

# Learning to Learn: Curriculum Selection via Soft Q-Learning for Sample-Efficient Language Models

Scott Viteri

Damir Vrabac

## 1 Overview

We propose an algorithm that enables language models to learn what data to learn from. Rather than training on randomly sampled data, the model takes actions that select training examples, receiving reward based on how well it currently predicts held-out data. A Q-function, parameterized as a small network reading the language model’s internal representations, learns to predict the long-run value of each candidate. The policy is derived from the Q-function via a Boltzmann distribution, eliminating the need for a separate policy network. The core hypothesis is that this learned curriculum selection will yield significantly better sample efficiency than random or heuristic curricula.

## 2 Technical Approach

### 2.1 Setup

- Language model (GPT-2) with parameters  $\theta$ , hidden dimension  $d = 768$
- Streaming data source, yielding fresh batches of  $N$  candidate context windows each step
- Fixed held-out evaluation set  $D$ , large; a fresh random subset  $\hat{D} \subset D$  of size  $M$  is sampled each step for reward computation
- Q-network  $Q_\phi$  with parameters  $\phi$ , implemented as a 2-layer MLP ( $d \rightarrow 32 \rightarrow 1$ ) reading the base model’s residual stream (see Section 2.3)
- Target Q-network  $Q_{\bar{\phi}}$  with parameters  $\bar{\phi}$ , a Polyak-averaged copy of  $Q_\phi$  used for stable bootstrap targets
- Boltzmann temperature  $\beta > 0$  controlling exploration
- Learning rate  $\alpha$  for the LM gradient step; learning rate  $\eta$  for the Q-network update
- Discount factor  $\gamma \in (0, 1)$  controlling the effective planning horizon
- Polyak averaging rate  $\tau \in (0, 1)$  for the target network update
- Gradient clipping threshold  $G$  (max global norm; applied to both LM and Q-network gradient steps)
- Optional: TD-into-theta flag and scaling coefficient  $\lambda_{\text{TD}}$  for representation shaping (Section 2.7)

### 2.2 State, Action, and Reward

The state  $s_k$  is the current model parameters  $\theta_k$ , which determine the model’s representations and therefore the features available to the Q-network. In practice,  $s_k$  is never represented explicitly—it is accessed implicitly through the base model’s hidden states when candidates are forwarded through it (Section 2.3).

The action  $a_k \in C_k$  is the choice of which training example to train on at step  $k$ , where  $C_k$  is the batch of  $N$  candidates available at that step. Since data is streamed, each step presents a fresh candidate batch, and each candidate is seen at most once. The Q-network generalises across batches via the base model’s hidden-state representation.

The reward is the held-out log-probability *after* the training update:

$$r_k = \frac{1}{M} \log p_{\theta_{k+1}}(\hat{D}_k), \quad (1)$$

where  $\hat{D}_k$  is a fresh random subset of  $D$  of size  $M$ , and  $\theta_{k+1}$  results from training on  $a_k$ . This measures the model’s absolute predictive fitness as a consequence of the selected action. Resampling  $\hat{D}_k$  each step prevents the Q-function from overfitting to a fixed evaluation subset.

The Q-function optimises the discounted return  $(1 - \gamma) \sum_k \gamma^k r_k$ . The weights  $(1 - \gamma) \gamma^k$  sum to one, so this is the *exponentially weighted average* of the reward sequence: the Q-value at any state equals the typical per-step reward expected over the next  $\sim 1/(1 - \gamma)$  steps under the current policy. Maximising this objective favours curricula that achieve high predictive fitness quickly—each step spent with poor predictions is penalised, with a horizon controlled by  $\gamma$ .

### 2.3 Q-Network Architecture

The Q-network reads the base model’s internal representations to assess candidate value. For each candidate  $x \in C_k$ :

1. Forward  $x$  through the base model with parameters  $\theta_k$ , extracting the last-layer hidden state at the final token position:  $h_x = \text{last\_hidden}(\theta_k, x) \in \mathbb{R}^d$ .
2. Detach  $h_x$  from the computation graph (no gradients flow into  $\theta$ ).
3. Pass through the Q-network:

$$Q_\phi(s_k, x) = W_2 \sigma(W_1 h_x + b_1) + b_2, \quad (2)$$

where  $\phi = \{W_1 \in \mathbb{R}^{32 \times d}, b_1 \in \mathbb{R}^{32}, W_2 \in \mathbb{R}^{1 \times 32}, b_2 \in \mathbb{R}\}$  and  $\sigma$  is ReLU.

The hidden state  $h_x$  implicitly encodes both the model’s current state (through  $\theta_k$ , which determines the representations) and the identity of the candidate (through the input  $x$ ). By default,  $h_x$  is detached from the computation graph, ensuring that  $\theta$  is updated only by the language modeling objective while  $\phi$  is updated only by the TD loss. This prevents the co-adaptation instability that arises when value gradients distort the base model’s representations. Section 2.7 describes an optional mode that relaxes this separation.

### 2.4 Normalised Discounted Soft Bellman Equation

The Q-function satisfies a normalised discounted soft Bellman equation:

$$Q(s_k, a_k) = (1 - \gamma) r_k + \gamma V_{\bar{\phi}}(s_{k+1}), \quad (3)$$

where  $\gamma \in (0, 1)$  is the discount factor and  $V_{\bar{\phi}}$  is the normalised soft value computed from the target network (see below). The  $(1 - \gamma)$  factor on the reward ensures that a constant reward  $r$  contributes  $r$  (not  $r/(1 - \gamma)$ ) to the Q-value fixed point. Combined with the normalised soft value (which subtracts  $\beta \log N$ ), a constant reward gives  $Q^* = r$  exactly. This keeps Q-values on the same scale as the reward and the MLP output in a numerically comfortable range.

