JOINT MOE SCALING LAWS: MIXTURE OF EXPERTS CAN BE MEMORY EFFICIENT

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ABSTRACT

Mixture of Experts (MoE) architectures have significantly increased computational efficiency in both research and real-world applications of large-scale machine learning models. However, their scalability and efficiency under memory constraints remain relatively underexplored. In this work, we present joint scaling laws for dense and MoE models, incorporating key factors such as the number of active parameters, dataset size, and the number of experts. Our findings provide a principled framework for selecting the optimal MoE configuration under fixed memory and compute budgets. Surprisingly, we show that MoE models can be more memory-efficient than dense models, contradicting conventional wisdom. Extensive empirical validation confirms the theoretical predictions of our scaling laws. These results offer actionable insights for designing and deploying MoE models in practical large-scale training scenarios.

1 INTRODUCTION

Recently, language models have grown increasingly large, a trend accelerated by Mixture of Experts (MoE) techniques (Fedus et al., 2022; Du et al., 2022). MoE models are now widely adopted (Jiang et al., 2024; Dai et al., 2024) and are generally considered compute-efficient (Ludziejewski et al., 2024; Clark et al., 2022), though often considered to be memory-inefficient Zadouri et al. (2023). However, the precise trade-offs between compute and memory efficiency remain unclear. Consider a motivating question: Is an MoE model the optimal choice when constrained by a fixed memory budget, such as a single H100 node? While computational efficiency is important, it does not directly determine the optimal number of experts. Increasing the number of experts has minimal impact on computation but can drastically raise memory requirements, often to a prohibitive level. To address this question, we derive a *joint* scaling law for both dense and MoE models, accounting for key factors such as the number of active parameters, dataset size, and number of experts. This framework provides a rigorous analysis of model performance under strict memory constraints. Our findings reveal that, contrary to common assumptions, MoE models can be more memory-efficient than dense models. Our work is the first to provide detailed guidance on selecting the optimal number of experts for MoE models, balancing both computational and memory constraints. Our conclusions are based on extensive large-scale experiments with over 280 models, scaled up to 5B parameters. In summary, the key contributions of this work are:

- We derive a joint scaling law for Mixture of Experts and dense models, $\mathcal{L}(N_{\text{act}}, D, \hat{E}) = \hat{E}^{\delta} N_{\text{act}}^{\alpha+\gamma \ln(\hat{E})} + b \hat{E}^{\omega} D^{\beta+\zeta \ln(\hat{E})} + c$ where \mathcal{L} is the final training loss, N_{act} is the number of active parameters, D is the dataset size, \hat{E} is the monotonic transformation of the number of experts, and c is the irreducible entropy of the dataset.
- Based on the proposed scaling law, we show that the choice of the optimal number of experts (including dense models with E = 1) depends on specific computational and memory constraints, see Figure 1. Moreover, we demonstrate how the optimal token-to-parameter ratio depends on E.
- We show that MoE can often be the preferred alternative to dense models, even if GPU memory is the constraining factor. We validate our theoretical findings by training a set of 1.1B-parameter models under identical compute and total memory budgets. The MoE models achieve a lower final loss, confirming their superior efficiency in practice.

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Figure 1: (a) The loss of memory-constrained models predicted using our scaling law under a fixed training budget of 10^{22} FLOPs. Each curve represents a different number of experts. Shaded areas present memory optimal number of experts for the corresponding parameter budgets. (b) Experimental validation of the thesis that MoE can be memory optimal. The marked area shows an interval in which a training compute-matched MoE achieves better loss than an overtrained dense model with the same number of total parameters (1.1B). The resulting MoE was trained for longer and had less active parameters, making it more practical.

2 JOINT MOE SCALING LAWS

We now derive the functional form of our joint scaling laws for both dense Transformers and MoE, relating the number of active model parameters N_{act} , training tokens D, and MoE experts E. We propose the form of our scaling law:

$$\mathcal{L}(N_{\text{act}}, D, \hat{E}) = a\hat{E}^{\delta}N_{\text{act}}^{\alpha+\gamma\ln(E)} + b\hat{E}^{\omega}D^{\beta+\zeta\ln(\hat{E})} + c.$$
(1)

We derive the formula based on the following observations. Assuming that if we fix the number of experts the model performance can be described using Equation 4 Hoffmann et al. (2022). Scaling in E can be described as a power law (Clark et al., 2022). Moreover, for a fixed dataset size, as model size increases, the benefit of using an MoE diminishes (Clark et al., 2022). On the other hand for a fixed model size, as the number of training tokens increases, the benefit of an MoE grows (Ludziejewski et al., 2024). To ensure flexibility in modeling these observations, we introduce an interaction with the exponents over N_{act} and D: $\mu(E) = \alpha + \gamma \ln(E), \nu(E) = \beta + \zeta \ln(E)$. See Sec. A in Appendix for more details. Empirically, we observe a good fit for our formula, as described in Section D.

