

Group-SAE: Efficient Training of Sparse Autoencoders for Large Language Models via Layer Groups

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Abstract

Sparse Autoencoders (SAEs) have recently been employed as a promising unsupervised approach for understanding the representations of layers of Large Language Models (LLMs). However, with the growth in model size and complexity, training SAEs is computationally intensive, as typically one SAE is trained for each model layer. To address such limitation, we propose *Group-SAE*, a novel strategy to train SAEs. Our method considers the similarity of the residual stream representations between contiguous layers to group similar layers and train a single SAE per group. To balance the trade-off between efficiency and performance, we further introduce *AMAD* (Average Maximum Angular Distance), an empirical metric that guides the selection of an optimal number of groups based on representational similarity across layers. Experiments on models from the Pythia family show that our approach significantly accelerates training with minimal impact on reconstruction quality and comparable downstream task performance and interpretability over baseline SAEs trained layer by layer. This method provides an efficient and scalable strategy for training SAEs in modern LLMs.

1 Introduction

Sparse Autoencoders (SAEs) (Makhzani and Frey, 2014) have recently emerged (Huben et al., 2024; Bricken et al., 2023) as a promising technique to tackle the polysemy of neurons in the activations of Large Language Models (LLMs) (Olah et al., 2020). SAEs decompose models’ activations into a sparse combination of human-interpretable directions, also called *features*. Despite the strengths in interpretability, SAEs face challenges that hinder their large-scale adoption (Sharkey et al., 2025). One of them is the high training and evaluation costs, which increase as model sizes and parameter counts grow. Notably, a

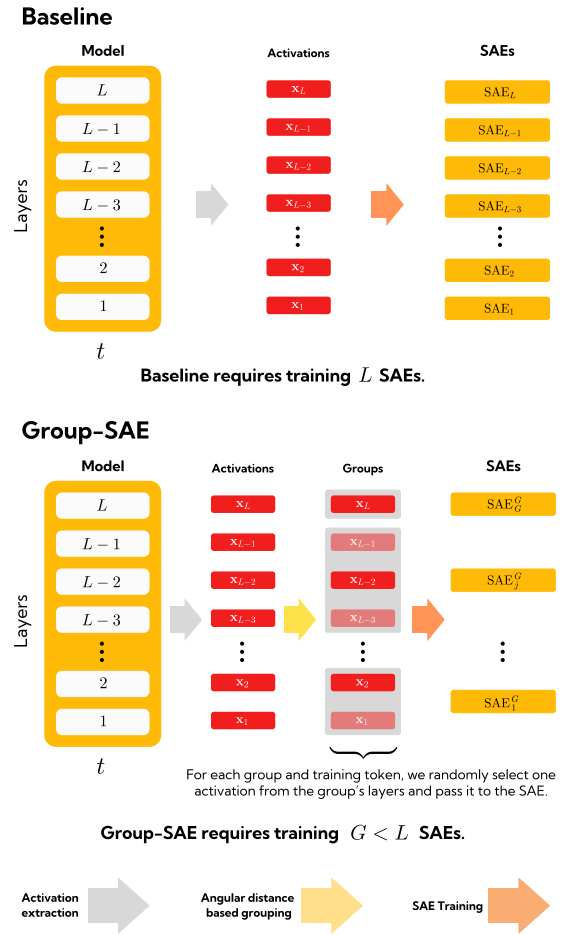


Figure 1: The illustration of our method. While standard training of SAEs requires training one per layer, our method first groups layers by angular similarity and then trains a single SAE for each group.

separate SAE is typically trained for each component (e.g., the output of the attention, the MLP, or a full transformer block) at every layer of an LLM, with a number of features that is a multiple of the dimensionality of the activation space of the model. For instance, a single SAE trained on the activations of a layer of Llama-3.1 8B (Grattafiori et al., 2024), with an expansion factor of 32, involves approximately $4096^2 \times 32 \times 2 \approx 1.073$ billion pa-

rameters. Such high computational demand increases training time and requires substantial hardware resources and energy consumption, making the approach increasingly impractical as models scale. Moreover, to make SAEs useful for interpretability, all their features have to be manually annotated. Even when using auto-interpretability techniques, this process can become unsustainable (Paulo et al., 2024b).

Facing such challenges, in this work we introduce **Group-SAE**, depicted in Figure 1, a method to reduce the computational overhead of training, evaluating, and interpreting SAEs. Our method leverages the similarity of the representations shared by close layers to reduce the total number of trained SAEs and uses a single SAE to reconstruct activations from different layers. The proposed technique follows primary observations that nearby neural network layers tend to learn similar levels of representations (Szegedy et al., 2014; Zeiler and Fergus, 2014; Jawahar et al., 2019). Shallow layers typically focus on capturing low-level features, while deeper layers are believed to learn high-level abstractions. In addition, Gromov et al. (2024) empirically shows that adjacent layers in LLMs could encode similar information.

Additionally, we introduce **AMAD** (Average Maximum Angular Distance), a novel empirical metric for selecting the optimal number of groups to partition a model’s layers—an important choice that balances SAEs quality and computational efficiency: more groups tend to improve reconstruction performance but reduce computational savings, while fewer groups offer greater efficiency at the cost of decreasing the quality of the reconstruction.

After thoroughly evaluating the reconstruction, downstream, and interpretability results of our methods on three models of varying sizes from the Pythia family (Biderman et al., 2023)—Pythia-160M, Pythia-410M, and Pythia-1B—we demonstrate that our method has several advantages over baselines. In particular, Group-SAE with AMAD finds an optimal tradeoff between training costs and quality of the SAE. It significantly reduces the number of trained SAEs, reducing training costs up to 50%. Moreover, such a novel approach only incurs a slight decrease in reconstruction quality and achieves comparable downstream performance. Additionally, Group-SAEs outperform standard SAEs matching the same computational cost. Finally, from an interpretability point of view, Group-SAEs offers the same, or even slightly better, level

of interpretability when compared with their baseline counterparts. Our **contributions** can be summarized as follows:

- We propose a novel method named **Group-SAE**, which partitions the layers of a model into groups and trains a single SAE for each group, thus significantly reducing the total number of SAEs to train.
- We introduce **AMAD** (Average Maximum Angular Distance), a new empirical metric for selecting the optimal number of groups, enabling an effective trade-off between computational efficiency and performance.

All the SAEs trained and used in our experiments, the code to train Group-SAE, and the code to replicate the experiments are all released as open source at <https://github.com/ghidav/group-sae>

2 Background and Related Work

2.1 Sparse Autoencoders

SAEs (Bricken et al., 2023) are a promising interpretability technique that decomposes dense LLM activations into a sparse combination of human-interpretable features. SAEs are based on two key intuitions. The first is the Linear Representation Hypothesis (LRH), which, supported by substantial empirical evidence (Mikolov et al., 2013; Nanda et al., 2023; Park et al., 2023), posits that Neural Networks (NNs) exhibit interpretable linear directions in their activation space. The second is the Superposition Hypothesis (SH), which assumes that observed NNs are dense compressions of a larger sparse model where each neuron corresponds to a specific feature (Elhage et al., 2022).

Within this framework, SAEs disentangle the effects of superposition, enabling the learning of interpretable linear directions in the model’s activations. Formally, given an activation $\mathbf{x} \in \mathbb{R}^n$, a SAE reconstructs it through two steps. First, it encodes the activation into the feature space as:

$$\mathbf{f}(\mathbf{x}) = \sigma (\mathbf{b}_e + \mathbf{W}_e (\mathbf{x} - \mathbf{b}_d)) \quad (1)$$

where $\mathbf{f}(\mathbf{x})$ represents feature activations, $\mathbf{b}_e \in \mathbb{R}^m$, $\mathbf{b}_d \in \mathbb{R}^n$ are bias terms, $\mathbf{W}_e \in \mathbb{R}^{m \times n}$ is the encoder matrix, and σ is an activation function. Typically, $m = c \cdot n$, with the expansion factor $c \in \{2^k \mid k \in \mathbb{N}_+\}$. $\sigma = \text{ReLU}$ was initially proposed (Bricken et al., 2023), with its

limitations that led to the development of two notable alternatives: TopK (Gao et al., 2024) and JumpReLU (Rajamanoharan et al., 2024).

The feature vector is then projected back into the model’s activation space:

$$\hat{\mathbf{x}} = \mathbf{b}_d + \mathbf{W}_d \mathbf{f}(\mathbf{x}) \quad (2)$$

where $\mathbf{W}_d \in \mathbb{R}^{n \times m}$ is the decoder matrix, with each column corresponding to a learned feature vector.

SAEs are trained to minimize the MSE between original activations and SAE reconstruction. To enforce feature sparsity, an additional penalty is usually included in the loss function, either as the L_1 norm (Bricken et al., 2023) or the L_0 norm (Rajamanoharan et al., 2024) of $\mathbf{f}(\mathbf{x})$, scaled by a positive factor λ , termed the *sparsity coefficient*. Formally, the loss function can be written as:

$$\mathcal{L}(\mathbf{x}) = \|\mathbf{x} - \hat{\mathbf{x}}\|_2^2 + \lambda \|\mathbf{f}(\mathbf{x})\|_s \quad (3)$$

with $s \in \{0, 1\}$. On the other hand, when using TopK (Gao et al., 2024), no additional loss components are needed, as the activation function inherently enforces sparsity.

