

CrystalFramer: Rethinking the role of frames for SE(3)-invariant crystal structure modeling



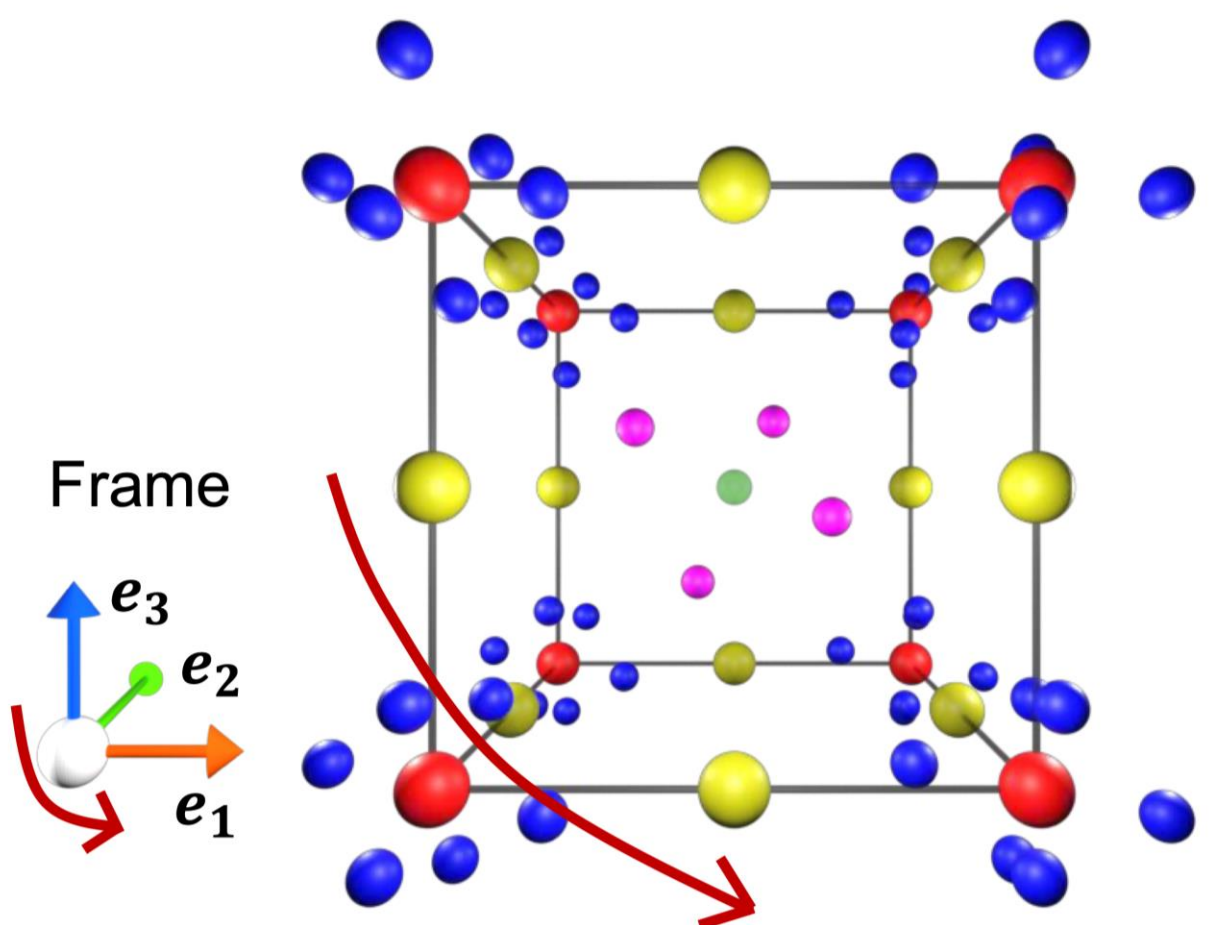
TL;DR: To make a GNN invariant to rotations, let's standardize the orientations of local atomic environments represented by internal self-attention weights!

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Frame-based SE(3)-invariant crystal structure modeling

The key to modeling crystal structures lies in learning **SE(3)-invariant (i.e., rotation & translation invariant) representations**.

💡 Why not normalize the orientation of the input structure?