3 COMPUTE AND MEMORY OPTIMALITY

In this section, we employ our scaling laws to derive recommendations on optimal settings in various training and inference scenarios.

Compute Optimality. A model is considered compute-optimal if, among models trained with the same compute budget F, it achieves the lowest loss. To find such an optimal configuration, we optimize the following: $\arg \min_{N_{act}, D, E} \mathcal{L}(N_{act}, D, E)$ s.t. $6N_{act}D = F$

Optimal N and D Depend on the Number of Experts. Assuming a given number of experts E, the compute-optimal training configuration can be achieved by selecting the appropriate trade-off between training tokens and model size. IsoFLOP slices comparing the predicted loss with dataset size for selected compute budgets are plotted in Figure 2(b).

For any fixed E our scaling law has the Chinchilla functional form of Equation 4. Thus, from Hoffmann et al. (2022), the compute-optimal number of tokens and active parameters for the budget F and

the number of experts E are given by $N_{\text{act}}^{\text{opt}}(F) = G\left(\frac{F}{6}\right)^a$, $D^{\text{opt}}(F) = G^{-1}\left(\frac{F}{6}\right)^b$, where $G = \left(\frac{\mu(E)m(E)}{\nu(E)n(E)}\right)^{\overline{\mu(E)}+\nu(E)}$ and $a = \frac{\nu(E)}{\mu(E)+\nu(E)}$, $b = \frac{\mu(E)}{\mu(E)+\nu(E)}$. We compare the optimal configurations for several compute budgets in Table 1.



Figure 2: (a): IsoFLOP profiles for selected training budgets. Compute-optimal points are marked. (b): Savings from switching from a compute-optimal dense model to MoE with the same total parameter count. (c): Compute-optimal training configurations for MoE models with 1×10^{21} training budget. As the number of experts increases, the optimal D^{opt} goes up, and $N_{\text{act}}^{\text{opt}}$ decreases.



Figure 3: Loss predicted for various expansion rates at a FLOPs budget $F = 5 \times 10^{22}$. The x-axis denotes the size of the corresponding dense model, possibly with KV cache. (a) The model size is simply the number of parameters. (b) The model size includes the KV cache (c) Additionally to KV cache, the training budget is reduced by the inference cost on 100B tokens.

Both from comparing the IsoFLOP slices and the values listed in the table from the joint scaling law, we can see that the compute-optimal configuration for a given compute budget clearly depends on E, with MoE models requiring comparatively larger datasets and correspondingly smaller numbers of active parameters.

Finding 1. More experts \rightarrow higher tokens-to-param ratio.

Assume a fixed compute budget. In this scenario, when increasing the number of experts, it is optimal to decrease the number of active parameters and increase the number of training tokens accordingly (Table 1).

Mixture of Experts is Compute Optimal. Now, we compare the performance across various numbers of experts, with respective values of tokens and active parameters optimized. As illustrated in Figure 2, we observe significant compute savings for MoE models compared to dense models, with a larger number of experts providing more pronounced benefits.

Finding 2. More experts \rightarrow better performance. For a given compute budget, increasing the number of experts always improves performance, provided the size of the model and the number of training tokens are adjusted (Figure 2b).

The higher efficiency of MoE in terms of training compute comes at a price of increased memory requirements. However, somewhat surprisingly, we find that MoE models can outperform dense models *of the same size* trained with the same amount of training compute.

Model Memory Optimality. Compute optimality alone is often insufficient, as a compute-optimal model may be too large for deployment or inefficient with small GPU batch sizes (He, 2022). A natural extension is model memory optimality, where a model is memory optimal if, among those trained with the same compute budget F and at most M parameters, it achieves the lowest loss: $\arg\min_{N_{act},D,E} \mathcal{L}(N_{act},D,E)$ s.t. $6N_{act}D = F$, $N_{total} \leq M$. Note that model memory-matched dense and MoE models differ in the number of active parameters—MoE uses just a fraction of them. Intuitively, it should thus have worse performance. At the same time, given some budget, it can be

trained on more tokens, lowering the loss. Our scaling laws suggest that MoE models can be model memory efficient. We validate this claim by training a 1.1B dense model and a model size and FLOP matched $E = \{2, 4\}$ counterparts (Figure 1). Significantly, the MoE models attains lower loss even if the dense model is overtrained (i.e., after passing its compute-optimal token count).

Finding 3. MoE can also be *memory*-efficient.

A total-parameter-matched MoE model can outperform a dense model trained with the same compute budget (Figure 1). Moreover, such an MoE model is more compute- *and* memory-efficient at inference.