2.2 Shared SAEs

While SAEs were originally designed to reconstruct activations from a single model component (e.g., the output of a specific layer, MLP, or Attention), subsequent approaches have explored their application to activations from multiple layers. For instance, Yun et al. (2023) and Lawson et al. (2024) employed a single SAE to reconstruct activations from all residual stream layers of a model, aiming to analyze how features evolve across layers. More recently, Lindsey et al. (2024) extended this concept by introducing *Crosscoder*, a modified SAE architecture that creates a unified representation of computations across multiple layers.

These methods are driven by empirical evidence suggesting that information in LLMs is often shared and rather redundant across nearby layers (Phang et al., 2021; Gromov et al., 2024). In this work, we leverage this principle to explore the optimal balance between performance and computational efficiency when applying SAEs to multiple layers.

2.3 Improving SAE efficiency

As highlighted by Sharkey et al. (2025), one of the major challenges of SAEs is their high training and evaluation costs. As previously mentioned,

SAEs scale alongside model size, making them impractical for low-resource settings. Furthermore, interpreting the meaning of SAE features presents an additional challenge. Even with automated techniques, interpretation costs can reach thousands of dollars (Paulo et al., 2024b).

To mitigate training costs, Gao et al. (2024) investigated the scaling laws of SAEs to determine the optimal balance between model size and sparsity. Recent work has also explored transfer learning as a means to enhance SAE training efficiency. For instance, Kissane et al. (2024) and Lieberum et al. (2024) demonstrated that SAE weights can be transferred between base and instruction-tuned versions of Gemma-1 (Team et al., 2024a) and Gemma-2 (Team et al., 2024b), respectively. Additionally, Ghilardi et al. (2024) showed that transferability also occurs within different layers of a single model, both in forward and backward directions.

3 Group-SAE

In our approach, a **Group-SAE** is defined as a sparse autoencoder that is trained to reconstruct the activations from multiple layers that have been grouped together, rather than training an individual SAE for each layer. This grouping leverages the observation that nearby layers tend to exhibit similar activation patterns (Gromov et al., 2024). A detailed analysis of this phenomenon can be found in Appendix C (Figures 5, 6, and 7).

3.1 Clustering layers into groups

For a model with L layers, there are theoretically $G! \cdot S(L, G)$ ways to partition the layers into G groups—where $S(L, G)$ denotes the Stirling number of the second kind. Because this number grows rapidly with model depth, we instead employ an agglomerative clustering strategy based on angular distances between layers to efficiently determine a suitable grouping. Specifically, following the formulation in (Gromov et al., 2024), we compute the mean angular distance between the residual activations of each layer using a subset of the training set used to train the SAEs (see Appendix C for detailed measurements)¹. We then apply a bottom-up hierarchical clustering method with complete linkage (Nielsen and Nielsen, 2016). At each step, the two groups with the smallest inter-group dis-

¹We precisely use 10M tokens from the training set used to train the SAEs, which amount to 1% of the total training tokens.

tance²are merged. This merging continues until exactly G groups remain, ensuring that within each group the maximum angular distance is minimized.

In addition to being motivated by recent work (Gromov et al., 2024; Li et al., 2025), we adopt angular distance as our similarity metric because it captures the directional component of activations, which is key to their sparse representation. As in our setting, feature directions are typically normalized to unit norm, so SAE feature activations are proportional to the cosine of that angle. Consequently, activations that are close in angular distance tend to activate similar features and can be effectively reconstructed by the same SAE. Empirical evidence supports this: reconstruction quality degrades when SAEs are applied to activations from more distant layers (Ghilardi et al., 2024), and this degradation correlates with larger angular distances.

3.2 Selecting the number groups G

The choice of G , the number of groups of layers, is an important choice to make in our method as it influences both computational savings and the quality of the SAE. To guide this choice, we propose an empirical score called the **Average Maximum Angular Distance** (AMAD), defined as

$$\text{AMAD}(G) = \frac{1}{G} \sum_{g=1}^G D_g, \quad (4)$$

where D_g is the maximum angular distance between any pair of activations within group g . AMAD thus quantifies, on average, the worst-case dissimilarity within groups. Intuitively, when G is small, each group aggregates more distant layers, which increases AMAD; conversely, when $G = L$ (one group per layer), AMAD becomes zero but no computational savings are achieved. The goal is therefore to select the smallest G such that the groups remain sufficiently homogeneous, i.e., $\text{AMAD}(G)$ stays below a target threshold. Formally, we select

$$\hat{G} = \min\{G \mid \text{AMAD}(G) < \theta\},$$

where θ is a distance threshold. Based on our empirical analysis (Section 4), we set $\theta = 0.2$, which provides a robust trade-off across model sizes. This

²In complete linkage, the inter-group distance is defined as $D(X, Y) = \max_{\mathbf{x} \in X, \mathbf{y} \in Y} d_{\text{angular}}(\mathbf{x}, \mathbf{y})$ for groups X and Y .

criterion ensures maximal grouping (hence computational savings) while avoiding the sharp increase in reconstruction error that arises when groups mix overly distant layers.

3.3 Computational Savings

The computational cost, in FLOPs, of training a SAE can be divided into two main components:

- *Activation caching (A):* FLOPs required to generate the model’s activations, which are used for training the SAE.
- *SAE training (T):* FLOPs involved in optimizing a single SAE using the cached activations.

Thus, the total cost of training SAEs across all residual stream layers of a model is given by $A + LT$. Since both baseline and Group-SAEs share the same architecture and undergo the same training process for a single SAE, the total cost of training all Group-SAEs is $A + GT$ ³.

The resulting compute savings, $\Delta(G)$, quantifying the relative change in total FLOPs when applying Group-SAEs instead of per-layer SAEs, is defined as:

$$\Delta(G) = 1 - \frac{A + GT}{A + LT}. \quad (5)$$

By definition, if $G = L$, then $\Delta(G) = 0$, meaning no savings. Conversely, as G decreases, savings increase, reaching a maximum of $(LT - T)/(A + LT)$ when $G = 1$, which approaches $(L - 1)/L$ as the ratio T/A increases.

Since our method does not alter either A or T , the efficiency gains of Group-SAEs are primarily determined by the G/L ratio.

4 Experiments

Our work is primarily focused on addressing the following research questions:

Q1 Do SAEs trained on groups of layers activations maintain reconstruction quality and downstream performance?

Q2 Does selecting the number of groups G based on the Average Maximum Angular Distance (AMAD) ensure an optimal balance between computational efficiency and model performance?

³We do not account for the cost of computing angular distance when selecting groups, as we rely on activations already sampled for training, making the additional computational overhead negligible.

Q3 How do Group-SAEs affect the interpretability of the SAE latent representations?

To address these questions, we compare the performance of standard SAEs and Group-SAEs across a range of metrics and alternative grouping strategies.

4.1 Experimental setting

We denote SAE_l as the baseline SAE trained to reconstruct the activations of layer l . For every $g = 1, \dots, G$, with $G \in \{1, \dots, L - 1\}$ and L being the number of layers of a model, let $[gG]$ represent the set of layers belonging to the g -th group within the partition of G groups. We then define SAE_g^G as the SAE trained to reconstruct the activations for all layers in $[gG]$. To ensure a fair comparison with baselines, we allocate 1 billion training tokens for each SAE_l and SAE_g^G . For baseline SAEs, activations are always taken from a single fixed layer. In contrast, for Group-SAEs, activations are drawn from a randomly selected layer within the set $[gG]$. In this way, we ensure that each Group- and baseline SAEs process exactly 1 billion tokens and activations.

Models, Dataset and Hyperparameters Following [Lawson et al. \(2024\)](#), we train both SAEs and Group-SAEs with the Fraction of Variance Unexplained (FVU) as reconstruction loss. Defined as

$$\text{FVU}(\mathbf{x}) = \frac{\|\mathbf{x} - \hat{\mathbf{x}}\|_2^2}{\text{Var}(\mathbf{x})}, \quad (6)$$

we prefer it to standard MSE loss as it accounts for the different magnitudes of activations coming from different layers of the model. We employ Top- K activation⁴ with $K = 128$ and expansion factor of $c = 16$ on the residual stream after the MLP contribution of three models of varying sizes from the Pythia family ([Biderman et al., 2023](#)): Pythia 160M, Pythia 410M, and Pythia 1B.

We follow established practice in SAE training ([Bricken et al., 2023; Gao et al., 2024; Rajamanoharan et al., 2024](#)) and use the same pre-training dataset as the models themselves. In particular, we sample 1 billion tokens from the Pile dataset ([Gao et al., 2020](#)) and process them with a context size of 1024.