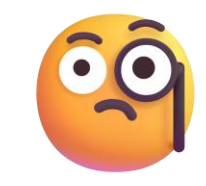
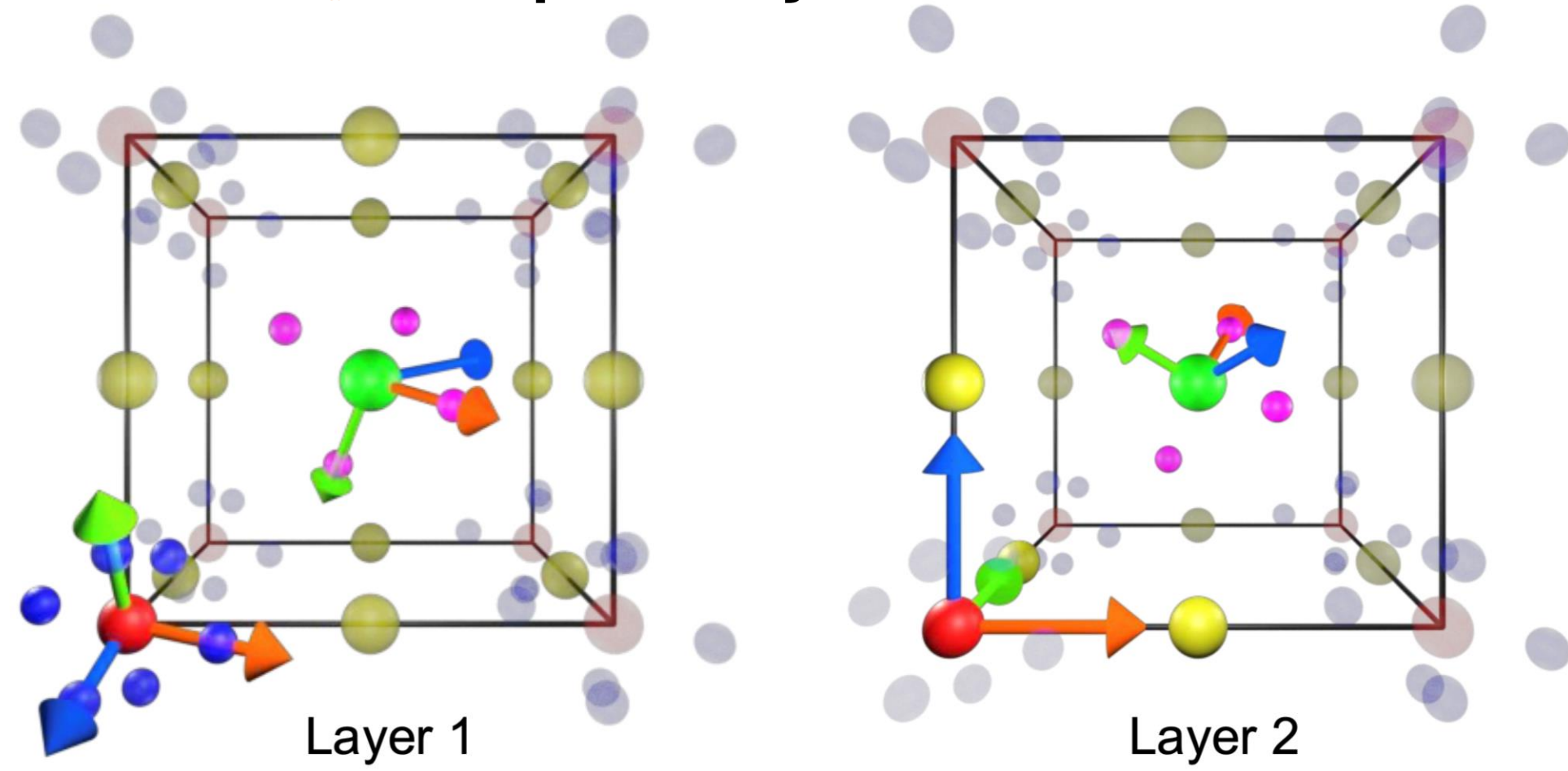


That's the "frame" (i.e., structure-aligned coordinate system)

- Ex 1) PCA for the positions of atoms in a unit cell (PCA frame)
- Ex 2) Lattice vectors of a unit cell (lattice frame)

- Can use richer information than interatomic distances
- No restriction on the network architectures
- Is it enough to just align with the structure?
- Is there no need to adapt the frame based on the task?

🚀 Proposed dynamic frames



The fundamental role of the frame is to *effectively incorporate relative positional information* into the message-passing layers for **modeling interatomic interactions**:

$$x'_i = \sum_j w_{ij} f_{i \leftarrow j}(x_i, x_j, r_{ij})$$



Let's consider **a frame aligned with the interatomic interactions** rather than with the structure itself.