The soft value function is normalised against the uniform prior:

$$V(s) = \beta \log \frac{1}{N} \sum_{a'} \exp(Q(s, a')/\beta) = \beta \left[ \log \sum_{a'} \exp(Q(s, a')/\beta) - \log N \right]. \quad (4)$$

The  $\log N$  subtraction removes the entropy bonus from the uniform distribution: when all Q-values are equal at  $q$ , we have  $V(s) = q$  rather than  $V(s) = q + \beta \log N$ . This is the *normalised* logsumexp, or equivalently the log-mean-exp. Without it, the entropy bonus  $\beta \log N$  would be amplified by  $\gamma/(1 - \gamma)$  at the Bellman fixed point, inflating Q-values far above the reward scale.

With both normalisations— $(1 - \gamma)$  on the reward and  $-\log N$  in the soft value—a constant reward  $r$  gives  $Q^* = r$  at the fixed point. Q-values live on the same scale as the reward (around  $-3$  to  $-4$  for GPT-2 perplexity), and the Q-network’s output is directly interpretable as the expected per-step reward over the planning horizon. At convergence,  $Q(s, a)$  represents a normalised discounted sum of future rewards from choosing  $a$  in state  $s$  and following the Boltzmann policy thereafter. With  $\gamma = 0.9$ , the effective horizon is  $\sim 10$  steps.

The policy  $\pi(a \mid s) \propto \exp(Q(s, a)/\beta)$  is unaffected by the  $\log N$  subtraction, since softmax is shift-invariant.

**Target network.** To stabilise the bootstrap target  $V(s_{k+1})$ , we maintain a target network  $Q_{\bar{\phi}}$  that is a Polyak-averaged copy of the online network  $Q_{\phi}$ :

$$\bar{\phi} \leftarrow \tau \phi + (1 - \tau) \bar{\phi}, \quad \tau \in (0, 1). \quad (5)$$

The soft value in the TD target is computed from  $Q_{\bar{\phi}}$ , not  $Q_{\phi}$ . This breaks the feedback loop where the network’s own output serves as its training target, preventing the oscillations and divergence that can occur with function approximation. The online network  $Q_{\phi}$  is still used for action selection (the Boltzmann policy) and for the prediction  $\hat{Q}$  in the TD loss. With  $\tau = 0.01$ , the target network tracks the online network with a lag of  $\sim 1/\tau = 100$  steps.

## 2.5 Algorithm

The key implementation insight is that each step’s forward pass of candidates through the base model serves two purposes: action selection for the *current* step and the bootstrap target for the *previous* step’s Q update. This avoids a redundant second forward pass.

The LM trains on a *policy-weighted mixture* of all  $N$  candidates at every step, rather than on a single sampled action. Define the Boltzmann weights:

$$\pi_i = \frac{\exp(Q_{\phi}(s_k, x_i)/\beta)}{\sum_{j=1}^N \exp(Q_{\phi}(s_k, x_j)/\beta)}, \quad x_i \in C_k. \quad (6)$$

The LM loss is the policy-weighted mixture:

$$\mathcal{L}_{\text{mix}}(\theta; C_k) = \sum_{i=1}^N \pi_i \ell(\theta, x_i), \quad (7)$$

where  $\ell(\theta, x_i)$  is the per-token cross-entropy loss on candidate  $x_i$ . This is precisely the *expected gradient* under the Boltzmann policy:

$$\nabla_{\theta} \mathcal{L}_{\text{mix}} = \mathbb{E}_{x \sim \pi(\cdot \mid s_k)} [\nabla_{\theta} \ell(\theta, x)], \quad (8)$$

computed exactly rather than estimated via a single Monte Carlo sample. This eliminates the variance from action sampling.

**Q-learning and the mixture LM update.** The Q-network update is unchanged by the mixture formulation: a discrete action  $a_k$  is still sampled from the Boltzmann policy, its hidden state  $h_{\text{prev}}$  is stored, and the standard TD update is applied at the next step. The Q-function continues to satisfy the discounted soft Bellman equation (3). The only difference from a single-action formulation is how the environment *transitions*: the LM parameters are updated via the full mixture rather than a single example.

**Off-policy interpretation.** From the Q-learner’s perspective, the situation is analogous to off-policy learning. The Q-learner observes: “I selected action  $a_k$ , the environment transitioned to state  $s_{k+1}$ , and I received reward  $r_k$ .” It does not need to know that the transition was caused by a mixture-weighted gradient step rather than a single-example step. Q-learning is inherently off-policy—the Bellman backup  $Q(s, a) = (1 - \gamma)r + \gamma V(s')$  is valid regardless of the behavior policy that generated the transition, as long as  $r$  and  $s'$  are the actual observed reward and next state.

Over many steps, the Q-function converges to predict the value of upweighting a given candidate within the mixture—which is exactly the marginal contribution signal needed to set the mixture weights well.

**Effective batch size.** The mixture-weighted update is equivalent to training on a soft batch of  $\sim 1/\|\pi\|_2^2$  effective examples per step (the inverse of the collision probability of the policy distribution). Early in training, when Q-values are similar, this approaches  $N$ ; as the policy sharpens, the effective batch size decreases toward 1.

**Computational cost per step.** One forward pass of  $N$  candidates through the base model (batched, no gradient) for Q-scoring;  $N$  forward-backward passes through the full model for the mixture LM loss (processed in mini-batches with gradient accumulation); one forward pass of  $M$  held-out examples for reward; one MLP forward-backward for the Q update; one MLP forward pass through the target network (negligible). The candidate extraction forward passes dominate; the mixture LM step adds roughly a  $2\times$  factor to total step time compared to a single-example LM step.