For each model, we compute all partitions $G \in \{1, \dots, L - 1\}$ and train a Group-SAE for all

groups of layers in them. We exclude the last layer from all partitions because it resides in the unembedding space and, based on our empirical findings, consistently exhibits a distinct reconstruction error pattern. As a result, it requires a separate SAE. Additionally, we compare our grouping strategy with two **baseline techniques** aimed to reduce the computational cost of training SAEs: (1) training Group SAEs on evenly spaced groups, and (2) training smaller SAEs on all layers. Hyperparameters for all the experiments and training details can be found in Appendix A and B respectively.

Evaluation. As in previous work ([Huben et al., 2024; Gao et al., 2024](#)), we evaluate quality of trained SAE across three key areas: reconstruction, downstream, and interpretability.

For both reconstruction and downstream evaluations, we use a subset of the Pile dataset (distinct from the training set) comprising 1 million tokens.

For reconstruction, we compare each SAE_g^G with its corresponding baseline SAE_l for every layer $l \in [gG]$. We report the average Fraction of Variance Unexplained (FVU, Equation 6) as our reconstruction metric.

To evaluate downstream performance, we measure the effect of replacing a layer’s activation with its SAE reconstruction on the next-token prediction. Specifically, we compute the average relative change in next-token Cross-Entropy:

$$\Delta \text{CE} = \frac{\text{CE}(\mathbf{M}(P \mid \mathbf{x}^l \leftarrow \hat{\mathbf{x}}^l)) - \text{CE}(\mathbf{M}(P))}{\text{CE}(\mathbf{M}(P))}, \quad (7)$$

where \mathbf{M} denotes the model, P is the input prompt, and $\mathbf{M}(P \mid \mathbf{x}^l \leftarrow \hat{\mathbf{x}}^l)$ indicates the model output when the true activation \mathbf{x}^l at layer l is replaced with the SAE reconstruction $\hat{\mathbf{x}}^l$.

For interpretability, we adopt the automated pipeline proposed by [Paulo et al. \(2024a\)](#). First, an *explainer* language model (LM) generates natural language explanations of the SAE latent representations. Then, a separate *scorer* LM evaluates these explanations. In our experiments, both the explainer and scorer are implemented using `gemini-2.0-flash-001`⁵. Specifically, for each SAE, we randomly sample 64 features and cache their latent activations over a 10M token sample from the Pile. For each latent, the explainer is shown 20 distinct examples, 10 activating the latent and 10 sampled randomly, each consisting of

⁴The Top- K activation function is directly applied on the features obtained with Equation 1, where $\sigma = \text{Top-}K \circ \text{ReLU}$.

⁵[https://deepmind.google/technologies/gemini/flash/](https://deepmind.google/technologies/gemini/)

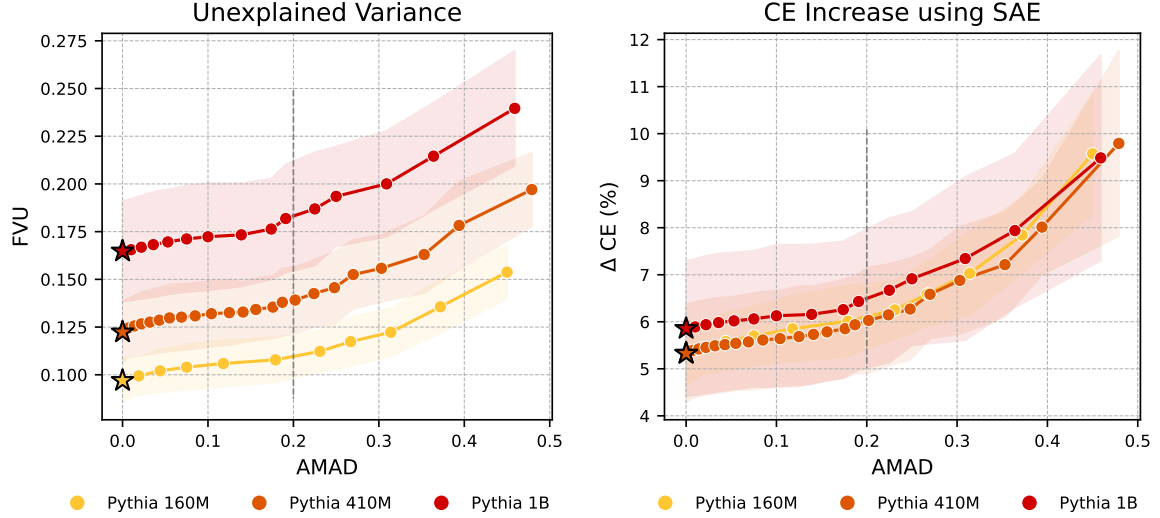


Figure 2: (Left) FVU and (Right) $\Delta\text{CE}(\%)$ over $\text{AMAD}(G)$ for every $G \in \{1, \dots, L - 1\}$. The highlighted star markers represent the baseline SAEs (i.e., with no grouping), while the other points correspond to Group-SAEs, ordered from left to right by increasing AMAD, which reflects a decrease in the number of groups. The shaded area indicates one std.

Table 1: FVU and ΔCE for different approaches across model sizes. Our proposed grouping strategy based on the AMAD achieves lower FVU and ΔCE compared to the baselines: Group SAEs with evenly spaced groups and smaller SAEs trained on all layers. Note that both the *Evenly Spaced* and *Smaller SAEs* strategies have the same number of training FLOPs as our AMAD-based grouping strategy. For each entry, the value in % shown to the right indicates the relative improvement over *Smaller SAEs (All layers)* baseline (positive = better).

Approach	Pythia-160M		Pythia-410M		Pythia-1B	
	FVU	$\Delta\text{CE}\%$	FVU	$\Delta\text{CE}\%$	FVU	$\Delta\text{CE}\%$
Group SAEs (AMAD with \widehat{G} groups)	0.108 (+6.1%)	6.01 (+18.5%)	0.138 (+5.5%)	5.94 (+16.3%)	0.182 (+3.2%)	6.43 (+20.6%)
Group SAEs (Evenly spaced with \widehat{G} groups)	0.114 (+0.9%)	5.40 (+26.7%)	0.145 (+0.7%)	6.01 (+15.4%)	0.189 (-0.5%)	6.63 (+18.1%)
Smaller SAEs (All layers)	0.115 (+0.0%)	7.37 (+0.0%)	0.146 (+0.0%)	7.10 (+0.0%)	0.188 (+0.0%)	8.10 (+0.0%)

32 tokens. Two binary scoring strategies are employed:

- *Detection*: A language model determines whether a given sequence activates an SAE latent according to the provided explanation.
- *Fuzzing*: Activating tokens are marked within each example, and a language model is prompted to assess whether the marked sentences are correctly identified.

Figure 8 in Appendix D shows a sentence example for each strategy. For every metric (FVU, ΔCE and Detection/Fuzzing) and for each $G \in \{1, \dots, L - 1\}$, we first compute all metrics at the layer level, then aggregate the results for each partition g within G by computing the mean and the standard deviation weighted by the number of layers in that partition.

4.2 Results

In the following paragraphs, we aim to empirically answer the research questions outlined in Section 4.

Q1: What is the impact of grouping layers (Group-SAEs) on reconstruction quality and downstream task performance? In Figure 2, we plot the average FVU and the cross-entropy difference (ΔCE) as functions of the AMAD for different group configurations. The highlighted star markers represent the baseline models (i.e., with no grouping), while the other points correspond to grouped models. The points are ordered from left to right by increasing AMAD, which reflects a decrease in the number of groups, with G ranging from $L - 1$ down to 1. From Figure 2, a notable turning point emerges around $\text{AMAD}(G) \approx 0.2$: increasing AMAD beyond this threshold leads to a more rapid loss in performance. In particular, train-

