format for structural representation, and CDK 2.8 was used to convert them into SMILES.<sup>29</sup>

In the device section, the platform stores detailed information on all OPV layers, including active-layer materials, additives, solvents, and processing conditions, which can be batch uploaded using standardized templates. Multiple experimental files can be uploaded simultaneously, enabling automated performance calculations and visualization. This automated workflow supports efficient integration and management of large-scale experimental datasets, thereby facilitating the application of artificial intelligence (AI) and machine learning (ML) techniques.

The dataset focuses on ternary organic photovoltaic systems composed of two donors and one acceptor. To acc

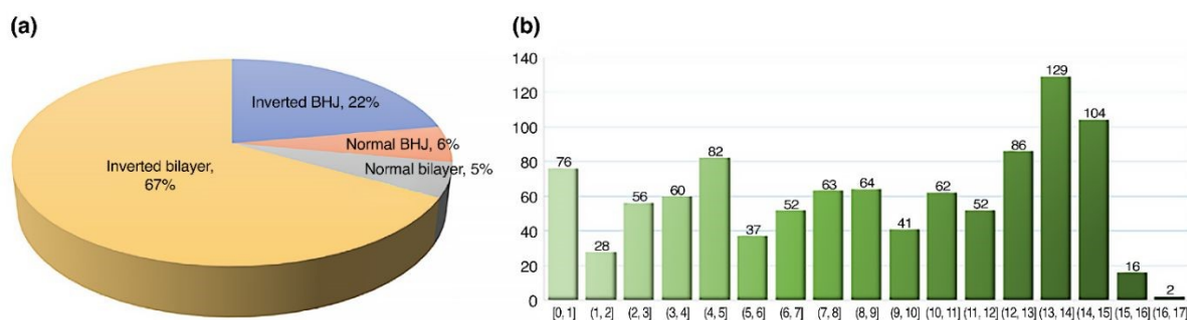


urately describe active-layer compositions, both the Donor1:Donor2 ratio and the overall (Donor1+Donor2):Acceptor ratio are recorded, enabling precise and reproducible representation of multicomponent formulations (Figure 2).



**Figure 2.** Schematic representation of donor–donor–acceptor (D1:D2:A) composition in a ternary organic photovoltaic system.

From the 13,430 devices stored on the established platform, non-fullerene data, data without missing values, and representative data selected by the experimenter from repeated experiments were selected. A dataset of 67 donors and 24 acceptors, totaling 1,010 D-A combinations (Table S1), was utilized for machine learning. The data included the OPV device architectures, donor and acceptor material structures, D:A ratios, device thicknesses, annealing conditions, and experimental PCE values (Figure 3).



**Figure 3.** (a) Distribution according to OPV device architecture; (b) distribution of PCE values.

In this study, we fragmented large organic molecules into structural subunits and extracted the molecular descriptors for each fragment. Calculations were performed for each fragment to further understand the chemical properties of the polymer material. The device information, process conditions, and polymer ratio information (D:A)



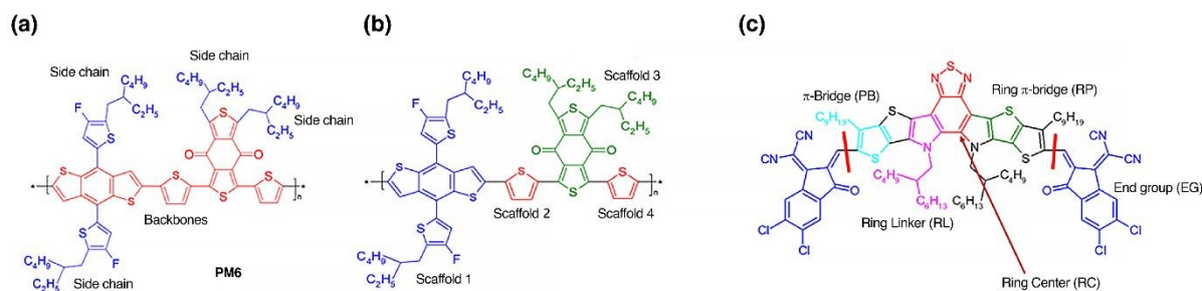
were added to the dataset. Furthermore, incorporating molecular symmetry, which has not been considered in previous studies, enables the extraction of structural characteristics that are not captured by conventional molecular descriptors. This approach enables the generation of diverse functions beyond device information and facilitates the processing of large molecules.