**Staleness.** The Q update at step  $k$  uses the target network  $Q_{\bar{\phi}}$  for the bootstrap value  $V_k$ , and the online network  $Q_{\phi}$  for action selection. The target network lags the online network by  $\sim 1/\tau$  steps, which is by design: this lag stabilises the TD target. The online Q-values  $q_k$  used for action selection have one-step staleness (computed before the  $\phi$  update), which is standard in online Q-learning and negligible given the small MLP learning rate  $\eta$ .

## 2.6 Uniform Mixture Variant (No Q-Weighting)

As an ablation, the flag `--no.q.weighting` replaces the Boltzmann policy weights with uniform weights:

$$\pi_i = \frac{1}{N}, \quad i = 1, \dots, N. \quad (9)$$

The LM loss becomes the unweighted average over all candidates:

$$\mathcal{L}_{\text{uniform}}(\theta; C_k) = \frac{1}{N} \sum_{i=1}^N \ell(\theta, x_i). \quad (10)$$

This decouples the LM training step from the Q-function entirely: the LM trains on an equal mixture of all candidates regardless of their Q-values. The Q-network still trains normally (receiving rewards, computing TD updates), but its output does not influence which data the LM learns from. This serves as a controlled ablation to isolate the effect of Q-weighted curriculum selection on sample efficiency.

## 2.7 TD Gradients into Theta (Representation Shaping)

By default, the hidden states  $h_x$  used by the Q-network are detached from  $\theta$ , so the TD loss  $\frac{1}{2}(\hat{q}-y)^2$  only updates the Q-network parameters  $\phi$ . This creates a one-directional dependency:  $\theta$  determines the representations that  $\phi$  reads, but  $\phi$ 's learning signal never shapes  $\theta$ . If  $\theta$  changes quickly (e.g. under a high LM learning rate), the Q-head may struggle to track the shifting representation landscape.

The flag `--td_into_theta` enables a bidirectional coupling: during the TD update, the previous action's hidden state is recomputed through the current  $\theta$  *with* gradients, so that  $\nabla_{\theta}\mathcal{L}_{\text{TD}}$  is nonzero and flows into  $\theta$  via the LM optimizer. This gives  $\theta$  a secondary objective: produce representations that are informative for Q-value prediction.

Concretely, at step  $k$  the Q update becomes:

1. Re-forward the previous action's tokens  $a_{k-1}$  through the current model  $\theta_k$  *with gradients*:

$$h_{\text{prev}} = \text{last\_hidden}(\theta_k, a_{k-1}) \in \mathbb{R}^d. \quad (\text{grad-connected to } \theta_k)$$

2. Compute the TD loss as before:

$$\mathcal{L}_{\text{TD}} = \frac{1}{2}(Q_{\phi}(h_{\text{prev}}) - y)^2, \quad y = (1 - \gamma)r_{\text{prev}} + \gamma V_{\bar{\phi}}(s_k).$$

3. Backpropagate:  $\nabla_{\phi}\mathcal{L}_{\text{TD}}$  updates  $\phi$  as usual;  $\lambda_{\text{TD}}\nabla_{\theta}\mathcal{L}_{\text{TD}}$  updates  $\theta$  via the LM optimizer. The coefficient  $\lambda_{\text{TD}}$  (`--td_lambda`, default 1.0) controls the relative strength of the TD signal on  $\theta$ . Setting  $\lambda_{\text{TD}} < 1$  downweights the TD contribution to avoid overwhelming the language modeling gradient. The LM optimizer thus receives gradients from two sources each step: the LM loss (from the mixture or retrieved-example training) and the scaled TD loss.

**Cross-step graph avoidance.** A naive implementation would require keeping the full computation graph from step  $k-1$  alive until step  $k$ , which is prohibitively expensive. Instead, we store the *token IDs* of the previous action (not the hidden state), and re-forward them through the current  $\theta_k$  at the next step. This costs one extra single-example transformer forward pass per step—negligible compared to the  $N$ -candidate or RAG forward passes—and avoids any cross-step graph retention.

**Relation to auxiliary tasks.** This is analogous to auxiliary-task representation learning in RL (Jaderberg et al., 2017; UNREAL): the TD loss acts as an auxiliary objective that encourages the base model to maintain representations useful for value prediction. Unlike standard auxiliary tasks, here the auxiliary signal is the Q-learning TD error itself, creating a closed loop between the value function and the representations it operates on.

**Note on the bootstrap target.** Only the Q-prediction side  $Q_{\phi}(h_{\text{prev}})$  carries gradients into  $\theta$ . The bootstrap target  $y = (1 - \gamma)r + \gamma V_{\bar{\phi}}(s')$  remains stop-gradient with respect to both  $\phi$  and  $\theta$ , as is standard in Q-learning. The target-network value  $V_{\bar{\phi}}$  uses detached hidden states.

### 3 Softmax-Relaxed Langevin-RAG Curriculum Selection

The discrete-Q approach (Section 2.5) scores a fixed batch of  $N$  candidates per step. This limits the search to the  $N$  examples that happen to appear in the current batch—a tiny fraction of the training corpus. To search over a much larger space of possible training data, we introduce a pipeline that uses the Q-network as an energy function for continuous optimisation via Stochastic Gradient Langevin Dynamics (SGLD), then retrieves real training examples from a large indexed corpus via Retrieval-Augmented Generation (RAG).

A naïve approach would run SGLD directly in the model’s embedding space  $\mathbb{R}^d$ , then snap the resulting continuous vectors to discrete tokens via nearest-neighbor lookup. However, this suffers from a *snap bottleneck*: Langevin dynamics pushes embeddings off the token manifold into regions of  $\mathbb{R}^d$  that are far from any real token, so nearest-neighbor snapping destroys the information that SGLD optimised. Our solution is to run SGLD in *logit space* over the vocabulary, using a differentiable softmax bridge to keep the transformer’s input always in the convex hull of real token embeddings. The gradient path from Q-value back through the transformer, through the softmax bridge, and into the logits is fully differentiable—no reparameterisation trick (e.g. Gumbel-Softmax) is needed, since we never backpropagate through a discrete sampling step.