Total Memory Optimality. During autoregressive generation, a decoder-only model processes a single token while storing activations (keys and values) for previous tokens in the KV cache. which yields the optimization criterion: $\arg\min_{N_{act},D,E} \mathcal{L}(N_{act},D,E)$ s.t. $6N_{act}D = F$, $N_{total} + 2TN_{blocks}d_{model} \leq M$. where T is the number of tokens in the cache (possibly within multiple sequences in the batch). Figure 1 (b) presents the optimal models for a given compute and varying memory constraints when the size of the KV cache is included. Importantly, MoE models compare more favorably to dense models in this graph, and as T increases, they outperform dense models at even smaller model sizes.

Inference Optimality. Large models, while capable, might also be too costly to run due to their high computational demand. To account for this drawback, we can further assume that a model will process some number of tokens, D_{inf} , throughout its lifetime and find the best model whose demands do not exceed some predefined joint training and inference budget: $\arg \min_{N_{act},D,E} \mathcal{L}(N_{act},D,E)$ s.t. $6N_{act}D+2N_{act}D_{inf} = F$. We find that in this scenario, MoE models outperform dense at smaller scales than in simple compute-optimality due to decreased inference FLOPs (see Fig, Figure 3 (c) in Appendix).

The notions of inference optimality and total memory optimality can naturally be combined. For practitioners, as a simplification of our analysis, we propose a general rule of thumb:

Rule of Thumb. An MoE model with $E \le 8$ experts, trained on *E*-times more tokens than a compute-optimal dense model, outperforms it while maintaining the same total parameter count.

Note that, in this scenario FLOPs matched MoE will generally have less than *E*-times larger dataset, but we wanted to keep this rule simple and conservative. Detailed comparisons and differences between memory and FLOPs matched models can be found on Figures 1 & 5.



4 CONCLUSION

In this work, we derived the joint scaling laws for Mixture of Experts, relating the loss of the model to the number of parameters, the number of training tokens, and the number of experts. By considering both compute and memory constraints, as well as the expected inference workload, we demonstrated that MoE models can outperform dense models even when constrained by memory usage or total parameters, contrary to common assumptions and intuitions that MoE models are more memory-intensive than dense models. Our analysis reveals how the optimal training strategies shift as the number of experts varies. This provides a principled framework for selecting MoE hyperparameters under given constraints, highlighting the trade-offs between memory and compute performance.

ACKNOWLEDGMENTS

We would like to express sincere gratitude to Szymon Antoniak and Piotr Padlewski for their detailed comments and invaluable discussions. We also thank Konrad Staniszewski for his feedback on the draft of this paper.

We gratefully acknowledge the Polish high-performance computing infrastructure PLGrid (HPC Center: ACK Cyfronet AGH) for providing computer facilities and support within computational grant no. PLG/2024/017060. This research was partially supported by the ERC PoC Grant EXALT no. 101082299, the National Science Centre (NCN) Grant no. 2020/37/B/ST6/04179, the National Science Centre (NCN) Preludium Grant no. 2022/45/N/ST6/02222, the "European Lighthouse of AI for Sustainability" - ELIAS grant no. 101120237, and the NCBiR grant POIR.01.01.01-00-0433/20. Part of the experiments utilized computational resources provided by Writer.

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A DERIVATION OF JOINT MOE SCALING LAW

We now derive the functional form of our joint scaling laws for both dense Transformers and MoE, relating the number of active model parameters N_{act} , training tokens D, and MoE experts E.

Fixed Number of Experts. Following Hoffmann et al. (2022) and established practice in the literature (Frantar et al., 2023; Kumar et al., 2024; Ludziejewski et al., 2024), we postulate the following form of the equation:

$$\mathcal{L}(N_{\rm act}, D, E) = m(E)N_{\rm act}^{\mu(E)} + n(E)D^{\nu(E)} + c(E),$$
(2)

assuming that if we fix the number of experts the model performance can be described using Equation 4. In the subsequent part, we will postulate how m, μ, n, ν, c depend on E, deriving the joint equation.

Constant Factor. c(E) represents irreducible loss caused by the inherent entropy of the dataset. Thus, it does not depend on the architecture (*E* in our case): c(E) := c.

Interaction of E with Model and Dataset Size. To quantify the interaction between the number of experts and other training parameters, we gather observations from related work:

- 1. Scaling in E can be described as a power law (Clark et al., 2022).
- 2. For a fixed dataset size, as model size increases, the benefit of using an MoE diminishes (Clark et al., 2022).
- 3. For a fixed model size, as the number of training tokens increases, the benefit of an MoE grows (Ludziejewski et al., 2024).