Because categorical features describing device structure and solvent type cannot be directly used in conventional machine learning models, they were converted into numerical values using label encoding. Integer labels were assigned as nominal identifiers without implying ordinal or physical meaning (**Table S2**). Missing values were imputed using the mode or median, depending on the feature.

## 2.2. Fragmentation of donors and acceptors

Most previous machine learning studies on organic photovoltaics (OPVs)<sup>30,31</sup> relied on the polymer fingerprints of the donor and acceptor materials in the active layer. The materials used as the OPV donors and acceptors are often polymers, single molecules, or macromolecules. This makes it difficult to infer their physicochemical properties using molecular descriptors. To overcome this limitation, we employed a fragmentation strategy to extract fragment-specific properties from the intrinsic molecular structure and subsequently recombine them, enabling machine learning models to better reflect the chemical characteristics derived from the materials themselves.

To digitally represent the donor and acceptor molecules in OPVs effectively, we developed our own fragmentation protocol. The fragmentation scheme was designed to simplify the polymer structures while retaining their essential chemical features, thereby producing independent fragments that contribute meaningfully to the overall molecular properties. (**Figure 4**).



**Figure 4.** (a) Definition and example of fragmentation of donor molecule (method 1). Red color represents the backbone, and blue color represents the side chains of the PM6 molecule. (b) Fragmentation using method 2. Each scaffold is expressed using a different color: blue for scaffold1, green for scaffold 3, red for scaffolds 2 and 4 of the PM6 molecule. (c) Definition of fragmentation of the acceptor molecule using the example of the Y6 molecule.

### [Fragmentation of Donor Molecules]

Donor molecules consist of a backbone and side chain. The backbone is defined as a continuous chain of arom



atic or non-aromatic rings, and the side chain consists of substituted rings, alkyl chains, or other functional groups. Donor fragmentation was performed using two methods.

Method 1: The backbone and side chains were explicitly separated and molecular descriptors were calculated independently for each structural unit.

Method 2 (scaffold-based fragmentation): The ring units that constitute the backbone are defined as scaffolds, and each scaffold represents a ring system connected by a single bond. Typically, a donor consists of two to eight scaffolds. For consistency, we standardized this number to four scaffolds. Each scaffold was further divided into a backbone and side chains.

[Fragmentation of the acceptor molecules]

The acceptor molecules contain a fused-ring (FR) system at their center and are divided into five parts based on this system.

1. Ring center (RC): The center of symmetry of the FR system.
2. Ring linker (RL): The ring units adjacent to the RC.
3. Ring  $\pi$ -bridge (RP): The  $\pi$ -bridge ring bonded to the outer ring; in some cases, the RP is absent.
4.  $\pi$ -Bridge (PB): The  $\pi$ -conjugated structure connecting the FR system and the end group; in some cases, the PB is considered a part of the FR system.
5. End group (EG): Terminal substituents located at the outermost positions of the acceptor molecule.

Through this fragmentation process, the acceptor molecule structure was defined as RC–RL–RP–PB–EG, centered on the central fused-ring system.

### 2.3. Extraction of molecular descriptors

The selection of appropriate molecular descriptors is a crucial step in developing reliable machine learning (ML) models. Considering computational efficiency and predictive performance, it is essential to employ descriptors that are both relevant and effective.

Accordingly, descriptors for each fragment unit were extracted using the RDKit and Stk libraries. For both Method 1 and Method 2, 217 physicochemical descriptors were computed using RDKit, whereas seven additional energy-related properties were obtained from Stk. These descriptors describe a wide range of molecular characteristics, ranging from basic molecular properties to complex structural characteristics. The complete list and aggregation methods are provided in **Table S3**.

### 2.4. Knowledge-based feature selection

[Operation of Descriptor]

Descriptors of each fragment extracted using RDKit and Stk were numbered and organized as shown in **Tables 1 and 2**, with the corresponding substructures illustrated in **Figures S2 and S3**.

Some molecules did not contain certain fragments and were therefore zero-coded. To mitigate this gap effect, pairwise computations between descriptors were performed.



This allowed us to maintain the effectiveness of each fragment-specific descriptor while compensating for the gap effect in fragment-free regions, resulting in molecular descriptors that utilize partial structures. The detailed computational operations for each descriptor type are provided in the Supporting Information **Table S3**.