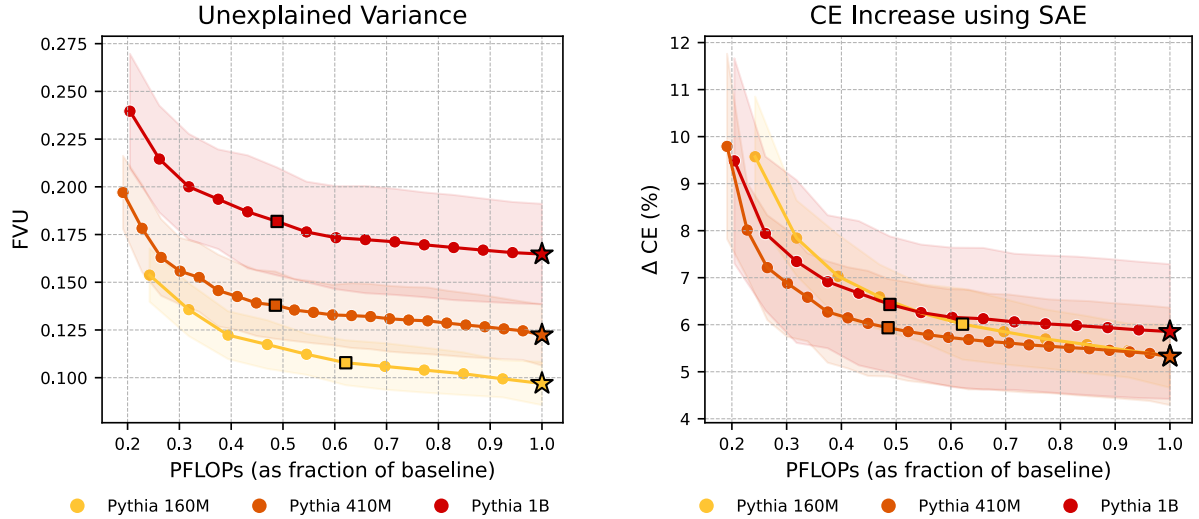


Figure 3: (Left) FVU and (Right) $\Delta\text{CE}(\%)$ over the fraction of training PFLOPs with respect to the baseline. The highlighted star markers represent the baseline SAEs (i.e., with no grouping), while the other points correspond to Group-SAEs, ordered from right to left by decreasing PFLOPs, which reflects a decrease in the number of groups. The highlighted square markers represent the Group-SAEs with a number of groups $\hat{G} = \min\{G \mid \text{AMAD}(G) < 0.2\}$.

ing a single SAE on all the model layers ($G = 1$), although achieving the best computational saving, also incurs the worst reconstruction and downstream performance.

To further validate our method, in Appendix E, we inspect the quality of features learned by Group-SAEs by measuring their similarity to the features learned by Baseline-SAEs. As expected, for each baseline SAE $_l$, we found average similarity to peak with the Group-SAE trained on a group containing l . Moreover, Appendix F analyzes how a Group-SAE’s features distribute across the activations of layers in its group: consistent with the analysis of Lindsey et al. (2024), individual features typically peak at a specific layer yet exhibit substantial spread to adjacent layers. This pattern supports our core hypothesis that while features may anchor to particular layers, they remain relevant across neighboring ones, enabling a single SAE to effectively reconstruct activations from multiple, similar layers.

Q2: Does selecting the number of groups G based on the AMAD ensure an optimal balance between computational efficiency and model performance? Motivated by the insights from the previous paragraph, the optimal G is chosen as $\hat{G} = \min\{G \mid \text{AMAD}(G) < 0.2\}$. In Figure 3 we show both FVU and ΔCE plotted against the fraction of PFLOPs relative to the baseline. Again,

star markers denote baseline SAEs, whereas circles represent Group-SAEs. Here, moving from right to left indicates reducing PFLOPs (i.e., training fewer SAEs overall). The points are ordered from right to left by decreasing PFLOPs, which reflects a decrease in the number of groups, from $L - 1$ down to 1. The highlighted square markers correspond to Group-SAEs with \hat{G} groups; they substantially reduce training costs up to more than 50% with only a moderate performance penalty: $\text{FVU}(\star) - \text{FVU}(\blacksquare) \approx -0.015$ and $\Delta\text{CE}(\star) - \Delta\text{CE}(\blacksquare) \approx -0.62$ for all three evaluated models.

To ensure that our grouping strategy and the selection of \hat{G} based on AMAD offer an effective trade-off between computational efficiency and performance, we compare them against two baselines: 1) *Evenly Spaced Group SAEs*: Group SAEs trained such that each partition contains nearly equal numbers of layers; 2) *Smaller SAEs*: A separate, smaller SAE is trained for each layer. All methods are adjusted to incur equal computational costs⁶. Results in Table 1 shows that the proposed method outperforms the two additional baselines across nearly all models and evaluation metrics, with only a single exception observed in the case of Pythia-160M.

⁶For Evenly Spaced Group SAEs, we use the same number of groups \hat{G} ; for Smaller SAEs, we set the expansion factor as $c' = c \cdot \hat{G}/T$, matching the FLOPs of a Group-SAE with \hat{G} groups.

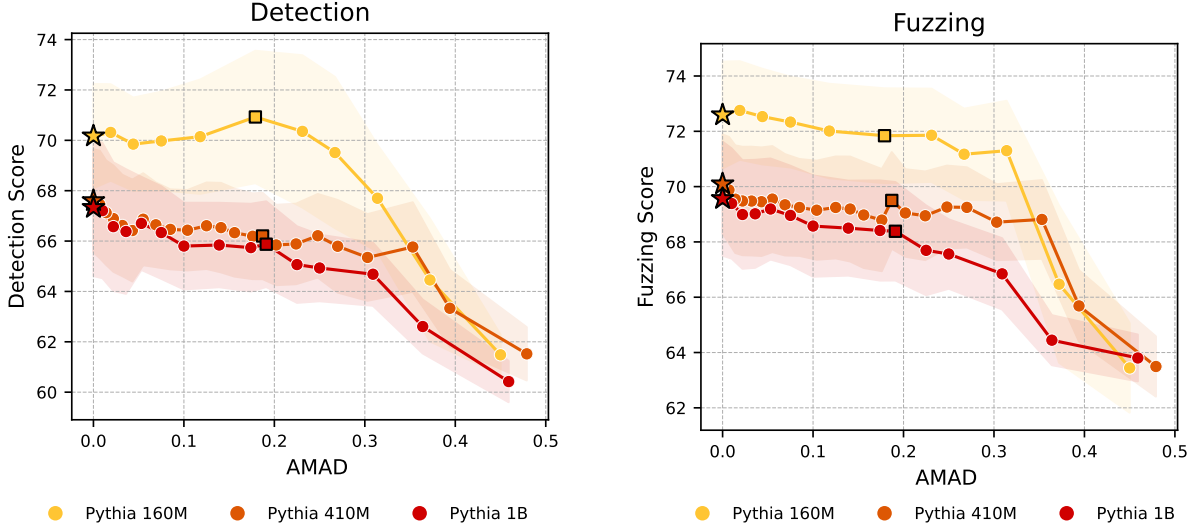


Figure 4: Auto-Interpretability scores following the automated pipeline defined by (Paulo et al., 2024b) over $\text{AMAD}(G)$ for every $G \in \{1, \dots, L - 1\}$. The highlighted star markers represent the baseline SAEs (i.e., with no grouping), while the other points correspond to Group-SAEs, ordered from left to right by increasing AMAD , which reflects a decrease in the number of groups. The highlighted square markers represent the Group-SAEs with a number of groups $\hat{G} = \min\{G \mid \text{AMAD}(G) < 0.2\}$. (Left) Detection and (Right) Fuzzing scores, as defined in the Evaluation paragraph of Section 4.

Table 2: Comparison of FLOPs (10^{18}) required for caching activations and training Baseline and Group SAEs on 1B tokens, covering all layers with an expansion factor of 16 and $\hat{G} = \min\{G \mid \text{AMAD}(G) < 0.2\}$.

Model	\hat{G}	A+LT	A+GT	$\Delta\%(\hat{G})$
Pythia 160M	6	1.34	0.77	+42.5%
Pythia 410M	9	4.73	2.21	+53.3%
Pythia 1B	6	12.48	5.77	+53.7%

Importantly, this exception does not arise from the idea of grouping layers but from the chosen grouping strategy. Indeed, our method consistently outperforms the standard per-layer approach with smaller Standard SAEs. We observe that these advantages are particularly noticeable for the ΔCE metric, related to the downstream performance. Additionally, Table 2 presents the computational costs and savings, as defined in Eq. 5 of Section 3.3, of Group-SAEs compared to the baselines when the optimal number of groups G is selected as \hat{G} .

Q3: How do Group-SAEs affect the interpretability of the SAE latent representations?

To answer, we employ the auto-interpretability pipeline proposed by (Paulo et al., 2024b). For each SAE latent, first, an *explainer* Language Model is asked to propose a natural language explanation of

it given both activating and non-activating examples. Then, given the explanation, a *scorer* Language Model is tasked with predicting the set of sentences that should activate the target latent (*detection*) and the sentences containing highlighted tokens that activate the target latent (*fuzzing*). In Figure 4 we plot both the detection and fuzzing scores for all the evaluated models. In the figures, square markers denote Group-SAEs with \hat{G} groups, while star markers indicate the baseline SAEs. We observe a similar trend as in reconstruction and downstream evaluations: detection and fuzzing scores improve more rapidly as $\text{AMAD}(G)$ decreases—provided it remains above the turning point—after which the scores plateau at an approximately constant level. This result further validates our selection of \hat{G} based on AMAD , suggesting that the interpretability of features in the baseline and Group-SAEs differs only marginally.

5 Conclusion

This work introduces a novel approach to efficiently train SAEs for LLMs by clustering layers based on their angular distance and training a single SAE for each group. Through this method, we achieved up to a 50% reduction in training costs without compromising reconstruction quality or performance on downstream tasks. The results demonstrate that

activations from adjacent layers in LLMs share common features, enabling effective reconstruction with fewer SAEs.