Motivated by Observation 1, we set $m(E) = aE^{\delta}$, $n(E) = bE^{\omega}$, reflecting the power-law relation between E and the loss. Additionally, to ensure flexibility in modeling Observations 2 and 3, we introduce an interaction with the exponents over N_{act} and D: $\mu(E) = \alpha + \gamma \ln(E), \nu(E) = \beta + \zeta \ln(E)$. Note that if we ignore the second and third terms in Equation 2, this yields a functional form identical to Equation 5. Empirically, we observe a good fit for our formula, as described in Section D. This shows that our proposed interactions between E, N_{act} , and D can accurately model the performance of MoE models.

Modeling of E. When the number of experts is small, a certain overhead, caused, for example, by interference from auxiliary losses, can overshadow the benefits of conditional computation. Additionally, using very large numbers of experts brings diminishing returns. To account for these phenomena, we follow Clark et al. (2022) and use a transformation of the number of experts \hat{E} given in Equation 6.

Joint MoE Scaling Law. Combining these observations, we derive the final form of our scaling law:

$$\mathcal{L}(N_{\rm act}, D, \hat{E}) = a\hat{E}^{\delta}N_{\rm act}^{\alpha+\gamma\ln(E)} + b\hat{E}^{\omega}D^{\beta+\zeta\ln(\hat{E})} + c.$$
(3)

We fit the coefficients in Equation 3 based on the results of our experiments; see Table 2. In Section 3, we present the outcomes and findings derived from the scaling laws. The details of the training runs, as well as the fitting procedure, are described in Section D.

B COMPUTE AND MEMORY OPTIMALITY RESULTS

The following table and plot analyze the compute-optimal configurations and performance characteristics of Mixture of Experts (MoE) models under various training budgets and memory constraints. The table presents optimal training configurations, while the subsequent plot illustrates the optimal number of experts for different model sizes. These analyses provide insights into how MoE models can be efficiently scaled while balancing computational and memory constraints.

C RELATED WORK

Mixture of Experts. Mixture of Experts (MoE) was introduced by Jacobs et al. (1991), who combined a gating network with a set of expert networks. Shazeer et al. (2017) applied MoE to an

Training Budget \Rightarrow	1 >	1×10^{20}		5×10^{20}		1×10^{21}	
Experts U	$N_{\rm act}^{\rm opt}$	D^{opt}		$N_{\rm act}^{\rm opt}$	D^{opt}	$N_{\rm act}^{\rm opt}$	D^{opt}
1	1.7 B	9.7 B		$4\mathbf{B}$	21 B	5.7 B	29.3B
2	$1.5\mathbf{B}$	11.4 B		3.5B	24B	5B	33B
4	1.2B	13.9B		3B	28B	4.4B	38B
8	990M	17B		2.5B	33.2B	3.8B	$44.3\mathbf{B}$
16	810M	20.7B		2.1B	39B	3.3B	51.2B

Table 1: Example compute-optimal training configurations for MoE models. For every training budget as the number of experts increases, the optimal D^{opt} also goes up while $N_{\text{act}}^{\text{opt}}$ decreases.



Figure 5: Investigation of the optimal number of experts for three different model sizes, 2B, 5B, and 10B; and in three different scenarios, from left to right: simply measuring the size of the model, including the size of a KV-cache with 32k tokens, and including the inference cost of processing 100B tokens.

LSTM-based model (Hochreiter & Schmidhuber, 1997), scaling the architecture up to 137 billion parameters. In Transformer-based LLMs, MoE is most often applied as a replacement for the feed-forward layer (Lepikhin et al., 2020; Shazeer et al., 2018). It replaces the feed-forward's MLP with a set of expert MLPs along with a router, which selects one or more MLPs for each token. With the recent surge in LLM research, MoE models are gaining even more traction. This is exemplified by the development of extremely large-scale models such as DeepSeek-R1 and Qwen2.5-Max (DeepSeek-AI et al., 2025; Team, 2024a). In our work, we use the standard Switch MoE layer (Fedus et al., 2022), which routes each token to one expert and encourages even token-to-expert assignment via the addition of a differentiable load-balancing loss.

Scaling Laws. Scaling laws refer to empirically derived equations that relate model loss to factors such as the number of parameters, the quantity of training data, or the computational budget. For dense Transformers, scaling laws were initially explored by Hestness et al. (2017) and Kaplan et al. (2020), who identified power-law relationships between the final loss, model size, and dataset size. Hoffmann et al. (2022) expanded this by incorporating variable cosine cycle lengths and adjusting the functional form of the equation:

$$\mathcal{L}(N_{\rm act}, D) = mN_{\rm act}^{\mu} + nD^{\nu} + c.$$
⁽⁴⁾

Scaling laws have also been applied to other architectures and training setups. Henighan et al. (2020) examined autoregressive modeling across multiple modalities, while Ghorbani et al. (2021) focused on machine translation. Frantar et al. (2023) studied the effects of pruning on vision and language Transformers, determining optimal sparsity given a fixed compute budget.