**Table 1.** Examples of blank code in donors

Donor	BB1	SC1	BB2	SC2	BB3	SC3	BB4	SC4
PBDB-T	1	3	2	0*	5	10	2	0*
PCE-10	1	3	8	21	1	3	8	21
PM6	1	1	2	0*	5	10	2	0*
P3HT	2	22	2	22	2	22	2	22

**Table 2.** Examples of blank code in acceptors

Acceptor	RC	RL	RP	PB	SC
Y6	11	5	1	1	2
Y6-BO	11	5	1	1	3
CTIC-4F	10	0*	0*	11	2
CTIC-4Cl	10	0*	0*	11	3

\* Zero code owing to the nonexistence of fragments

#### [Local Symmetry States]

Materials used in OPVs (both polymers and small molecules) are often found to have symmetrical compound groups incorporated into their partial or entire structures for various reasons, such as their synthetic design or enhanced device performance. We noted the symmetry states of the scaffold of the donor molecule fragment and the ring centers of the acceptor molecules. Symmetry patterns from the donor and acceptor symmetry states were applied to the ML models.

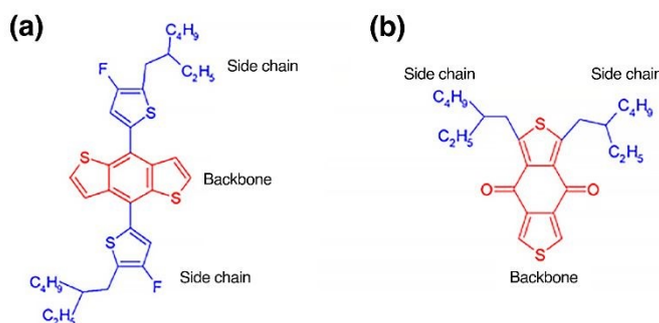
In this paper, local symmetry is defined in a restricted sense as either two-fold rotational ( $C_2$ ) or mirror-plane ( $\sigma_v$ ) symmetry, as illustrated in **Figure 5**. For donor molecules, local symmetry describes the symmetry of side-chain groups with respect to the molecular backbone. Donors are categorized into four scaffolds, yielding four symmetry states per donor; when two donor components are present, this results in a total of eight donor symmetry states. For acceptor molecules, local symmetry characterizes the symmetry of the end groups with respect to the ring center, with an additional symmetry state assigned at the ring center, leading to twenty symmetry states in total (**Table S4**).

The symmetry descriptors employed in this study are molecular-level features derived from specific structural motifs, rather than explicit representations of solid-state packing. Although thin-film packing is governed by processing conditions and intermolecular interactions, these symmetry states capture structural characteristics that



may indirectly relate to packing-relevant tendencies, while remaining general and scalable for machine-learning models.

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**Figure 5.** Examples of side chain symmetry: (a)  $C_2$  symmetric side chains, (b)  $\sigma_v$  symmetric side chains.

## 2.5. Selection of machine learning algorithms

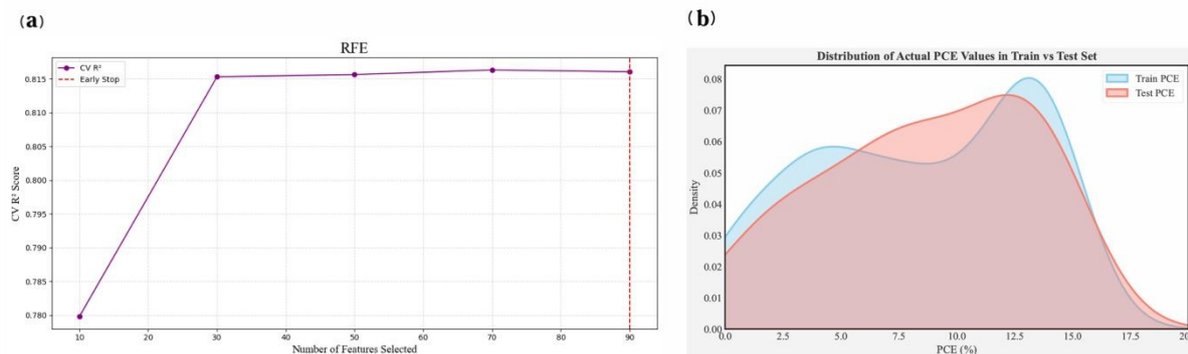
All machine-learning (ML) models were implemented within a unified Python framework using molecular descriptors generated with RDKit. XGBoost was selected as the primary model, and its performance was compared with random forest (RF), support vector regression (SVR), LightGBM (LGBM), and multilayer perceptron (MLP) models. Model performance was evaluated using the coefficient of determination ( $R^2$ ) under 5-fold cross-validation.