Our findings also show that the SAEs trained on grouped layers perform comparably to layer-specific SAEs in terms of reconstruction and downstream metrics. Furthermore, the automated interpretability evaluations confirmed the interpretability of the features learned by our SAEs, underscoring their utility in disentangling neural activations.

The methodology proposed in this paper opens avenues for more scalable interpretability tools, facilitating deeper analysis of LLMs as they grow in size. Future work will focus on further optimizing the number of layer groups and scaling the approach to even larger models.

Limitations

Although we evaluated our approach across various groups and model sizes, our primary focus here is on experiments using a fixed expansion factor of $c = 16$ and TopK as activation function. Even if we don't expect the choices of these hyper-parameters to influence the results of this work, we left investigations of this phenomenon for future work.

We also limit the scope of our study to models from the Pythia family trained on the Pile dataset. While using the pre-training dataset for SAE training is standard practice (Bricken et al., 2023; Gao et al., 2024), evaluating on additional datasets could provide stronger evidence of generality. We leave such cross-dataset evaluation to future work. Furthermore, architectural and training differences across model families may influence the behavior of Group-SAEs. We defer a comprehensive cross-model analysis to future research. Exploring the generality of our findings across diverse architectures, such as Gemma, LLaMA, Qwen, or Mistral, is an important next step. Finally, our interpretability evaluation remains limited, primarily due to the high economic cost of annotating large numbers of features. While we observe promising patterns, a more comprehensive and systematic interpretability analysis is left for future work.

Reproducibility statement

To support the replication of our empirical findings on training SAEs via layer groups and to enable further research on understanding their inner works, we release the code and SAEs used in this study.⁷

⁷<https://github.com/ghidav/group-sae>

Acknowledgements

The work has received funding from the European Union's Horizon Europe research and innovation programme under grant agreements No. 101189771 (DataPACT) and No. 101070284 (enRichMyData), and the Italian PRIN project Discount Quality for Responsible Data Science (202248FWFS). Additionally, we acknowledge and thank Nscale for providing the compute resources (8 AMD Mi250x GPUs) used for all SAE training and most evaluations in this paper. We are especially grateful to Karl Havard for leading this partnership, Konstantinos Mouzakitis for his technical assistance, Brian Dervan for structuring our collaboration, and the entire Nscale team for their support.

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A Hyperparameters

We train both SAEs and Group-SAEs using Top- K activation⁸ with $K = 128$ and expansion factor of $c = 16$ on the residual stream after the MLP contribution of three models of varying sizes from the Pythia family (Biderman et al., 2023): Pythia 160M, Pythia 410M, and Pythia 1B. To train all the SAEs, we sample 1 billion tokens from the Pile dataset (Gao et al., 2020) and process them with a context size of 1024. We use Adam optimizer (Kingma and Ba, 2017) with default β parameters and set the learning rate equal to $2e-4/\sqrt{(m/2^{14})}$ as specified in Gao et al. (2024). We use a batch size of 131072, 65536 and 32768 for the three models, respectively, to maximize computational usage. Following (Bricken et al., 2023) we constrain the decoder columns (i.e. the feature directions) to have unit norm. Additionally, we normalize the activations to have mean squared ℓ_2 norm of 1 during SAE training, as specified in (Rajamanoharan et al., 2024), by first estimating the norm scaling factor over 5 million tokens of our train set.

Table 3: Pythia model details.

Pythia model	Non-Embedding Params	Layers	Model Dim	Heads
160M	85,056,000	12	768	12
410M	302,311,424	24	1024	16
1.0B	805,736,448	16	2048	8

Table 4: Training and fine-tuning hyperparameters

Hyperparameter	Value
c	16
Top- K K	128
α_{aux}	1/32
Hook name	resid-post
Batch size	131'072 (Pythia-160M) 65'536 (Pythia-410M) 32'768 (Pythia-1B)
Adam (β_1, β_2)	(0.9, 0.999)
Context size	1024
lr	$2e-4/\sqrt{(m/2^{14})}$
lr scheduler	constant
Dead latents threshold	10M
# tokens (Train)	1B
Checkpoint freq	100K
Decoder column normalization	Yes
Activation normalization	Mean squared ℓ_2 norm equal to 1 during SAE training
FP precision	32
Prepend BOS token	No

The experiments were carried out on a cluster of 8 AMD MI250X. The longest experimental run took approximately 24 hours. Our experiments were carried out using PyTorch (Paszke et al., 2019) and the sparsify library.⁹ We performed our data analysis using NumPy (Harris et al., 2020) and Pandas (Wes McKinney, 2010). Our figures were made using Matplotlib (Hunter, 2007) and Seaborn (Waskom, 2021).

⁸The Top- K activation function is directly applied on the features obtained with Equation 1, where $\sigma = \text{Top-}K \circ \text{ReLU}$.

⁹<https://github.com/EleutherAI/sparsify>

B SAEs Training Details

Following [Lawson et al. \(2024\)](#), given $\mathbf{X}, \widehat{\mathbf{X}} \in \mathbb{R}^{B \times n}$ being the input activation batch and its SAE reconstruction, respectively, we train our SAEs with the following loss:

$$\mathcal{L}(\mathbf{X}) = \text{FVU}(\mathbf{X}, \widehat{\mathbf{X}}) + \alpha_{\text{aux}} \cdot \text{AuxK}(\mathbf{X}, \widehat{\mathbf{X}}) \quad (8)$$

The first term of the loss is the Fraction of Variance Unexplained, or:

$$\text{FVU}(\mathbf{X}, \widehat{\mathbf{X}}) = \frac{\|\mathbf{X} - \widehat{\mathbf{X}}\|_F}{\|\mathbf{X} - \overline{\mathbf{X}}\|_F} \quad (9)$$

where $\|\cdot\|_F$ is the Frobenius norm and $\overline{\mathbf{X}} = \frac{1}{B} \mathbf{1}_B \mathbf{1}_B^\top \mathbf{X}$ is a matrix where each row corresponds to the mean of \mathbf{X} along the batch dimension. The second term of the loss is an auxiliary loss to prevent the formation of dead latents during training and is defined as:

$$\text{AuxK}(\mathbf{X}, \widehat{\mathbf{X}}) = \frac{\|\mathbf{E} - \widehat{\mathbf{E}}\|_F}{\|\mathbf{X} - \overline{\mathbf{X}}\|_F} \quad (10)$$

Here, $\mathbf{E} = \mathbf{X} - \widehat{\mathbf{X}}$ is the reconstruction error of the main model, and $\widehat{\mathbf{E}}$ is its reconstruction using the top- K_{aux} dead latents. A dead latent $\mathbf{f}_i(\mathbf{x})$ is a latent that didn't fire, i.e. $\mathbf{f}_i(\mathbf{x}) = 0$, for a predefined number of tokens (10M in our experiments). Following [Gao et al. \(2024\)](#), we choose K_{aux} as the minimum between the number of dead latents and $m/2$, and $\alpha = 1/32$.

To ensure a fair comparison with baselines, we allocate 1 billion training tokens for each SAE_l and SAE_g^G . For baseline SAEs, activations are always taken from a single fixed layer. In contrast, for Group-SAEs, activations are drawn from a randomly selected layer within the set $[g_G]$. In this way, we ensure that each Group- and baseline SAEs process exactly 1 billion tokens and activations.

C Angular Distances and Layers Groups

We use the same angular distance formulation of Gromov et al. (2024):

$$d_\theta(\mathbf{x}^i, \mathbf{x}^j) = \frac{1}{\pi} \arccos \left(\frac{\mathbf{x}^i \cdot \mathbf{x}^j}{\|\mathbf{x}^i\|_2 \|\mathbf{x}^j\|_2} \right) \quad (11)$$

for every $i, j \in \{1, \dots, L\}$, where \mathbf{x}^l are the l -th residual stream activations after the MLP's contribution.

Figures 5–7 visualize the *pairwise* average angular distances between residual-stream activations across all layers for each Pythia model (computed on 10M training tokens). Values are scaled to [0, 1] (0 = identical directions, 0.5 = orthogonal, 1 = opposite), with block structure revealing contiguous regions of high similarity that motivate layer grouping. Tables 5–7 then report the grouping solutions as we vary the number of groups G (up to $L - 1$): the *Groups* row lists, for each layer index, the assigned group ID, and the accompanying *AMAD* value (Average Maximum Angular Distance, Eq. 4) summarizes within-group compactness. As G increases, AMAD typically decreases, reflecting finer partitions that better capture the block-diagonal structure observed in the distance matrices.