Dynamic frames: attention-based local frames

Suppose a message-passing layer in a general form: $x'_i = \sum_j w_{ij} f_{i \leftarrow j}$.

Interaction weights w_{ij} can be interpreted as a **"mask" representing the local environment of the structure viewed from target atom i** .

Weighted PCA frame

For each atom i , compute the weighted covariance matrix of the direction vectors \bar{r}_{ij} pointing toward the surrounding atoms j .

$$\Sigma_i = \sum_j w_{ij} \bar{r}_{ij} \bar{r}_{ij}^T$$

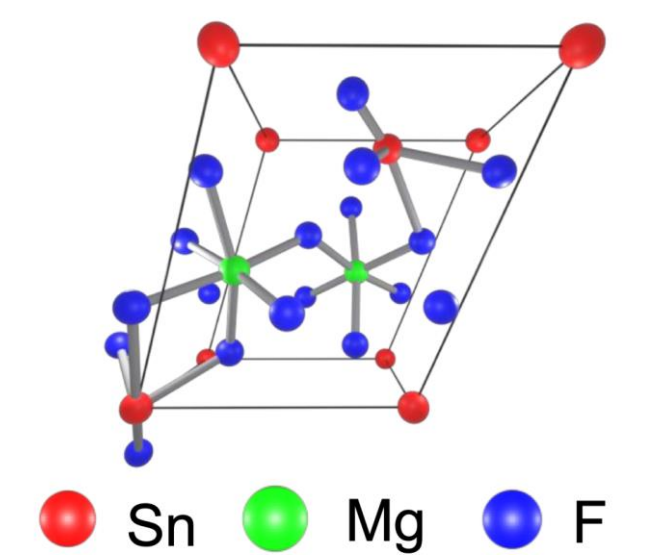
Set the orthonormal eigenvectors $[e_1, e_2, e_3]$ of the matrix as the three axes of the frame.

Max frame

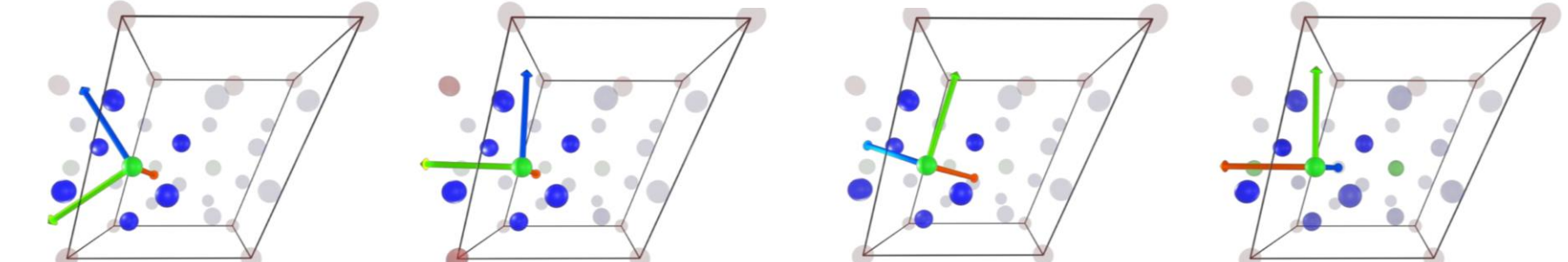
(Weight-sorted selection & orthogonalization)

1. Set the first axis e_1 to the direction \bar{r}_{ij} pointing toward the atom j with the highest weight w_{ij} .
2. To ensure diversity, select the direction \bar{r}_{ij} for the second axis candidate \hat{e}_2 by maximizing the adjusted weight $(1 - |e_1 \cdot \bar{r}_{ij}|)w_{ij}$, which penalizes alignment with e_1 .
3. Apply Gram-Schmidt orthogonalization to obtain $e_2 = \hat{e}_2 - (e_1 \cdot \hat{e}_2)e_1$.
4. Set $e_3 = e_1 \times e_2$ to form a right-handed orthonormal system.