Clark et al. (2022) investigated scaling in MoE models, varying model size and the number of experts on a fixed dataset, and concluded that routed models are more efficient only up to a certain size. Their formula took the form:

$$\mathcal{L}(N_{\rm act}, \hat{E}) = a\hat{E}^{\delta} N_{\rm act}^{\alpha + \gamma \ln(E)},\tag{5}$$

where \hat{E} is a monotonic transformation of the number of experts E defined as:

$$\frac{1}{\hat{E}} = \frac{1}{E - 1 + \left(\frac{1}{E_{\text{start}}} - \frac{1}{E_{\text{max}}}\right)^{-1}} + \frac{1}{E_{\text{max}}}.$$
(6)

These analyses have since been extended by Ludziejewski et al. (2024) and Dai et al. (2024), who considered variable dataset size as well as the granularity of experts. In our work, we keep the experts non-granular; however, we treat the number of experts and the number of training tokens as variables. Sardana et al. (2024) assumes a fixed joint inference and training budget. We make similar assumptions; however, we consider accelerator memory as a limiting factor and extend the analysis to MoE models, which can serve as a more compute-friendly alternative to dense models. Yun et al. (2024) have focused on MoE inference optimality and measuring real hardware efficiency.

D FITTING THE SCALING LAW

In this section, we present details of experiments and procedure of fitting the scaling law parameters, see Table 2 in Appendix. Those results are based on an extensive large-scale empirical evidence, including over 280 models with up to 5B parameters, trained on a variety of compute budgets. For a full list of experiments, see Appendix H.

D.1 MODEL HYPERPARAMETERS

The selection of hyperparameters and training details is crucial for ensuring the robustness of scaling laws (Porian et al., 2025; Pearce & Song, 2024). In our work, we employ a set of best practices and modern design choices, aiming to provide accurate predictions applicable to real-life practice.

All models used in this study are decoder-only Transformers trained on the highly filtered FineWeb-Edu (Penedo et al., 2024). We use a Transformer model with Switch (Fedus et al., 2022) layers, using standard values of router z-loss 0.001 and load balancing loss 0.01. The GPT-2 tokenizer (Radford et al., 2018) is employed. For better stability, weight initialization follows a truncated normal distribution with a reduced scale of 0.1, as suggested by Fedus et al. (2022). Mixed precision training is used, with the attention mechanism, position embeddings RoPE Su et al. (2023) and router always maintained at high precision. The models use the SwiGLU activation (Shazeer, 2020) with hidden size equal to $3d_{model}$ and activate one expert per token (unless the token is dropped due to limited capacity). For evaluation, we increase the capacity factor to ensure dropless processing of the tokens.

Batch Size Ramp-up. Performance of a deep learning optimization procedure can suffer as a result of using an exceedingly large batch size (McCandlish et al., 2018). To mitigate this potential issue, especially early in the training, we employ batch-size ramp-up. Similar strategies are used in contemporary LLM training runs (Rae et al., 2022; Dubey et al., 2024). We increase the batch size from 64K to 128K after 0.5B training tokens and further to 256K after 1B training tokens. Instead of using noise scale as a critical batch size predictor (McCandlish et al., 2018) we opted for a straightforward grid to directly predict a transition point after which increased batch size does not impair performance.

Learning Rate Scaling. Kaplan et al. (2020) have shown that scaling laws for hyperparameters can be used to adjust them according to the size of the model in the case of dense Transformers. For MoE models, we find the literature inconclusive–while some (Dai et al., 2024) pretrain MoEs with lower LR than corresponding dense models, others (Zoph et al., 2022) report better performance when finetuning MoEs with higher learning rates. To fill this gap, we derive a scaling law for the peak learning rate for MoE based on the number of active non-embedding parameters $N_{act \setminus e}$ and the number of experts E:

$$LR(N_{act \setminus e}, E) = \exp(8.39 - 0.81 \ln(N_{act \setminus e}) - 0.25 \ln(E)), \tag{7}$$

and use this equation to set the learning rate in our main scaling laws experiments. We fit the coefficients of this equation using the least squares method, minimizing the error between the prediction and the optimal learning rate from the experiment grid. Contrary to Kaplan et al. (2020), we use a linear transformation of the parameter count to predict the logarithm of the learning rate, instead of directly predicting the learning rate. This approach allows us to avoid the breakdown of the formula above 10^{10} parameters mentioned in their work, where the predicted learning rate



Figure 6: (a) Quality of the fit. The maximum absolute error on the held-out extrapolation is 0.018. (b) Predicted loss compared with an observed loss for E = 1. (c) Predicted loss (dashed line) compared with an observed loss for E = 4. We can see that on the training dataset, the error increases in an undertrained setting (D/N < 1 - more tokens than parameters). However, this scenario is never practical from our perspective.

becomes negative. This phenomenon is independent of the actual fit and is simply a property of the formula used. Besides being well-defined in the extrapolation, we argue that optimal learning rates visibly follow this logarithmic trend, as seen in Figure 7 in Appendix.