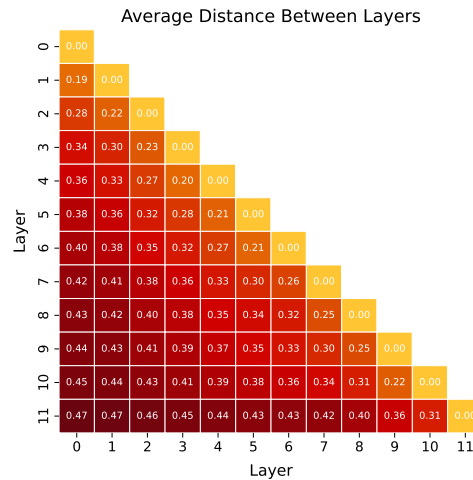


Figure 5: Average angular distance between all layers of the Pythia-160M model, as defined in Equation 11. The angular distances are computed over 10M tokens from the training dataset. The angular distances are bounded in [0, 1], where an angular distance equal to 0 means equal activations, 0.5 means activations are perpendicular and an angular distance of 1 means that the activations point in opposite directions.

G	Groups	AMAD
1	0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0	0.450
2	0, 0, 0, 0, 0, 0, 1, 1, 1, 1	0.372
3	2, 2, 2, 1, 1, 1, 0, 0, 0, 0	0.314
4	2, 2, 2, 0, 0, 0, 1, 1, 3, 3	0.267
5	0, 0, 0, 4, 2, 2, 1, 1, 3, 3	0.231
6	3, 3, 5, 4, 4, 2, 2, 0, 0, 1, 1	0.179
7	3, 3, 5, 1, 1, 2, 2, 6, 4, 0, 0	0.118
8	3, 3, 5, 1, 1, 0, 0, 6, 4, 7, 2	0.075
9	1, 1, 5, 0, 0, 8, 7, 6, 4, 3, 2	0.044
10	0, 0, 5, 9, 7, 8, 3, 6, 4, 1, 2	0.019

Table 5: Layer groups for every G up to $L - 1$ for Pythia-160M

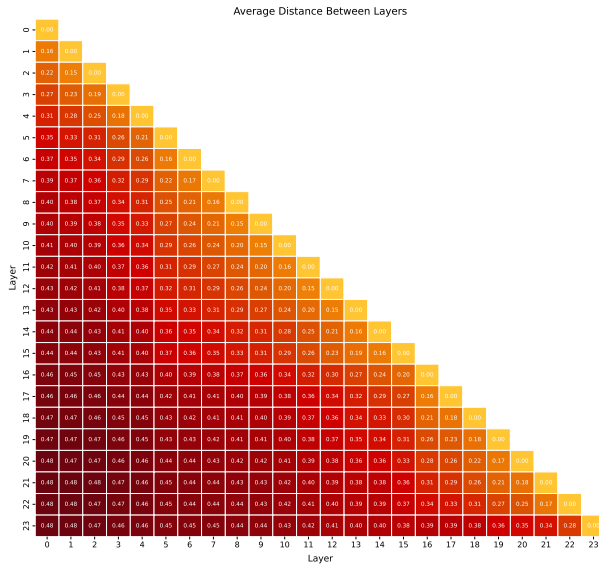


Figure 6: Average angular distance between all layers of the Pythia-410M model, as defined in Equation 11. The angular distances are computed over 10M tokens from the training dataset. The angular distances are bounded in $[0, 1]$, where an angular distance equal to 0 means equal activations, 0.5 means activations are perpendicular and an angular distance of 1 means that the activations point in opposite directions.

Table 6: Layer groups for every G up to $L - 1$ for Pythia-410M

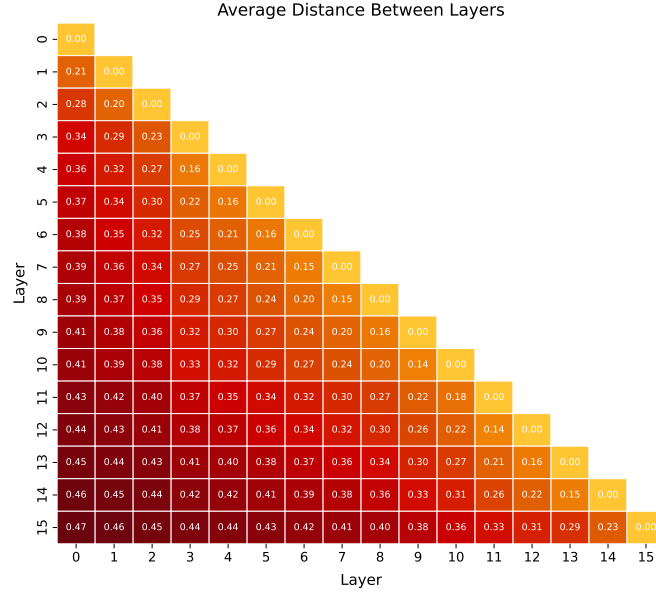


Figure 7: Average angular distance between all layers of the Pythia-1B model, as defined in Equation 11. The angular distances are computed over 10M tokens from the training dataset and are bounded in $[0, 1]$. An angular distance equal to 0 means equal activations, 0.5 means activations are perpendicular and an angular distance of 1 means that the activations point in opposite directions.

G	Groups	AMAD
1	0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0	0.459
2	0, 0, 0, 0, 0, 0, 0, 0, 1, 1, 1, 1, 1, 1	0.364
3	1, 1, 1, 1, 2, 2, 2, 2, 0, 0, 0, 0, 0, 0	0.309
4	0, 0, 0, 0, 2, 2, 2, 2, 1, 1, 1, 3, 3	0.250
5	4, 4, 1, 1, 1, 2, 2, 2, 2, 0, 0, 0, 3, 3	0.225
6	4, 4, 1, 1, 1, 0, 0, 0, 0, 2, 2, 5, 5, 3, 3	0.191
7	1, 1, 0, 0, 4, 4, 2, 2, 6, 6, 5, 5, 3, 3	0.174
8	0, 0, 7, 3, 3, 4, 4, 2, 2, 6, 6, 5, 5, 1, 1	0.139
9	8, 4, 7, 3, 3, 1, 1, 2, 2, 6, 6, 5, 5, 0, 0	0.100
10	8, 9, 7, 1, 1, 0, 0, 2, 2, 6, 6, 5, 5, 4, 3	0.075
11	8, 9, 7, 0, 0, 10, 3, 2, 2, 6, 6, 5, 5, 4, 1	0.053
12	8, 9, 7, 11, 6, 10, 3, 0, 0, 2, 2, 5, 5, 4, 1	0.036
13	8, 9, 7, 11, 6, 10, 3, 12, 5, 0, 0, 2, 2, 4, 1	0.022
14	8, 9, 7, 11, 13, 10, 3, 12, 5, 6, 2, 0, 0, 4, 1	0.010

Table 7: Layer groups for every G up to $L - 1$ for Pythia-1b

D Auto Interpretability

To evaluate the interpretability of features of baseline and Group SAEs, we adopt automated pipeline from (Paulo et al., 2024b), focusing on *detection* and *fuzzing* scores. First, an *explainer* language model (LM) generates natural language explanations of the SAE latent representations. Then, a separate *scorer* LM evaluates these explanations.

Then, detection scoring assesses whether a language model can identify entire sequences that activate a specific latent, given its interpretation. This method evaluates the model’s ability to distinguish between activating and non-activating contexts, offering insights into the precision and recall of the interpretation. Fuzzing scoring, on the other hand, operates at the token level, prompting the model to pinpoint specific tokens within sequences that trigger latent activations. This approach closely mirrors simulation scoring and is particularly effective in evaluating the model’s token-level understanding of latent activations.

In our experiments, we use `gemini-2.0-flash-001` as the base model for both the explainer and the scorer. For each SAE, we randomly select 64 features and cache their latent activations across 10M tokens from the Pile (Gao et al., 2020). To generate annotations, we present the explainer with 20 distinct examples per feature—10 that activate the latent and 10 randomly sampled—each comprising 32 tokens.

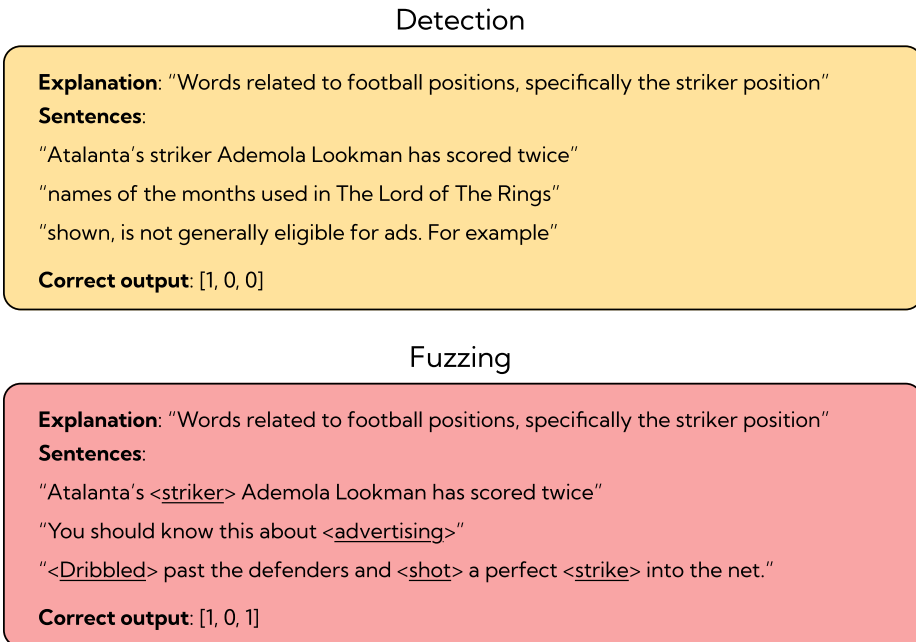


Figure 8: Examples of each of the auto-interpretability techniques: Detection and Fuzzing. In detection, the objective is to find the sentences in which the feature is active. In fuzzing, the objective is to spot the highlighted tokens referring to the target feature.