#### 3.1 Overview

The pipeline proceeds in four stages per training step:

1. **Softmax-relaxed SGLD in logit space** (Section 3.2): Maintain  $K$  parallel chains, each a tensor of logits  $\boldsymbol{\lambda} \in \mathbb{R}^{L \times V}$  over the vocabulary at every token position. At each Langevin step, form soft token embeddings via  $\text{softmax}(\boldsymbol{\lambda}/\tau_t) \cdot W_e$ , feed through the transformer and Q-network, backpropagate  $\partial Q/\partial \boldsymbol{\lambda}$ , and apply the Langevin update. After all SGLD steps, collect one sample per chain.
2. **Token extraction** (Section 3.3): Extract discrete tokens via arg max of the logits—no lossy nearest-neighbor snap.
3. **RAG retrieval** (Section 3.4): Embed the query sentences with a sentence-transformer and retrieve top- $k$  real training examples from a FAISS index.
4. **LM training**: Train the language model on the retrieved examples (uniform or Q-weighted loss).

The key insight is that the softmax constraint  $\text{softmax}(\boldsymbol{\lambda}/\tau) \cdot W_e$  keeps the transformer’s input in the *convex hull* of real token embeddings at all times. As SGLD progresses and the softmax sharpens, each position’s distribution concentrates around a specific token—the one that maximises the Q-value in context. The arg max at the end simply reads off which token dominates, rather than performing a lossy projection.

#### 3.2 SGLD in Logit Space

**Optimisation variable.** Let  $\boldsymbol{\lambda} \in \mathbb{R}^{L \times V}$  be a matrix of unconstrained logits, where  $L$  is the query sequence length and  $V$  is the vocabulary size. Each row  $\boldsymbol{\lambda}_i \in \mathbb{R}^V$  parameterises a categorical distribution over tokens at position  $i$ .

**Softmax bridge.** Given a softmax temperature  $\tau_t > 0$ , the soft token probabilities and soft embeddings are:

$$\boldsymbol{\pi}_i = \text{softmax}(\boldsymbol{\lambda}_i/\tau_t), \quad \tilde{\mathbf{e}}_i = \sum_{v=1}^V \pi_{i,v} W_e[v] = \boldsymbol{\pi}_i^\top W_e, \quad (11)$$

where  $W_e \in \mathbb{R}^{V \times d}$  is the frozen token embedding matrix. This is a plain softmax—no Gumbel noise is added, because the gradient path  $\boldsymbol{\lambda} \rightarrow \text{softmax} \rightarrow \tilde{\mathbf{e}} \rightarrow \text{transformer} \rightarrow Q$  is fully differentiable and we never need to backpropagate through a discrete sampling or argmax operation. The soft embedding  $\tilde{\mathbf{e}}_i$  is a convex combination of real token embeddings, weighted by the softmax probabilities. When  $\tau_t$  is large,  $\boldsymbol{\pi}_i$  is diffuse and  $\tilde{\mathbf{e}}_i$  is a broad average; when  $\tau_t$  is small,  $\boldsymbol{\pi}_i$  concentrates on one token and  $\tilde{\mathbf{e}}_i \approx W_e[v^*]$  for the dominant token  $v^*$ .

**Energy function.** The energy is defined on the logits via the softmax bridge:

$$E(\boldsymbol{\lambda}; \tau_t) = \frac{1}{\tau_{\text{sld}}} Q_\phi(\text{transformer}(\tilde{\mathbf{e}}(\boldsymbol{\lambda}, \tau_t))), \quad (12)$$

where  $\text{transformer}(\tilde{\mathbf{e}})$  feeds the soft embeddings as `inputs_embeds`, extracts the last-layer hidden state at the final position, and passes it through the Q-network. The SGLD temperature  $\tau_{\text{sld}}$  controls how peaked the Boltzmann distribution  $p(\boldsymbol{\lambda}) \propto \exp(E(\boldsymbol{\lambda}))$  is.

**Langevin update.** SGLD produces approximate samples via:

$$\boldsymbol{\lambda}_{t+1} = \boldsymbol{\lambda}_t + \frac{\varepsilon}{2} \nabla_{\boldsymbol{\lambda}} E(\boldsymbol{\lambda}_t; \tau_t) + \sqrt{\varepsilon} \boldsymbol{\eta}_t, \quad \boldsymbol{\eta}_t \sim \mathcal{N}(0, \sigma^2 I), \quad (13)$$

where  $\varepsilon$  is the step size and  $\sigma$  the noise scale. The gradient  $\nabla_{\boldsymbol{\lambda}} E$  is computed by backpropagating from the Q-value through the Q-network, the transformer, the softmax bridge, and into the logits. Gradient checkpointing is used for the transformer blocks to keep memory bounded.

**Temperature annealing.** The softmax temperature  $\tau_t$  is annealed linearly from  $\tau_{\text{start}}$  (high, exploratory) to  $\tau_{\text{end}}$  (low, sharp) over the SGLD steps:

$$\tau_t = \tau_{\text{start}} + \frac{t}{T_{\text{max}} - 1} (\tau_{\text{end}} - \tau_{\text{start}}). \quad (14)$$

This creates a natural coarse-to-fine search: early steps explore broadly (high  $\tau$ , soft distributions), while later steps refine toward specific tokens (low  $\tau$ , peaked distributions). The annealing schedule is analogous to simulated annealing, but applied to the softmax bridge rather than the SGLD noise.