Finding 4. More experts \rightarrow lower learning rate. Increasing the number of experts in MoE model should be accompanied by lowering the learning rate accordingly (Figure 7 in Appendix).

The second difference between our formula and the one by Kaplan et al. (2020) is incorporating the number of experts, allowing us to model the optimal behavior of this hyperparameter across dense models and different MoEs. This is an important detail that allows unbiased comparison among different models, ensuring that each one is optimally tuned. Furthermore, it allows us to answer the question of whether MoE should be trained with a lower or higher LR. While our formula accommodates both scenarios, we can clearly see in Figure 7 in Appendix that increasing E requires lower learning rates, resulting in a negative value for the coefficient. Moreover, we verify this thesis by tuning the fit on E = 1 and E = 8, and validating it on interpolation E = 4 and extrapolation E = 32. In both cases, the validation predicts the optimal learning rate for the model configuration or a value with practically the same performance. In Figure 8 in Appendix, we perform an ablation of this additional power law on E by repeating our entire fitting procedure without the E component. This shows, especially with the extrapolation on E = 32, that dependence on E is crucial, and its omission can impair the performance of MoEs. Further details about our scaling rule for learning rates can be found in the plots in Appendix G.

Learning Rate Schedule. Hägele et al. (2024) suggest that a constant learning rate schedule can yield similar performance to other established methods, such as the cosine schedule. At the same time, it offers a valuable advantage when varying training duration, as intermediate checkpoints can be reused when training models for a longer time. With a cosine schedule, intermediate checkpoints can introduce bias into the fit, according to the analysis of Kaplan et al. (2020) by Hoffmann et al. (2022). We employ a constant learning rate schedule with a linear warmup over the initial 130M tokens and with a linear decay from the peak learning rate to 0 over the final 20% of tokens. For each model size, longer runs reuse intermediate checkpoints from the shorter ones.

D.2 OPTIMIZATION OF FORMULA COEFFICIENTS

Following Hoffmann et al. (2022), we use the LBFGS algorithm to optimize the coefficients of formula 3. See Appendix F for details. We observe a good fit with $\text{RMSE}_v = 0.0039$ on a held-out set of our 30 runs with the lowest loss, and $\text{RMSE}_t = 0.0062$ on the training dataset. To further verify the validity of our formula, we train separate Chinchilla scaling laws 4 for different *E* using the same hyperparameters and the corresponding subset of the initializations grid. This approach serves as a

lower bound for loss of our joint formula on the training dataset, as it can emulate its coefficients; however, it is more prone to overfitting because effectively more parameters are utilized. Using this approach, we obtain lower error on the training dataset of $\text{RMSE}_t^{\text{sep}} = 0.0059$ and marginally higher on the validation $\text{RMSE}_v^{\text{sep}} = 0.0041$. We believe this is strong confirmation that our joint formula is actually describing how variable E influences training. In Figure 6, we visually verify the extrapolation of the joint fit. Prediction errors are categorized by different numbers of experts, highlighting that our joint formula is not biased for any specific E.

E TECHNICAL DETAILS

E.1 COUNTING PARAMETERS

There are many ways the size of a model can be measured. The two most important distinctions are whether total or active parameters are counted and whether the parameters in the embedding and unembedding layers are counted. Various papers assume different notations, notably Kaplan et al. (2020) use nonembedding parameters while Hoffmann et al. (2022) opt for the parameter count including embedding and unembedding. Throughout our work, we try to make it clear which way of counting we are using in each particular instance. When no additional information is given, $N_{\rm act}$ and $N_{\rm total}$ denote respectively active and total parameters, including the embedding and unembedding.

If we let d_{model} be the hidden dimension of a model, and d_{vocab} be the vocabulary size (50,257 in our case), then the following relations hold:

$$N_{\text{total}} = 2d_{\text{model}}d_{\text{vocab}} + (4+9E)N_{\text{blocks}}d_{\text{model}}^2 \tag{8}$$

$$N_{\rm act} = 2d_{\rm model}d_{\rm vocab} + 13N_{\rm blocks}d_{\rm model}^2 \tag{9}$$

E.2 COUNTING FLOPs

Basing on Sardana et al. (2024), we assume the cost of training to be $F_{\text{training}} = 6N_{\text{act}}D_{\text{training}}$, and the cost of inference to be $F_{\text{inference}} = 2N_{\text{act}}D_{\text{inference}}$. Due to the relatively small number (≤ 32) of experts used with implicit expert granularity of 1.0 (Ludziejewski et al., 2024), we can consider the memory and FLOPs cost of routing to be negligible, following Clark et al. (2022).