E Feature Similarity Analysis

Following Sharkey et al. (2023), we adopt Mean Maximum Cosine Similarity (MMCS) to assess the extent to which Baseline and Group SAEs learn similar feature directions. For any two SAEs, SAE_i and SAE_j , we compute the MMCS between their decoder matrices $\mathbf{W}_d^i, \mathbf{W}_d^j \in \mathbb{R}^{n \times m}$ as these matrices encode the directions of the learned features:

$$\text{MMCS}(\mathbf{W}_d^i, \mathbf{W}_d^j) = \frac{1}{m} \sum_{k=1}^m \max_{l \in \{1, \dots, m\}} \cos(\tilde{\mathbf{w}}_k^i, \tilde{\mathbf{w}}_l^j) \quad (12)$$

where $\tilde{\mathbf{w}}_k^i$ and $\tilde{\mathbf{w}}_l^j$ are the k -th and l -th columns of the normalized decoder matrices $\tilde{\mathbf{W}}_d^i$ and $\tilde{\mathbf{W}}_d^j$, respectively. The directionality of the maximum operation is important for interpretation: we first find, for each feature in SAE_i , the most similar feature in SAE_j (by cosine similarity), and then average these maximum similarities across all features of SAE_i . In our analysis, we specifically compute $\text{MMCS}(\mathbf{W}_d^{\text{Baseline}}, \mathbf{W}_d^{\text{Group}})$, meaning that the resulting value represents the average highest similarity that each Baseline SAE feature has with any feature in the Group SAE.

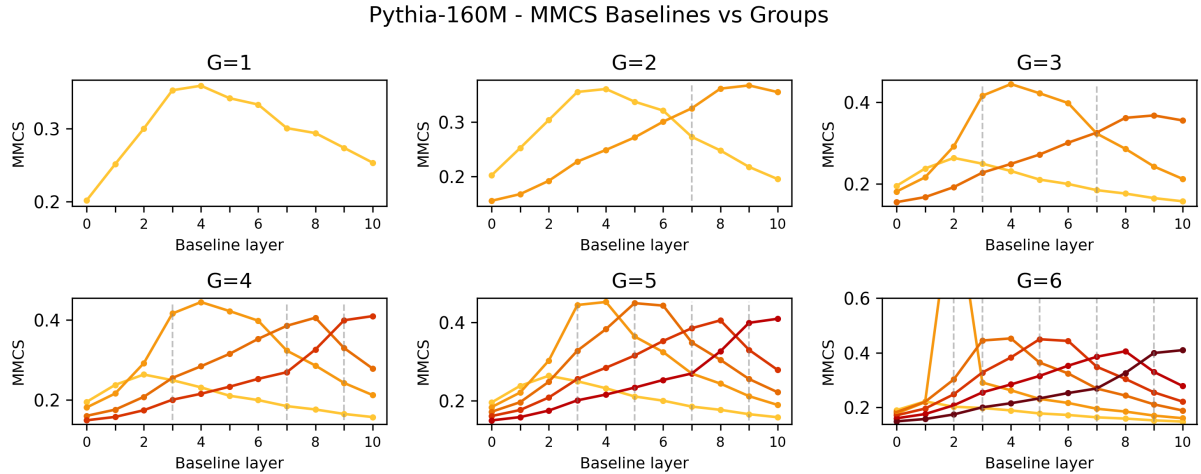


Figure 9: Mean Maximum Cosine Similarity (MMCS) between all the learned features of baseline and group SAEs for each group $G \in \{1, \dots, \hat{G}\}$ of Pythia-160M. Colors represent the different Group SAEs of a given partition.

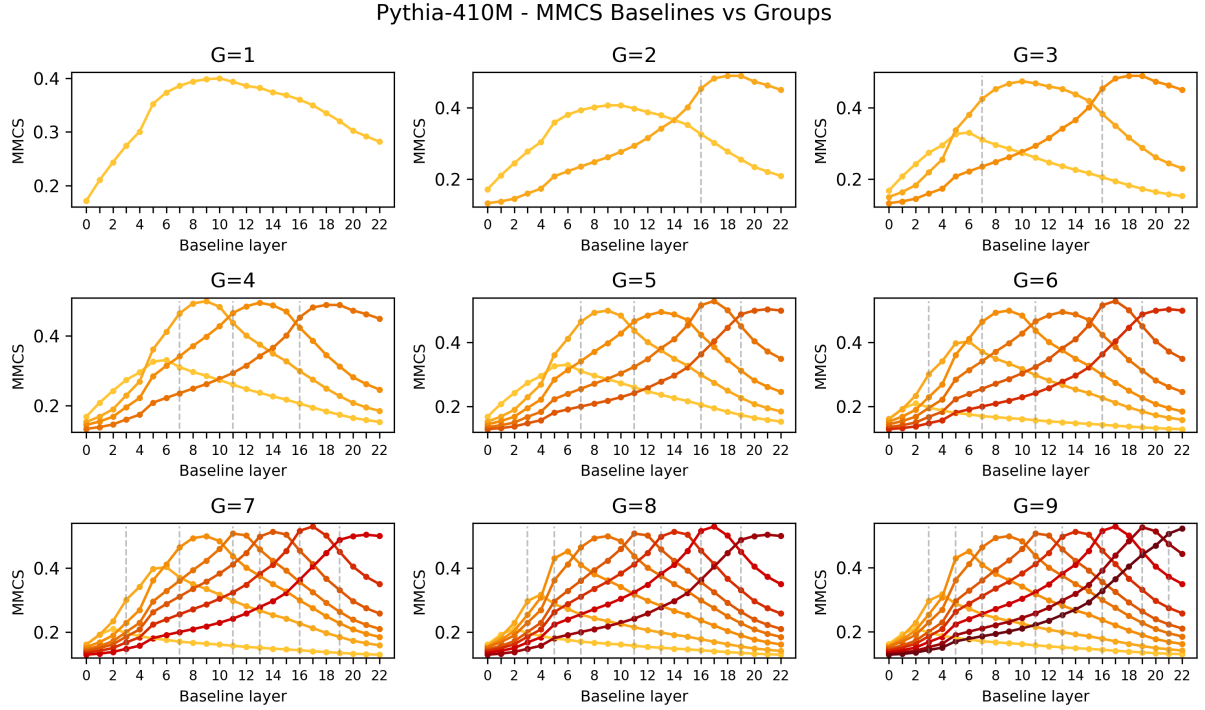


Figure 10: Mean Maximum Cosine Similarity (MMCS) between all the learned features of baseline and Group SAEs for each group $G \in \{1, \dots, \hat{G}\}$ of Pythia-410M. Colors represent the different Group SAEs of a given partition.

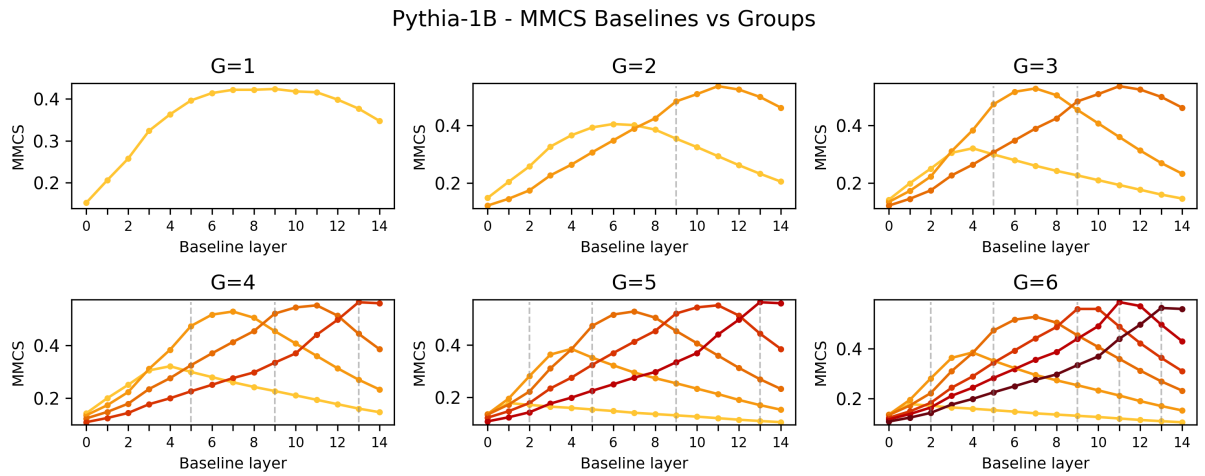


Figure 11: Mean Maximum Cosine Similarity (MMCS) between all the learned features of baseline and Group SAEs for each group $G \in \{1, \dots, \hat{G}\}$ of Pythia-1B. Colors represent the different Group SAEs of a given partition.