**Non-stationarity.** Because  $\tau_t$  changes the energy landscape at each step, the SGLD chain does not converge to a stationary Boltzmann distribution. The procedure is better understood as *optimisation with injected noise* (gradient ascent on Q, regularised by Langevin noise for diversity) rather than as approximate posterior sampling. The noise prevents the chains from collapsing to a single mode, while the annealing schedule progressively sharpens the softmax so that the final logits represent well-defined token choices.

**Initialisation.** Each chain’s logits are initialised as approximate one-hots: for each position, a random token is chosen uniformly from the vocabulary and assigned a high logit value (e.g. 10), with all other logits at 0. This ensures the initial soft embeddings are very close to real token embeddings, starting SGLD on the token manifold.

**Gradient clipping.** Per-chain gradient norms (flattened across all  $L \times V$  logit entries) are clipped to a threshold  $G$  for stability.

**Collection.** All  $T_{\max}$  SGLD steps run to completion, then the final logits from each of the  $K$  chains are collected, yielding  $K$  samples (one per chain). The number of SGLD steps serves as a burn-in period: enough steps for the chains to climb the Q-landscape and for the softmax temperature to anneal to  $\tau_{\text{end}}$ .

### 3.3 Token Extraction

After SGLD, discrete tokens are extracted via argmax of the collected logits:

$$t_i = \arg \max_{v \in \{1, \dots, V\}} \lambda_{i,v}, \quad i = 1, \dots, L. \quad (15)$$

Unlike the legacy approach (nearest-neighbor cosine similarity in  $\mathbb{R}^d$ ), this is *exact*: the argmax token *is* the dominant mode of the softmax distribution that SGLD optimised. No information is lost in the continuous-to-discrete conversion, because the optimisation variable (logits) directly parameterises the discrete choice.

**Sharpness diagnostic.** The quality of token extraction is measured by the *softmax entropy*  $H = -\sum_v \pi_v \log \pi_v$  and the *max probability*  $\max_v \pi_v$  at the final softmax temperature. Low entropy / high max-probability indicates that the SGLD has converged to well-defined token choices. For reference, a uniform distribution over  $V = 50,257$  tokens has entropy  $\log V \approx 10.8$ ; a perfectly sharp distribution has entropy 0 and max-probability 1.

### 3.4 RAG Retrieval

**Index construction and rolling refresh.** At startup we stream through the training corpus, extract  $I$  fixed-length token windows, embed them with a pretrained sentence-transformer (all-MiniLM-L6-v2,  $d_{\text{st}} = 384$ , normalised), and store the resulting vectors in a FAISS `IndexFlatIP` (flat brute-force inner product—no tree or quantisation). The embeddings, token IDs, and decoded texts are held in a fixed-size *ring buffer* of capacity  $I$ .

Each training step,  $n_{\text{ingest}}$  fresh windows are consumed from the data stream, embedded, and written into the ring buffer at the current write head, overwriting the oldest entries. The FAISS index is then rebuilt from the updated embedding matrix. Because `IndexFlatIP` is just a dense matrix, rebuilding for  $I = 50,000$  vectors of dimension 384 amounts to a single `memcpy` ( $< 5$  ms). With  $n_{\text{ingest}} = 100$  and  $I = 50,000$ , each step replaces 0.2% of the index. The full buffer turns over after 500 steps—the same total throughput as a bulk refresh every 500 steps, but spread smoothly to avoid the LM loss jumps caused by a sudden distribution shift.

**Query and retrieval (each step).** The  $K$  discrete query sequences from the token extraction step are decoded to text and embedded with the same sentence-transformer. For each query, the top- $k$  nearest neighbours in the FAISS index are retrieved by cosine similarity. By default, one result is *softmax-sampled* from the top- $k$  per query rather than taking the deterministic top-1:

$$p_j \propto \exp(\text{score}_j / \tau_{\text{rag}}), \quad j = 1, \dots, k, \quad (16)$$

where  $\tau_{\text{rag}}$  is a sampling temperature (default 0.1). This gives diversity: instead of every query selecting its single nearest neighbour (which often overlap), each query draws from its local neighbourhood, producing a more varied retrieved set. Results are deduplicated across all queries to produce  $R \leq K$  unique training examples.



### 3.5 Softmax-Relaxed Langevin-RAG Algorithm

**Why logit-space SGLD avoids the snap bottleneck.** In the legacy embedding-space approach, Langevin dynamics in  $\mathbb{R}^d$  per position pushes embeddings into regions far from any token (snap cosine similarity  $\approx 0.22$ , barely above the random baseline of 0.17), and nearest-neighbor snapping destroys the optimised information. In the softmax-relaxed approach, the softmax constraint ensures the transformer’s input is always a convex combination of real token embeddings. The Q-network’s gradients flow through the softmax into the logits, directly shaping which tokens are favoured. As  $\tau_t$  decreases, the softmax sharpens and the dominant token emerges naturally—arg max simply reads it off.

**LM training (uniform or Q-weighted).** The LM can train on retrieved examples with either *uniform* loss ( $w_i = 1/R$ ) or *Q-weighted* loss ( $w_i = \text{softmax}(Q(h_i)/\beta)$ ). In the uniform case, the curriculum signal is entirely in *which* examples are retrieved. In the Q-weighted case, examples the Q-network considers more valuable receive a larger share of the gradient, combining selection and weighting.

**Q-learning is off-policy.** The Q-network update is identical to the discrete-Q version: observe reward  $r_k$ , compute the TD target from the target network, and minimise the squared TD error. The Q-learner does not need to know how the training data was selected—it only observes the resulting reward. This makes the pipeline compatible with the same normalised discounted soft Bellman equation (Eq. (3)).