E.3 MODEL CONFIGS

The vast majority of our experiments use a simple rule for scaling the config, i.e. $N_{\text{blocks}} = N_{\text{heads}} = d_{\text{model}}/64$ and assume these relations hold in all calculations. We base this rule on findings by Kaplan et al. (2020).

F FIT DETAILS

Table 2: Fitted coefficients of our joined formula.

a	α	δ	γ	b	β	ω	ζ	E_{start}	E_{\max}	c
35.91	0.1889	0.2285	-0.0098	35.98	0.1775	-0.5529	0.0259	2.0732	290.4521	1.3637

Following Hoffmann et al. (2022), we use the LBFGS algorithm with a learning rate of 1e-4 and weight decay of 1e-5 to fit the coefficients of Equation 3, optimizing the Huber loss with $\delta = 0.01$ over the set of our training runs described in table in Appendix H. Instead of removing outliers and underperforming models from the training set, we underweight them proportionally to the loss. Optimization hyperparameters were manually tuned to minimize error over the training dataset. The final fitted coefficients of Equation 3 are within the boundaries of the grid of initializations given by: $\alpha \in \{0.05, 0.25, 0.5\}, \beta \in \{0.05, 0.25, 0.5\}, A \in \{30, 100, 300\}, B \in \{30, 100, 300\}, C \in \{0.5, 1, 2\}, \delta \in \{-0.5, 0, 0.5\}, \gamma \in \{-0.5, 0, 0.5\}, \omega \in \{-0.5, 0, 0.5\}, \zeta \in \{-0.5, 0, 0.5\}$. The selected coefficients were those with the lowest score, defined as the sum of RMSE on the training and a held-out extrapolation validation set. The formula in Equation 3 was calculated in

Table 3: The fitted coefficients of our joint formula, Equation equation 3, reduced to the Chinchilla scaling law, Equation equation 4, for a given number of experts, E. We observe that the dataset exponent, ν , increases significantly. This is one of the reasons why compute-optimal parameter-to-token ratios change with E.

E	m	μ	n	ν	c
1	30.3640	0.1817	53.9838	0.1965	1.3637
2	27.7982	0.1780	66.8401	0.2065	1.3637
4	24.8462	0.1731	87.7022	0.2192	1.3637
8	21.8330	0.1676	119.9126	0.2338	1.3637
16	19.0159	0.1617	167.5073	0.2494	1.3637
32	16.5424	0.1557	234.6726	0.2652	1.3637

logarithm, without any exponentials, using only linear transformations and the logsumexp operation. It was optimized to predict the logarithm of L, and parameters a, b, and c were optimized in logarithm. All these steps were taken to increase numerical stability and were essential for proper convergence.

G LEARNING RATE SCALING FIT



Figure 7: Visualization of the fit $(E \in \{1, 8\})$ of our LR scaling rule, interpolation (E = 4) and extrapolation (E = 32).



Figure 8: Ablation for the LR scaling rule fit without considering the number of experts E. While performance on the training set $(E \in \{1, 8\})$ looks acceptable, the extrapolation on E = 32 is clearly suboptimal, validating the need for considering E.