F Feature Distribution Analysis

Following (Lawson et al., 2024), we perform a study to understand how features distribute across layers of a given group. Previous work from (Lindsey et al., 2024) showed that activations of a given feature usually peak at a specific layer. To measure this phenomenon, for each Group SAE of a given partition in G groups, we sample 1 million tokens from the test set and compute feature distributions across the layers of its group.

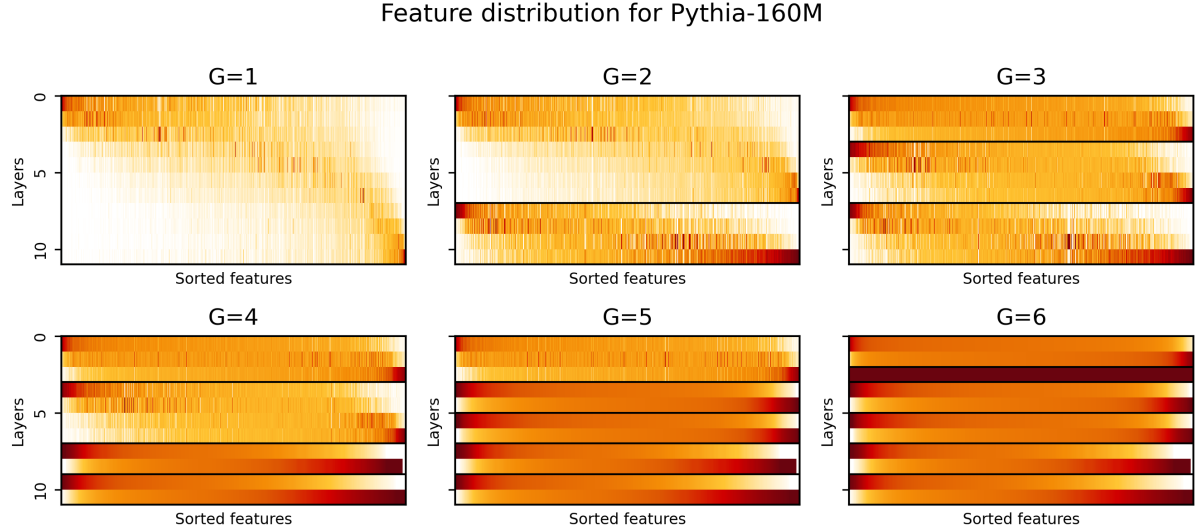


Figure 12: Pythia-160M feature activations distribution for every group $G \in \{1, \dots, \hat{G}\}$ over 1 million tokens from the test set. Darker regions indicate higher feature activation density.

Heatmaps in Figure 12, 13, and 14 show distributions of features activations for all the models and Group SAEs of partitions from 1 to \hat{G} . In the images, we sort the features by the average layer they activate the most. Darker regions indicate higher feature activation density. Looking at the charts several considerations can be drawn:

- Features activating for the first and last layers of a given group tend to be more specific for that layers (i.e. their activation frequencies peak at those layers).
- Features at early layers of a model are more spread across their respective group.
- Bigger models tend to have features more spread across the layers of a given group with respect to smaller models.

In summary, while feature distributions tend to peak at a specific layer (with this being more evident in smaller models and later layers), they also spread across close ones. This result agrees with findings from (Lindsey et al., 2024) while still leaving the potential for Group SAEs to make SAE training more efficient.

Feature distribution for Pythia-410M

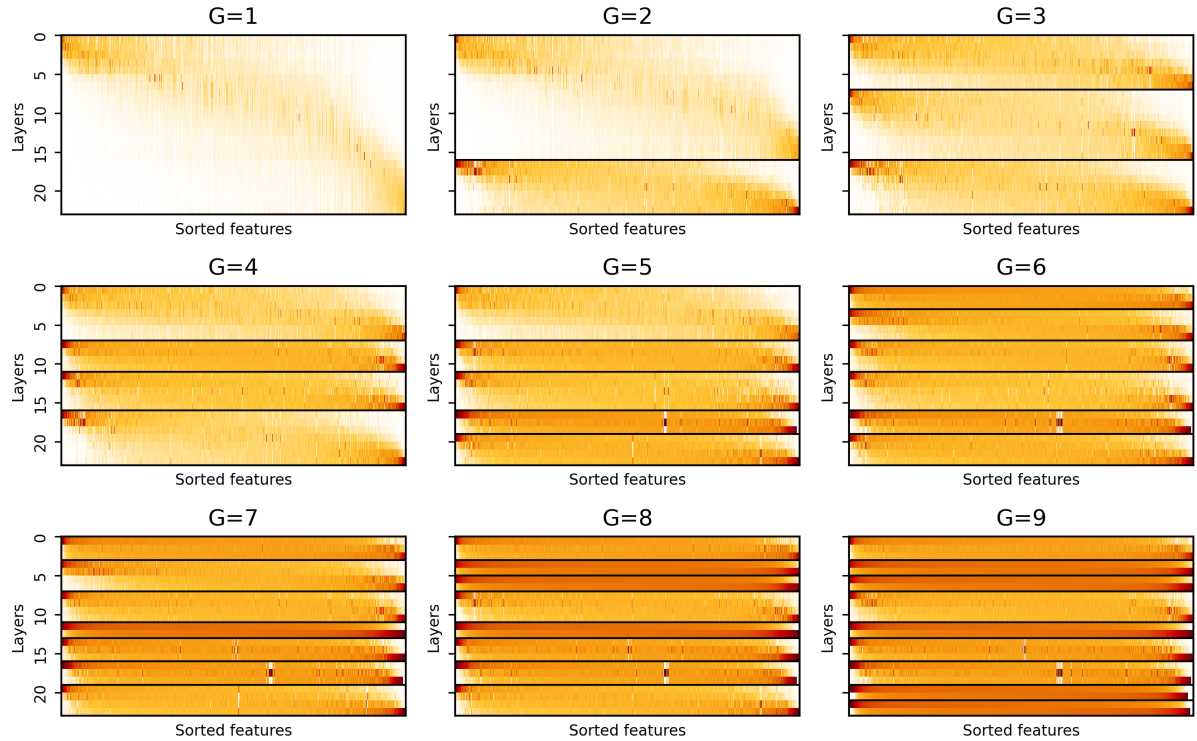


Figure 13: Pythia-410M feature activations distribution for every group $G \in \{1, \dots, \hat{G}\}$ over 1 million tokens from the test set. Darker regions indicate higher feature activation density.

Feature distribution for Pythia-1B

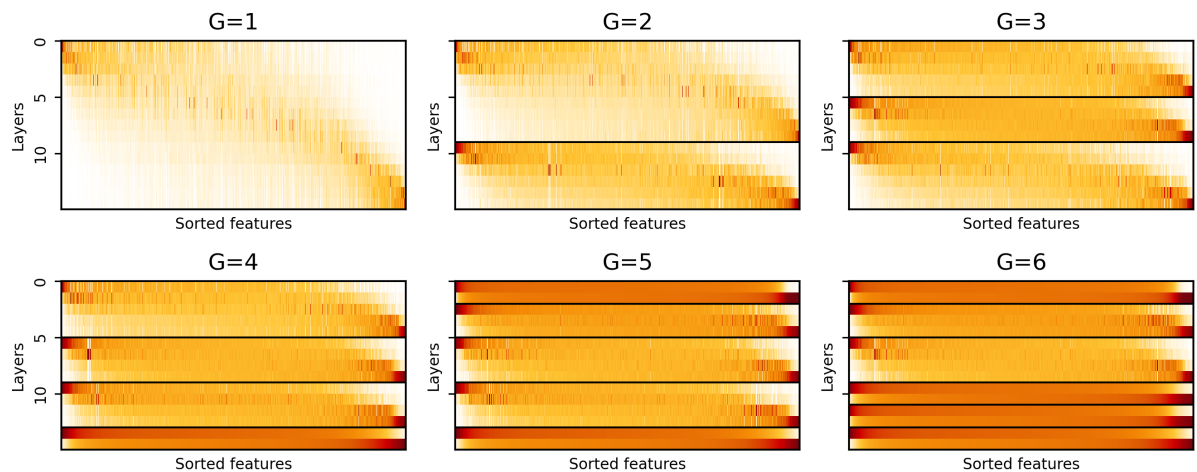


Figure 14: Pythia-1b feature activations distribution for every group $G \in \{1, \dots, \hat{G}\}$ over 1 million tokens from the test set. Darker regions indicate higher feature activation density.