**Shared structure.** The same GPT-2 model (parameters  $\theta$ ) is used for both the SGLD energy computation (with the Q-head) and the LM training. The Q-head is a small MLP ( $d \rightarrow 32 \rightarrow 1$ ) that reads the transformer’s last-layer hidden state. This maximises parameter sharing: the Q-network’s gradient signal during SGLD flows through the same transformer representations that get updated during LM training.

### 3.6 SGLD Diagnostics

Monitoring SGLD health is critical for tuning the Langevin hyperparameters. We track:

- **Q gain ( $\Delta Q$ ):** The difference between Q-values of collected samples and Q-values at initialisation. Positive  $\Delta Q$  indicates that SGLD is climbing the Q landscape.
- **Random-Q baseline:** Q-values of fresh random logits, evaluated each step. This provides an *unbiased* reference—the Q-values of collected samples are biased high by construction.
- **Softmax sharpness:** The mean entropy  $H = -\sum_v \pi_v \log \pi_v$  and mean max-probability  $\max_v \pi_v$  of the softmax distribution at the final temperature  $\tau_{\text{end}}$ . Low entropy / high max-probability indicates convergence to well-defined token choices. This replaces the snap cosine similarity metric of the legacy approach and is more interpretable: a max-probability of 0.9 means the dominant token accounts for 90% of the soft embedding’s weight.
- **Diversity:** Pairwise cosine similarity of collected soft embedding vectors (lower = more diverse), and pairwise Jaccard similarity of extracted token sets.
- **Gradient health:** Mean gradient norm and fraction of steps where gradient clipping activates.
- **Per-phase timing:** Wall-clock seconds for SGLD, RAG, LM training, and reward evaluation.

### 3.7 Interpretation as Free-Energy Minimisation

Both the discrete-Q and Langevin-RAG pipelines can be understood through a unified free-energy lens. In the discrete-Q mode, the Boltzmann policy over a finite candidate set  $C_k$  is the exact solution to a free-energy minimisation problem at each state:

$$\pi^*(\cdot | s) = \arg \min_{\pi} \left[ -\mathbb{E}_{x \sim \pi} [Q_{\phi}(s, x)] + \beta \text{KL}(\pi \| p_0) \right], \quad (17)$$

where  $p_0$  is the uniform distribution over  $C_k$ . Since  $p_0$  is uniform,  $\text{KL}(\pi \| p_0) = \log N - H(\pi)$ , and the objective reduces (up to constants) to:

$$\pi^*(\cdot | s) = \arg \min_{\pi} \left[ -\mathbb{E}_{x \sim \pi} [Q_{\phi}(s, x)] - \beta H(\pi) \right]. \quad (18)$$

The first term encourages selecting high-value candidates; the second encourages exploration. The temperature  $\beta$  controls the tradeoff.

In the Langevin-RAG mode, SGLD approximately samples from the Boltzmann distribution  $p(\lambda) \propto \exp(E(\lambda))$  over the continuous logit space. The Langevin noise plays the role of the entropy term—it prevents the chains from collapsing to a single mode, maintaining diversity among the  $K$  samples. However, because the softmax temperature  $\tau_t$  is annealed during the run, the energy landscape is non-stationary and the chain does not converge to a fixed Boltzmann distribution. In practice the procedure functions as Q-guided optimisation with noise-injected diversity, rather than as exact posterior sampling.

**ELBO interpretation.** Define a latent variable model where the “evidence” is the event that the current trajectory is optimal, with likelihood  $p(\text{optimal} | x) \propto \exp(Q(s, x)/\beta)$ . The policy  $\pi$  acts as a variational posterior over actions, and the free energy is the negative ELBO:

$$\log p(\text{optimal} | s) \geq \mathbb{E}_{x \sim \pi} \left[ \frac{Q(s, x)}{\beta} \right] - \text{KL}(\pi \| p_0). \quad (19)$$

The Boltzmann policy makes this bound tight. The normalised soft value  $V(s) = \beta[\log \sum_x \exp(Q(s, x)/\beta) - \log N]$ —which appears as the bootstrap term in the TD target—is the log-evidence minus  $\beta \log N$ , measuring how much value the current candidate batch offers above the uniform baseline.

## 4 Evaluation

### 4.1 Primary Metric: Sample Efficiency

The central question is: how many training examples does the model need to reach a given performance level?

We will measure:

- Perplexity on a fixed evaluation set as a function of training examples seen
- Performance on downstream tasks (e.g., MMLU, HellaSwag) as a function of training examples

### 4.2 Baselines

- **Random curriculum:** Uniform sampling from candidates
- **Loss-based curriculum:** Prioritise high-loss examples
- **Uncertainty-based curriculum:** Prioritise examples with high model uncertainty
- **Competence-based curriculum:** Examples ordered by difficulty (requires difficulty labels)

### 4.3 Analysis

Beyond aggregate metrics, we will examine:

- **Curriculum structure:** Does the learned curriculum exhibit interpretable patterns? Developmental stages? Topic clustering?
- **Exploration dynamics:** How does the policy entropy evolve over training? Does the temperature  $\beta$  produce reasonable exploration?
- **Q-function interpretability:** Which candidates receive high Q-values at different stages of training? Does the Q-function learn to identify examples that are valuable given the model’s current state?
- **Q-value dynamics:** How do Q-values evolve over training? Convergence of Q-values indicates the value function has stabilised; the spread (standard deviation) reflects how strongly the Q-function discriminates between candidates.

## 5 Relation to Prior Work

### 5.1 Curriculum Learning

Graves et al. (2017) use multi-armed bandits to select tasks; Jiang et al. (2018) train a separate “mentor” network to weight examples. Our approach differs by using the language model’s own representations as features for the Q-function and applying soft Q-learning with temporal credit assignment, rather than treating selection as a stateless bandit problem.