H EXPERIMENTS LISTING

N _{total}	N_{attn_heads}	N _{blocks}	d_{model}	$N_{\rm act}$	E	D
5.0B	16	16	1024	321M	32	16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
3.8B	28	28	1792	1.3B	4	11.1B, 5.6B, 2.8B, 2.0B
3.3B	11	21	1408	683M	8	16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
3.0B	26	26	1664	1.1B	4	80.0B, 64.0B, 48.0B, 32.0B, 16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
2.7B	36	36	2304	2.7B	1	9.2B, 5.5B, 2.8B, 2.0B, 1.4B, 980M
2.6B	30	30	1920	1.6B	2	5.4B, 2.7B
2.6B	16	16	1024	321M	16	16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
2.2B	28	28	1792	1.3B	2	18.6B, 11.1B, 5.6B, 4.0B, 2.8B, 2.0B
2.1B	12	12	768	169M	32	8.0B, 4.0B, 2.0B, 1.0B, 500M
2.1B	10	16	1280	469M	8	32.0B, 16.0B, 8.0B, 4.0B, 2.0B, 1.0B
1.9B	22	22	1408	709M	4	35.3B, 12.2B, 10.6B, 7.7B, 5.3B, 3.8B
1.8B	11	21	1408	683M	4	8.0B, 16.0B, 4.0B, 2.0B, 1.0B, 500M
1.8B	26	26	1664	1.1B	2	16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
1.6B	30	30	1920	1.6B	1	5.4B, 2.7B
1.4B	16	16	1024	321M	8	16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
1.3B	28	28	1792	1.3B	1	6.5B, 3.3B, 18.6B, 11.1B, 5.6B, 4.0B, 2.8B, 2.0B
1.3B	10	10	640	118M	32	4.0B, 2.0B, 1.0B, 500M
1.2B	10	16	1280	469M	4	32.0B, 16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
1.1B	12	12	768	169M	16	8.0B, 4.0B, 2.0B, 1.0B, 500M
1.1B	26	26	1664	1.1B	1	14.0B, 12.0B, 10.0B, 80.0B, 64.0B, 48.0B, 32.0B
1.1B	26	26	1664	1.1B	1	16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
1.1B	22	22	1408	709M	2	3.8B, 49.8B, 24.9B, 12.5B, 6.2B, 3.1B, 1.6B, 778M
1.1B	22	22	1408	709M	2	21.8B, 18.7B, 15.6B, 35.3B, 12.2B, 10.6B, 7.7B, 5.3B
1.1B	18	18	1152	426M	4	31.0B, 25.9B, 20.7B, 10.4B, 5.2B, 2.6B, 1.3B
1.1B	11	21	1408	683M	2	32.0B, 16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
890M	24	24	1536	890M	1	9.9B, 5.0B
850M	20	20	1280	555M	2	16.0B, 8.0B
774M	16	16	1024	321M	4	16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
709M	22	22	1408	709M	1	35.3B, 12.2B, 10.6B, 7.7B, 5.3B, 3.8B, 12.5B, 6.2B
705M	10	16	1280	469M	2	32.0B, 16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
683M	11	21	1408	683M	1	32.0B, 16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
671M	10	10	640	118M	16	4.0B, 2.0B, 1.0B, 500M
664M	8	8	512	79M	32	2.0B, 1.0B, 500M
615M	12	12	768	169M	8	8.0B, 4.0B, 2.0B, 1.0B, 500M
555M	20	20	1280	555M	1	16.0B, 8.0B
472M	16	16	1024	321M	2	16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
469M	10	16	1280	469M	1	32.0B, 16.0B, 8.0B, 4.0B, 2.0B, 1.0B, 500M
3/6M	10	10	640	118M	8	4.0B, 2.0B, 1.0B, 500M
362M	8	8	512	/9M	16	2.0B, 1.0B, 500M
360M	12	12	/68	169M	4	8.0B, 4.0B, 2.0B, 1.0B, 500M
321M	10	10	1024	321M	1	10.0B, 8.0B, 4.0B, 2.0B, 1.0B, 300M
289M	11	11	/04	142M	4	4.5B, 2.5B, 1.1B
283M	9	12	3/0	9/M	ð	3.3B, 1.7B
282M	13	13	832 769	201M	2	0.4B, 5.2B, 1.0B, 800M 8 0B, 4 0B, 2 0B, 1 0B, 500M
200M	12	12	/08 640	109M	2 4	6.0D, 4.0D, 2.0D, 1.0D, 500W
220IVI 211M	10	10	512	70M	4	4.0D, 2.0D, 1.0D, 500M
∠11WI 160M	0 10	0 12	512 769	150M	0	2.0D, 1.0D, 300000 $8.0B, 4.0B, 2.0B, 1.0B, 5000M$
109IVI 154M	12	12	640	109IVI 110N/	1	4 OR 2 OR 1 OR 500M
134M	10	10	512	110IVI 70M	2 1	4.0D, 2.0D, 1.0D, 300000 2 0B 1 0B 500M
119JW	0 10	10	640	118M	+ 1	4 OB 2 OB 1 OB 500 M
98M	10 8	10	512	79M	2	2 0B 1 0B 500M
70M	o Q	o Q	512	79M	∠ 1	2.0B, 1.0B, 500M
1 2111	0	0	512	1 2111	1	2.00, 1.00, 30000

I LIMITATIONS AND FUTURE WORK

In our work, we focus on the standard MoE variant, where the size of the expert is the same as the size of the feed-forward layer of a corresponding dense model. Some recent findings (Dai et al., 2024; Ludziejewski et al., 2024; Muennighoff et al., 2024; Team, 2024b) indicate that fine-grained MoE

models are more efficient and, most probably, would enhance our reported benefits of using MoE. Similarly, adopting a dropless MoE (Gale et al., 2022) approach instead of relying on a capacity factor could lead to further improvements. We leave the integration of those MoE improvements for future work. Moreover, our Chinchilla-based optimality analysis uses FLOPs, that may not reflect wall-clock training time of models with different architectures. While analyzing total parameter, instead of active parameter matched models partly alleviates this issue because of the same memorybottleneck, various implementations and distributed training algorithms are not considered in this work. We assumed, the Chinchilla scaling law equation 4 as the basis of our formulas. While this is well-grounded in literature, this formula is known to have limitations, especially for a wide range of token-to-parameter ratios. We observed this also in some of our experiments, as outliers often are highly under or over-trained.