The dataset comprised 1,010 experimentally reported OPV devices and was split into training (80%) and test (20%) sets. To assess model robustness and minimize bias from a single data split, eight independent random seeds were used for repeated training and evaluation.

The initial feature space consisted of 1,054 molecular descriptors, resulting in a high feature-to-sample ratio. To reduce overfitting and computational cost, recursive feature elimination (RFE) was applied, with feature selection guided by cross-validated  $R^2$ . Feature removal was terminated once performance stabilized over consecutive iterations. Performance convergence was observed when approximately 90 descriptors remained (**Figure 6a**). Within this region, the final feature set was selected by identifying the subset that achieved the highest cross-validated  $R^2$  while maintaining consistent descriptor composition across successive RFE steps. Based on this criterion, 70 descriptors were retained for the final models.

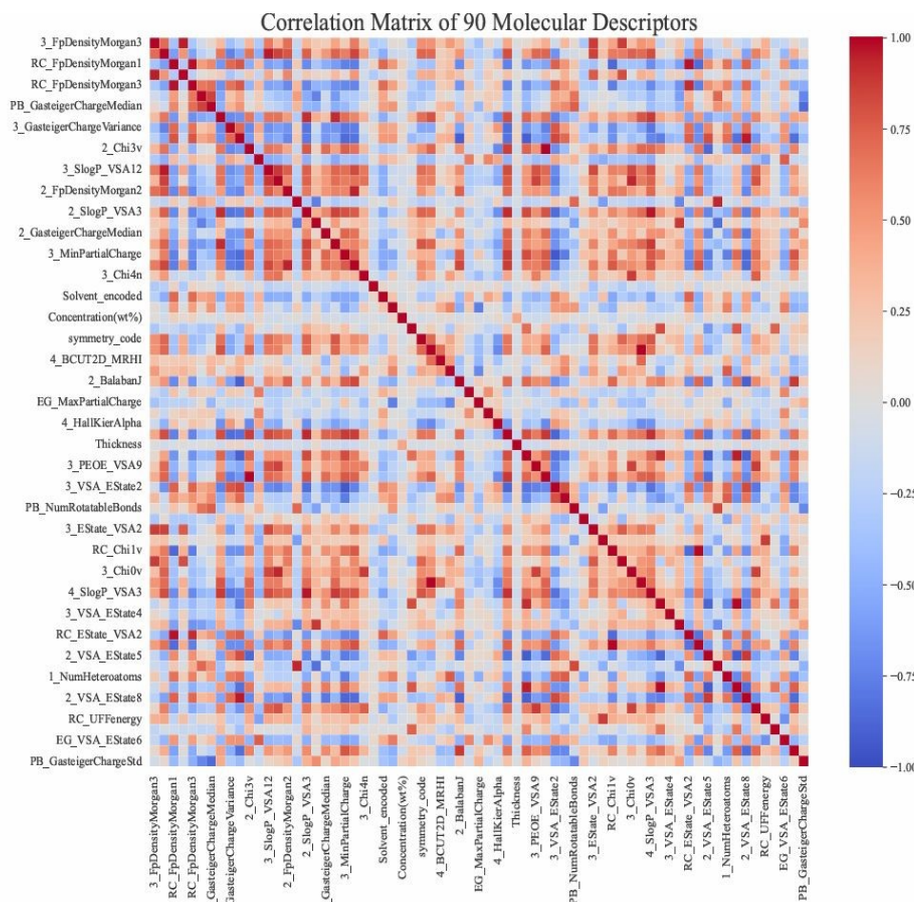
This reduction in feature dimensionality led to a substantial decrease in training time (from 101.7 s to 10.5 s) while maintaining comparable predictive performance ( $R^2$  of 0.848 and 0.839 before and after feature selection, respectively). The final descriptor sets for Methods 1 and 2 are summarized in **Tables S5 and S6**.





**Figure 6.** (a) Recursive feature elimination (RFE); (b) train and test dataset distribution in OPV prediction.

A Y-scrambling test was performed to assess dataset reliability. The original models achieved an average R<sup>2</sup> of 0.81, whereas the scrambled models yielded R<sup>2</sup> values close to zero (**Figure S8**), indicating that the predictions arose from genuine structure–property relationships rather than spurious correlations. Kernel density estimation (KDE) analysis showed similar PCE distributions for the training (n = 808) and test (n = 202) sets, confirming statistically balanced data partitioning. The small positive density near PCE = 0 originates from the smoothing nature of KDE rather than the presence of zero-valued samples (**Figure 6b**).