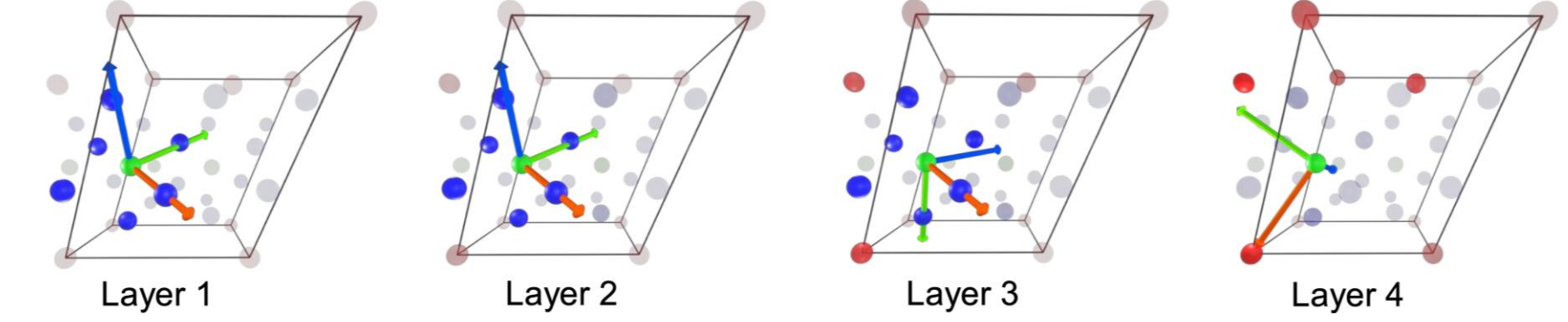
Crystal structure (MgSnF₄)