### 5.2 Meta-Learning

MAML (Finn et al., 2017) learns initializations for fast adaptation. We share the computational structure—reasoning about the effect of gradient updates—but learn *what* to train on rather than *where* to start.

### 5.3 RL as Inference

The control-as-inference framework (Levine, 2018; Rawlik et al., 2013) casts optimal control as probabilistic inference, with the Q-function serving as an energy function and the optimal policy as the corresponding Boltzmann distribution. Our algorithm instantiates this framework in the curriculum selection setting: the Q-network provides unnormalized log-probabilities over actions, and the Boltzmann policy is derived analytically without requiring a separate policy network. This is the soft Q-learning approach of Haarnoja et al. (2017), applied to curriculum selection with a standard discount factor for Bellman contraction.

### 5.4 Soft Q-Learning

Soft Q-learning (Haarnoja et al., 2017) augments the standard Bellman equation with an entropy bonus, yielding a Boltzmann policy without requiring a separate policy network. SAC (Haarnoja et al., 2018) extends this to continuous actions with a learned temperature. Our setting is simpler: the action space is a finite set of  $N$  candidates, so the Boltzmann policy and soft value can be computed exactly via softmax and logsumexp. Following standard practice in deep Q-learning (Mnih et al., 2015; Lillicrap et al., 2016), we use a Polyak-averaged target network for the bootstrap value, normalise the reward by  $(1 - \gamma)$ , and subtract  $\log N$  from the soft value to keep Q-values on the same scale as the reward.

## 5.5 Intrinsic Motivation

Our objective relates to compression progress (Schmidhuber) and active inference, but avoids the memory requirements of the former and the dark room problem of the latter by using a fixed held-out set as a proxy for predictive success. The absolute log-probability reward (1) provides a direct measure of predictive fitness, connecting to the epistemic homeostasis motivation: the agent is penalised for *being* in a state of poor prediction, not merely for failing to improve.

## 5.6 Connection to Prior Work

This proposal builds directly on Markovian Transformers for Informative Language Modeling (<https://arxiv.org/abs/2404.18988>), which introduces a framework for training language models with RL to produce causally load-bearing Chain-of-Thought reasoning. Both projects share a common structure: use RL to learn intermediate representations that improve prediction on held-out data. In the Markovian Transformers work, the learned object is a CoT:

Question  $\rightarrow$  CoT  $\rightarrow$  Answer

In the current proposal, the learned object is a curriculum:

Model State  $\rightarrow$  Selected Data  $\rightarrow$  Improved Predictions

The Markovian Transformers work demonstrates that this general approach is tractable and yields large gains on reasoning benchmarks (e.g., GSM8K: 19.6%  $\rightarrow$  57.1%). The current proposal extends this framework from learning *what to say* to learning *what to study*.

# 6 Broader Motivation

Language models trained to predict text develop remarkable capabilities from a simple objective. Yet they require extensive post-training to behave agentially and arguably lack a kind of global coherence. One hypothesis: the training process is purely observational—the model never takes actions that affect what it observes.

This project is a stepping stone toward studying whether learned curriculum selection produces qualitatively different agents. The immediate goal is demonstrating sample efficiency gains. The longer-term question is whether controlling one’s own learning process contributes to the coherence and agency that current models seem to lack.

## 6.1 Long-Term Direction: Formalizing Homeostasis

Biological agents are shaped by survival pressures. Hunger, pain, and fatigue are not arbitrary reward signals—they are tied to the organism’s continued existence. Current approaches to intrinsic motivation (curiosity, empowerment, compression progress) capture aspects of adaptive behavior but lack this grounding in self-preservation.

A long-term goal of this research program is to formalize homeostasis and survival into a simple, biologically plausible metric that could serve as a foundation for intrinsic motivation in artificial systems. The current project—learning to select data that improves future prediction—is a minimal step in this direction: the agent takes actions that maintain its predictive capacity, a kind of epistemic homeostasis. The absolute reward formulation makes this connection explicit: the agent is directly penalised for poor predictive fitness at every moment, not merely for failing to improve.

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**Algorithm 1** Curriculum Selection via Normalised Discounted Soft Q-Learning
 

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```

1: Initialise: LM parameters  $\theta$ ; Q-network parameters  $\phi$ ; target parameters  $\bar{\phi} \leftarrow \phi$ 

2: // Bootstrap step (no Q update)
3:  $C_0 \leftarrow \text{NEXTBATCH}(\text{stream}, N)$ 
4:  $H_0 \leftarrow \text{DETACHEDHIDDEN}(\theta, C_0)$  ▷  $[N, d]$  — no grad into  $\theta$ 
5:  $q_0 \leftarrow Q_\phi(H_0)$  ▷  $[N, 1]$ 
6:  $\pi_0 \leftarrow \text{SOFTMAX}(q_0/\beta)$  ▷ Boltzmann policy weights
7:  $a_0 \leftarrow \text{SAMPLE}(\pi_0)$ 
8:  $h_{\text{prev}} \leftarrow H_0[a_0]$  ▷ Store hidden state of selected action
9:  $\theta \leftarrow \theta - \alpha \nabla_\theta \sum_i \pi_{0,i} \ell(\theta, C_0[i])$  ▷ Mixture LM update, clipped to  $\|\nabla\| \leq G$ 
10:  $r_{\text{prev}} \leftarrow \frac{1}{M} \log p_\theta(\hat{D}_0)$  ▷ Reward after update

11: for  $k = 1, 2, \dots$  do // Main loop
12:    $C_k \leftarrow \text{NEXTBATCH}(\text{stream}, N)$ 
13:    $H_k \leftarrow \text{DETACHEDHIDDEN}(\theta, C_k)$  ▷ Serves dual purpose below
14:    $q_k \leftarrow Q_\phi(H_k)$ 