**Figure 7.** Pearson correlation matrix of the final 70 selected descriptors.

To assess multicollinearity among the selected descriptors, Pearson correlation analysis was performed on the f



inal 70-feature set. The mean and median absolute correlation coefficients ( $|r|$ ) were 0.344 and 0.295, respectively, indicating moderate overall correlations. Of the 2,415 possible descriptor pairs, only 25 pairs ( $\approx 1.0\%$ ) exhibited very high correlation ( $|r| > 0.9$ ), suggesting limited redundancy and minimal multicollinearity in the final feature set (**Figure 7**).

**Table 3.** R<sup>2</sup> scores for Method 1 and Method 2

Random state	Method 1	Method 2
0	0.879	0.841
42	0.854	0.885
150	0.782	0.842
500	0.892	0.841
790	0.847	0.758
1000	0.826	0.854
7500	0.881	0.830
10000	0.829	0.820
Average R <sup>2</sup>	0.849 ± 0.034	0.834 ± 0.034

[Comparative Analysis of ML Models for OPV Performance Prediction]

Two fragmentation strategies (Methods 1 and 2) were evaluated using machine learning models. As shown in **Table 3**, the XGBoost model achieved average R<sup>2</sup> values of 0.849 and 0.834 for Methods 1 and 2, respectively, indicating comparable predictive performance. Results for the other models are provided in **Table S7**.

The similar performance of the two strategies suggests that, for large donor–acceptor molecules where conventional RDKit descriptors are insufficient, fragmentation into smaller chemically meaningful units followed by recombination yields informative representations that enhance machine-learning training. Model robustness and generalization were further assessed using 5-fold cross-validation (**Tables S8, S9**).

**Table 4.** R<sup>2</sup> scores of ML methods

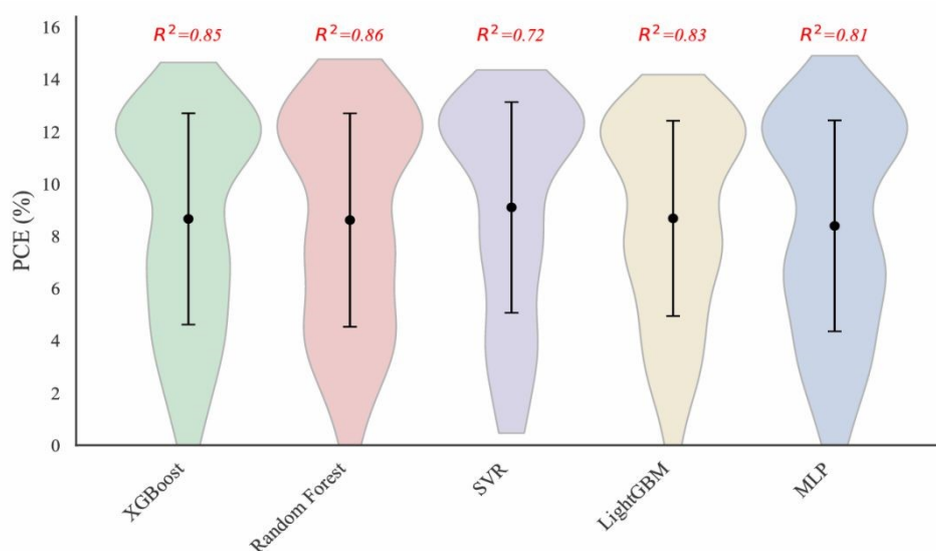
Random state	XGB	MLP	RF	SVR	LGBM
0	0.879	0.760	0.877	0.713	0.831
42	0.854	0.811	0.869	0.725	0.828
150	0.782	0.770	0.796	0.657	0.800
500	0.892	0.845	0.881	0.818	0.841



790	0.847	0.735	0.838	0.657	0.812
1000	0.824	0.752	0.823	0.692	0.780
7500	0.881	0.866	0.874	0.813	0.853
10000	0.829	0.738	0.829	0.650	0.748
Average $R^2$	$0.849 \pm 0.034$	$0.785 \pm 0.047$	$0.849 \pm 0.029$	$0.670 \pm 0.054$	$0.812 \pm 0.032$

**Table 4** summarizes the performance of the five machine-learning models using Method 1. XGBoost and RF achieved the highest average  $R^2$  values (0.849), followed by LGBM (0.812) and MLP (0.785), which exhibited moderate performance. SVR showed comparatively lower performance (0.718), consistent with its limited scalability in high-dimensional feature spaces.