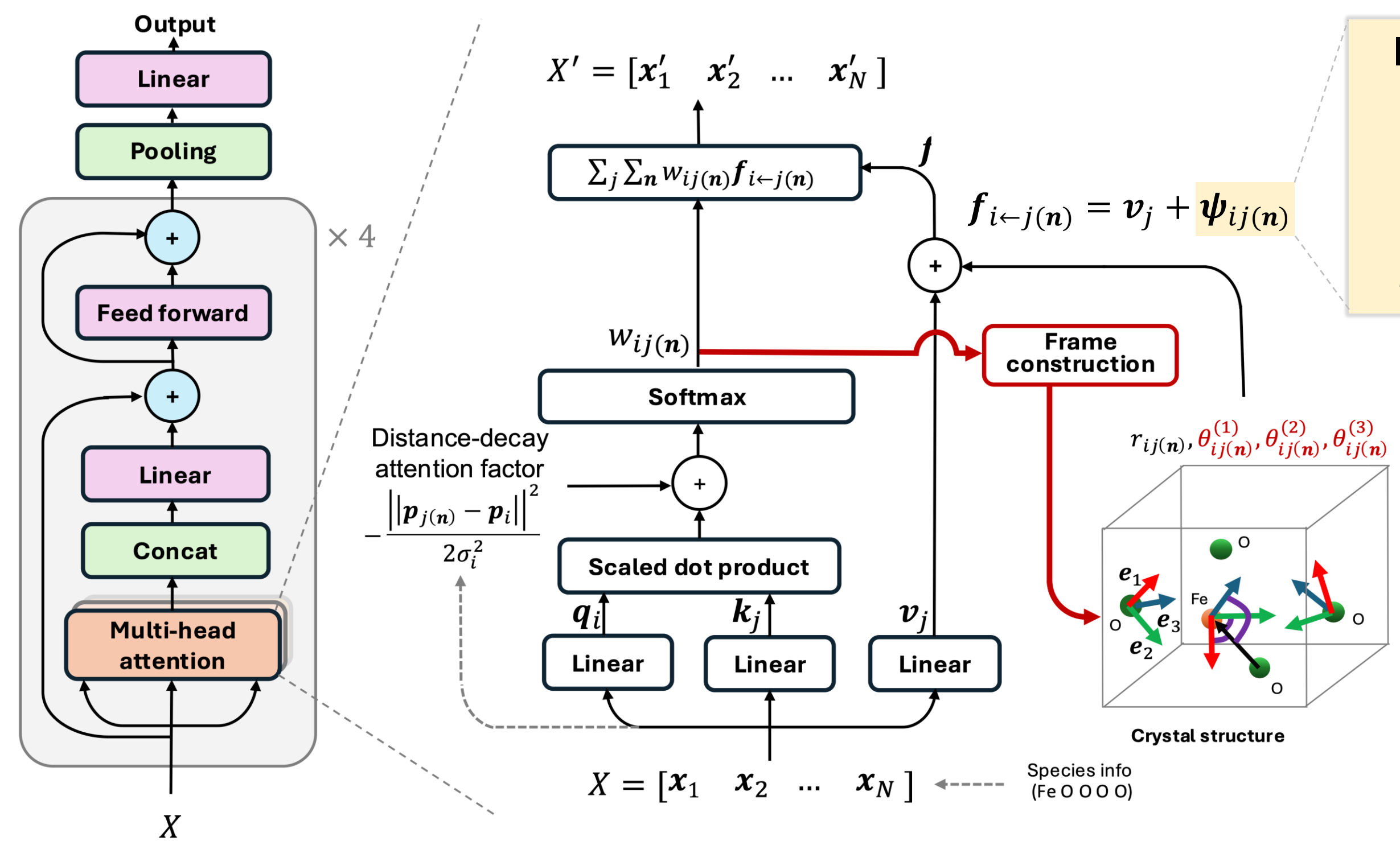
Weighted PCA frames



Max frames



CrystalFramer: Crystalformer (Taniai et al., 2024) + Dynamic frames



Invariant edge feature

Projection onto the frame coordinate system

$$\psi_{ij(n)} = W_0 b_{\text{dist}}(r_{ij(n)}) + \sum_{k=1,2,3} W_k b_{\text{angl}}(e_k^T \bar{r}_{ij(n)})$$

Distance encoding

Angle encoding

$b(x)$: Gaussian basis functions that convert a scalar value into a soft one-hot vector

In each layer, a frame is **dynamically constructed based on the atoms that the network is currently attending to**. The normalized relative positions $\bar{r}_{ij(n)}$ are projected onto the frame coordinate system as cosines of angles between frame axes e_k and $\bar{r}_{ij(n)}$, and used as **position encodings** $\psi_{ij(n)}$ for the Value vectors v_j .

Physical property prediction tasks

Method	Materials Project (MEGNET's snapshot)				JARVIS-DFT 3D				
	E form	Bandgap	Bulk mod.	Shear mod.	E form	E total	Bandgap(opt)	Bandgap(mbj)	E hull
	eV/atom	eV	log (GPa)	log (GPa)	eV/atom	eV/atom	eV	eV	eV
MatFormer (Yan et al., 2023)	0.021	0.211	0.043	0.073	0.0325	0.035	0.137	0.30	0.064
PotNet (Lin et al., 2023)	0.0188	0.204	0.040	0.065	0.0294	0.032	0.127	0.27	0.055
eComFormer (Yan et al., 2024)	0.0182	0.202	0.0417	0.0729	0.0284	0.032	0.124	0.28	0.044
iComFormer (Yan et al., 2024)	0.0183	0.193	0.0380	0.0637	0.0272	0.0288	0.122	0.26	0.047
Crystalformer (Taniai et al., 2024)	0.0186	0.198	0.0377	0.0689	0.0306	0.0320	0.128	0.274	0.0463
- w/ PCA frames (Duval et al., 2023)	0.0197	0.217	0.0424	0.0719	0.0325	0.0334	0.144	0.292	0.0568
- w/ lattice frames (Yan et al., 2024)	0.0194	0.212	0.0389	0.0720	0.0302	0.0323	0.125	0.274	0.0531
- w/ static local frames	0.0178	0.191	0.0354	0.0708	0.0285	0.0292	0.122	0.261	0.0444
- w/ weighted PCA frames (proposed)	0.0197	0.214	0.0423	0.0715	0.0287	0.0305	0.126	0.279	0.0444
- w/ max frames (proposed)	0.0172	0.185	0.0338	0.0677	0.0263	0.0279	0.117	0.242	0.0471
CrystalFramer (default)	0.0172	0.185	0.0338	0.0677	0.0263	0.0279	0.117	0.242	0.0471
CrystalFramer (lightweight)	0.0176	0.191	0.0341	0.0654	0.0268	0.0279	0.117	0.262	0.0467

Method	Time/epoch	Test/mater.	# Params.
MatFormer	60 s	20.4 ms	2.9 M
PotNet	43 s	313 ms	1.8 M
iComFormer	59 s	54.8 ms	5.0 M
Crystalformer	32 s	6.6 ms	853 K
CrystalFramer (default)	74 s	16.8 ms	952 K
CrystalFramer (lightweight)	43 s	15.2 ms	878 K



Incorporating dynamic frame-based edge features substantially enhances the prediction performance.



High model efficiency with only a small increase in parameters compared to the baseline Crystalformer.

The default ver encodes an angle into a 64-D vector using 64 Gaussian basis functions, while the lightweight ver uses a 16-D vec.