15:   // Q update for previous transition (target network for bootstrap)
16:    $\bar{q}_k \leftarrow Q_{\bar{\phi}}(H_k)$  ▷ Target network Q-values, no gradient
17:    $V_k \leftarrow \beta [\log \sum_{a'} \exp(\bar{q}_k[a']/\beta) - \log N]$  ▷ Normalised soft value from target network
18:    $y \leftarrow (1 - \gamma) r_{\text{prev}} + \gamma V_k$  ▷ Normalised TD target, stop-gradient
19:    $h_{\text{prev}} \leftarrow \text{HIDDEN}(\theta, a_{\text{prev}})$  ▷ Detached by default; with-grad if --td_into_theta (Sec. 2.7)
20:    $\hat{q} \leftarrow Q_\phi(h_{\text{prev}})$  ▷ Re-evaluate previous action with current  $\phi$ 
21:   Update  $\phi$  (and optionally  $\theta$ ) to minimise  $\frac{1}{2}(\hat{q} - y)^2$ , clipped to  $\|\nabla\| \leq G$ 
22:    $\bar{\phi} \leftarrow \tau \phi + (1 - \tau) \bar{\phi}$  ▷ Polyak update

23:   // Action selection for current step (for Q-learning backup)
24:    $\pi_k \leftarrow \text{SOFTMAX}(q_k/\beta)$  ▷ Boltzmann policy weights
25:    $a_k \leftarrow \text{SAMPLE}(\pi_k)$ 
26:    $h_{\text{prev}} \leftarrow H_k[a_k]$ 

27:   // LM mixture update and reward
28:    $\theta \leftarrow \theta - \alpha \nabla_\theta \sum_i \pi_{k,i} \ell(\theta, C_k[i])$  ▷ Mixture LM update, clipped to  $\|\nabla\| \leq G$ 
29:    $r_k \leftarrow \frac{1}{M} \log p_\theta(\hat{D}_k)$  ▷ Fresh  $\hat{D}_k \subset D$ , reward after update
30:    $r_{\text{prev}} \leftarrow r_k$ 
31: end for

```

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**Algorithm 2** Curriculum Selection via Softmax-Relaxed Langevin-RAG

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1: Initialise: LM parameters  $\theta$ ; Q-network  $\phi$ ; target  $\bar{\phi} \leftarrow \phi$ 
2: Build FAISS ring buffer of capacity  $I$  from first  $I$  corpus windows (Section 3.4)

3: for  $k = 0, 1, 2, \dots$  do

4:   // Stage 1: Softmax-relaxed SGLD in logit space
5:   Initialise  $K$  chains:  $\lambda_0^{(j)} \leftarrow \text{one-hot-logits}(\text{random tokens}), j = 1, \dots, K$ 
6:   for  $t = 0, 1, \dots, T_{\max} - 1$  do
7:      $\tau_t \leftarrow \tau_{\text{start}} + \frac{t}{T_{\max}-1}(\tau_{\text{end}} - \tau_{\text{start}})$  ▷ Anneal softmax temperature
8:      $\tilde{\mathbf{e}}^{(j)} \leftarrow \text{softmax}(\lambda_t^{(j)} / \tau_t) \cdot W_e$  ▷ Soft token embeddings, Eq. (11)
9:      $Q^{(j)} \leftarrow Q_\phi(\text{transformer}(\tilde{\mathbf{e}}^{(j)}))$  ▷ Energy via Eq. (12)
10:     $\lambda_{t+1}^{(j)} \leftarrow \lambda_t^{(j)} + \frac{\varepsilon}{2} \text{clip}(\nabla_{\lambda} E, G) + \sqrt{\varepsilon} \eta_t$  ▷ Langevin update, Eq. (13)
11:  end for
12:  Collect final logits  $\lambda_{T_{\max}}^{(1)}, \dots, \lambda_{T_{\max}}^{(K)}$  ▷ One sample per chain

13:  // Stage 2: Token extraction (argmax — no snap)
14:  For each collected  $\lambda$ , compute  $t_i = \arg \max_v \lambda_{i,v}$  via Eq. (15)

15:  // Stage 3: RAG retrieval + rolling refresh
16:  Embed query texts  $\{\text{decode}(\mathbf{t})\}$  with sentence-transformer
17:  Retrieve  $R$  unique training examples via softmax-sampling from top- $k$  (Eq. (16))
18:  Ingest  $n_{\text{ingest}}$  fresh windows into ring buffer; rebuild FAISS index

19:  // Stage 4: Q update (for previous transition, if  $k > 0$ )
20:   $H_k \leftarrow \text{DETACHEDHIDDEN}(\theta, \text{retrieved examples})$  ▷  $[R, d]$ 
21:  Compute  $V_k$  from  $Q_{\bar{\phi}}(H_k)$  as in Eq. (3)
22:  Update  $\phi$  to minimise  $\frac{1}{2}(Q_\phi(h_{\text{prev}}) - [(1 - \gamma)r_{\text{prev}} + \gamma V_k])^2$ 
23:   $\bar{\phi} \leftarrow \tau \phi + (1 - \tau) \bar{\phi}$ 

24:  // Stage 5: LM training
25:   $\theta \leftarrow \theta - \alpha \nabla_{\theta} \sum_{i=1}^R w_i \ell(\theta, x_i)$  ▷  $w_i = 1/R$  (uniform) or  $w_i = \pi_i$  (Q-weighted)

26:  // Stage 6: Reward
27:   $r_k \leftarrow \frac{1}{M} \log p_{\theta}(\hat{D}_k)$  ▷ Held-out evaluation
28: end for
```

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