Repeated experiments across eight random states yielded low standard deviations (0.029–0.054), indicating stable predictions with minimal sensitivity to data partitioning and a low risk of overfitting. These results are visualized using violin plots in **Figure 8**.



**Figure 8.** Comparative predictive performance of machine learning models for PCE.

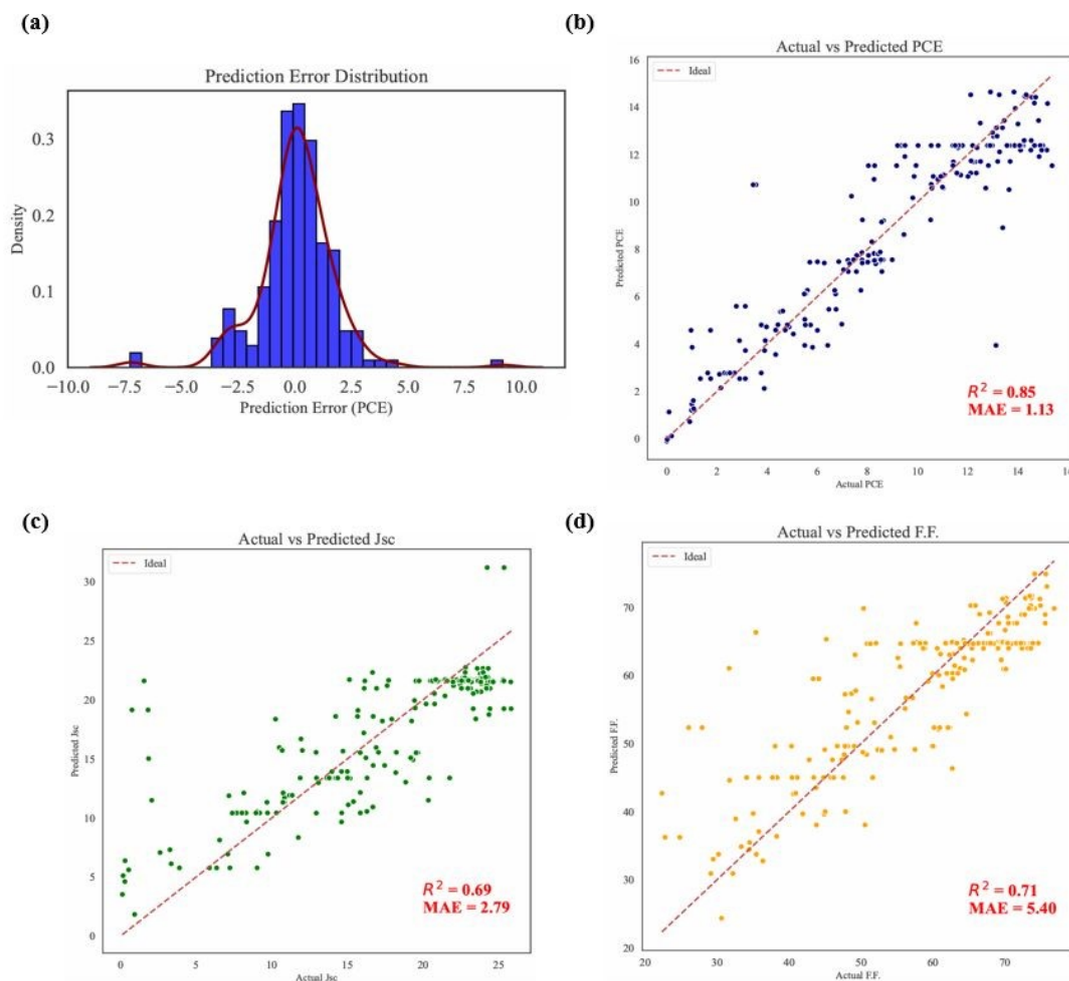
Using the experimental dataset, we further evaluated the model predictions for PCE and additionally explored the applicability of the model to Jsc and FF as auxiliary targets. The distribution of PCE prediction errors approximated a Gaussian profile, indicating minimal systematic bias (**Figure 9a**). Scatter plots comparing experimental and predicted values further illustrate the predictive capability of the model, yielding  $R^2$  values of 0.69 for Jsc and 0.71 for F.F (**Figures 9b–d**). The corresponding prediction errors, quantified by MAE and RMSE are summarized in **Table S10** for the XGBoost model evaluated on the test set.

In contrast, predictions for the open-circuit voltage ( $V_{oc}$ ) showed relatively low accuracy ( $R^2 = 0.2–0.4$ ). This li



mitation<sup>32-34</sup> is attributed to the descriptor set used in this study, which primarily captures molecular and electronic structure information but does not explicitly account for interfacial disorder, charge-transfer state energetics, or nonradiative recombination processes.

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**Figure 9.** (a) Distribution of prediction error for PCE, (b) scatter plots comparing predicted and actual values of PCE, (c) scatter plots comparing predicted and actual values of the short-circuit current density ( $J_{sc}$ ), and (d) scatter plots comparing predicted and actual values of the fill factor (F.F).

Finally, a comparison with two previous studies on fragmentation methods highlights the methodological novelty of our approach (Table 5). Rather than directly comparing identical datasets, this analysis focuses on differences in molecular representation and feature construction strategies. Kim et al.<sup>15</sup> used predefined, synthesis-oriented molecular fragments that efficiently organize structural diversity but treat fragments largely independently, whereas Wu et al.<sup>14</sup> employed a function-driven fragmentation scheme based on the electron push-pull principle, emphasizing electronic roles with limited structural uniformity across molecular classes.

In contrast, our study adopts a chemically meaningful fragmentation framework that incorporates explicit local symmetry information and integrates fragment-level features into a unified molecular representation. Combined with experimentally measured device parameters and polymer-specific chemical descriptors, this approach simultaneously captures electronic, structural, and processing effects, leading to improved predictive accuracy and en



hanced generalizability. Our model was evaluated using a held-out test set from an in-house experimental database, while the results from Kim *et al.*<sup>15</sup> and Wu *et al.*<sup>14</sup> were taken from their respective publications, each based on independently constructed datasets and evaluation protocols.

To further assess the generalization ability of the model, 130 data<sup>35-64</sup> points independently collected from the literature and not used during model training were extracted and evaluated using the same XGBoost model. Despite the inherent heterogeneity of literature data, including variations in material combinations and experimental conditions, the model achieved an  $R^2$  of 0.62, an MAE of 2.11, and an RMSE of 2.63, indicating reasonable predictive performance on unseen data (**Figure S9**).

**Table 5.** Comparison of ML model performances

<b>Our research</b>	<b>Algorithm</b>	XGBoost	RF	SVR	LGBM	MLP
	<b>Performance (r)</b>	0.93	0.94	0.85	0.91	0.93
<b>Kim <i>et al.</i> [15]</b>	<b>Algorithm</b>	XGBoost	RF	GBDT	LGBM	AdaB
	<b>Performance (r)</b>	0.85	0.86	0.86	0.86	0.86
<b>Yao Wu <i>et al.</i> [14]</b>	<b>Algorithm</b>	LR	RF	MLR	BRT	ANN
	<b>Performance (r)</b>	0.54	0.70	0.59	0.71	0.60

## 2.6. Feature importance analysis using SHAP

To enhance interpretability and quantitatively assess the contribution of individual features to PCE predictions, we performed a SHAP (SHapley Additive exPlanations) analysis on the XGBoost model trained using Method 1.

The SHAP analysis (**Figure 10**) identifies key features governing PCE prediction, which are primarily associated with molecular stability, local electronic structure, and processing conditions, highlighting the multiscale nature of performance-determining factors in organic photovoltaic devices.

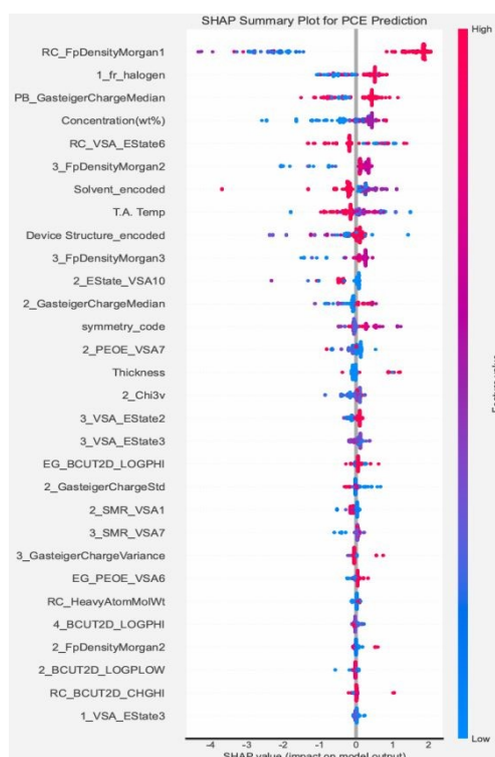
Among the top five features, RC\_FpDensityMorgan1 shows the strongest impact, indicating that dense and electronically coherent core environments favor efficient charge transport and reduced energetic disorder. 1\_fr\_halogen further highlights the positive role of halogen substitution, consistent with enhanced intermolecular interactions and frontier orbital tuning. PB\_GasteigerChargeMedian emphasizes the importance of balanced local charge distribution in promoting charge separation while suppressing recombination. The prominence of solution concentration(wt%) reflects its role as a key processing parameter that mediates thin-film formation and structure–property coupling, with its influence becoming significant in specific molecular contexts. RC\_VSA\_EState6 indicates that exposed electronic environments at the molecular core critically affect interfacial charge-transfer processes.



Beyond intrinsic molecular descriptors, several processing parameters including concentration, thermal annealing temperature, solvent type, device structure, and active-layer thickness also appear among the influential features. Although their average SHAP magnitudes are smaller, these parameters can become decisive depending on molecular structure, suggesting that they act as conditional modifiers of morphology, crystallinity, and charge-transport pathways. Notably, symmetry-related descriptors also contribute non-negligibly, implying that fragment-level local symmetry influences molecular packing regularity and orientational degeneracy.

Consistent with the SHAP results, the feature-importance rankings obtained from the XGBoost, RF, and LGBM models also highlight RC\_FpDensityMorgan1 and 1\_fr\_halogen as major contributors (**Figure S10, S11**), supporting the reliability of the SHAP-based interpretation across different learning algorithms.

Overall, the SHAP analysis demonstrates that PCE is governed by a synergistic interplay between molecular electronic structure, topology, symmetry, and processing conditions. The fragment-based, symmetry-aware descriptor framework effectively captures these coupled effects, providing both high predictive accuracy and physically interpretable structure–process property relationships.



**Figure 10.** SHAP-based feature importance summary for PCE prediction.

### 3. CONCLUSIONS

In this study, we established an OPV data platform to systematically collect experimental data and developed machine-learning models for predicting device performance. Fragment-based representations of donor and acceptor molecules were constructed using two fragmentation strategies, and 224 physicochemical descriptors were generated using RDKit and stk.

Among five machine-learning models evaluated, XGBoost showed the best performance with an  $R^2$  of 0.849.



Robust validation using five-fold cross-validation and multiple random states confirmed the reliability of the models. SHAP analysis identified local molecular symmetry as a key factor influencing PCE, highlighting its important role in OPV performance.

Future work will extend this framework by refining symmetry definitions and integrating external literature data to explore new high-efficiency donor–acceptor combinations.



## Data availability

The computational codes developed and used for this study are openly accessible via GitHub at [<https://github.com/juhyun7749/OPV-ML-2025>].

To ensure long-term preservation and reproducibility, a fixed version of the repository corresponding to the manuscript has been archived in Zenodo (DOI : <https://doi.org/10.5281/zenodo.18688051>)

All relevant datasets required to reproduce the analyses and figures presented in this article are included in the Zenodo archive.

The archived materials are provided under an open license to facilitate reuse and future research based on this work. The calculated molecular descriptors for the OPV donor and acceptor materials used in this study are openly available.

## Code availability

<https://github.com/juhyun7749/OPV-ML-2025>

## Author contributions

**J. L.:** Conceptualization, Data curation, Formal analysis, Methodology, Writing – original draft.

**H. B.:** Machine learning modeling, Methodology, Writing – review & editing.

**H. S.:** Code review, Validation.

**H. L.:** Experimental investigation, Sample preparation.

**F. A.:** Materials synthesis, Experimental data analysis.

**D. K.:** Conceptualization, Supervision, Project administration.

All authors contributed to manuscript review and revision.

## Competing Interests

The authors declare no competing interests.

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## Data availability

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The computational codes developed and used for this study are openly accessible via GitHub at [<https://github.com/juhyun7749/OPV-ML-2025>].

To ensure long-term preservation and reproducibility, a fixed version of the repository corresponding to the manuscript has been archived in Zenodo (DOI : <https://doi.org/10.5281/zenodo.18688051>)

All relevant datasets required to reproduce the analyses and figures presented in this article are included in the Zenodo archive.

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