

STABILIZING THE KUMARASWAMY DISTRIBUTION*

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Abstract. Large-scale latent variable models require expressive continuous distributions that support efficient sampling and low-variance differentiation, achievable through the reparameterization trick. The Kumaraswamy (KS) distribution is both expressive and supports the reparameterization trick with a simple closed-form inverse CDF. Yet, its adoption remains limited. We identify and resolve numerical instabilities in the log-pdf, CDF, and inverse CDF, exposing issues in libraries like PyTorch and TensorFlow. We then introduce simple and scalable latent variable models to improve exploration-exploitation trade-offs in contextual multi-armed bandits and enhance uncertainty quantification for link prediction with graph neural networks. We find these models to be most performant when paired with the stable KS. Our results support the stabilized KS distribution as a core component in scalable variational models for bounded latent variables.

Key words. Kumaraswamy Distribution, Catastrophic Cancellation, Latent Variable Models, Variational Inference, Multi-Armed Bandits

MSC codes. 65G50, 65C20, 62F15, 68T07, 62H12

1. Introduction. Probabilistic models use probability distributions as building blocks to model complex joint distributions between random variables. Such distributions can model unobserved ‘latent’ variables \mathbf{z} , or observed ‘data’ variables \mathbf{x} . Bounded interval-supported latent variables are central to many key applications, such as unobserved probabilities (e.g., user clicks in recommendation systems or links between network nodes), missing measurements in control systems (e.g., joint angles in $[0, 2\pi]$), and stochastic policies over bounded actions in reinforcement learning (e.g., motor torque in $[-10, 10]$).

To meet the demands of large-scale latent variable models, distributions supported on bounded intervals must satisfy the following criteria: (i) support the reparameterization trick through an explicit reparameterization function, such as a closed-form inverse CDF, enabling low-variance gradient estimation and efficient sampling; (ii) provide sufficient expressiveness to capture complex latent spaces; and (iii) offer simple distribution-related functions (log-pdf, explicit reparameterization function, and gradients) that allow fast and accurate evaluation. In Section 2.1, we argue that the Kumaraswamy (KS) distribution uniquely meets these criteria, yet remains surprisingly underused. In Section 3, we demonstrate that the KS distribution-related functions exhibit numerical instabilities concealed by standard parameterizations and exacerbated in large-scale latent variable models.

In this paper, we make the following technical contributions:

- We introduce an unconstrained logarithmic parameterization of the KS’s log-pdf, CDF, inverse CDF, and gradients, which isolate the dominant numerical instabilities, allowing application of recently developed stabilization techniques (Section 3).
- We propose the Variational Bandit Encoder (VBE), addressing exploration-exploitation trade-offs in contextual Bernoulli multi-armed bandits (Section 4.2).
- We propose the Variational Edge Encoder (VEE) for improved uncertainty quantification in link prediction with graph neural networks (Section 4.3).

The VBE and VEE are scalable latent variable models with bounded interval-supported

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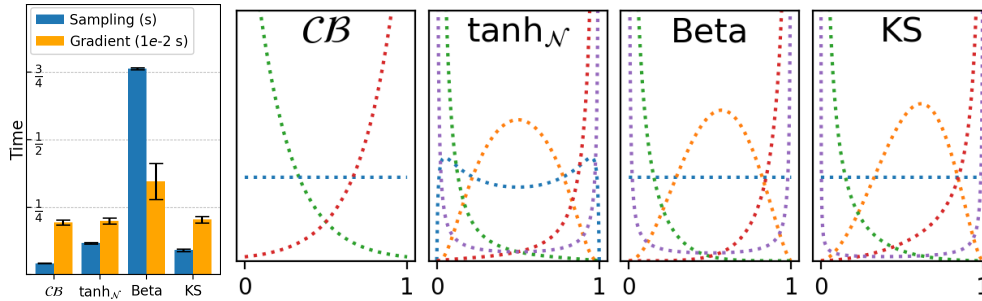


FIG. 1. Comparison of relevant bounded interval-supported distributions. Left: Time for sampling and differentiating through samples. The Beta lacks explicit reparameterization, and has slower sampling and gradients. Right: Expressiveness in terms of attainable prototypical shapes.

latent variables. Unlike traditional methods which tend to model global latent variables (Section 5.1), such as the parameters of a shared neural network (NN), the VBE and VEE define local latent variables per bandit arm or network link. This allows the models to incorporate prior knowledge precisely where domain expertise tends to reside—at a granular level, such as the expected reward of a specific arm or the probability of a particular link. Our numerical experiments demonstrate that both models perform best when paired with the stabilized KS distribution in their variational posterior, reinforcing its role as a core component in large-scale bounded latent variable modeling.

2. Background. The KS distribution [12, 16] has pdf $f(x) = abx^{a-1}(1-x)^{b-1}$, CDF $F(x) = 1 - (1-x)^b$, and inverse CDF $F^{-1}(u) = (1-u^{b^{-1}})^{a^{-1}}$, all defined for $x, u \in (0, 1)$ and parameterized by $a, b > 0$. The differential entropy of a KS with parameters a, b is

$$\begin{aligned} \mathcal{H}(\text{KS}) &:= - \int_0^1 f(x) \log f(x) dx \\ &= 1 - b + (1 - a) \left(\phi^{(0)}(b^{-1} + 1) + \gamma \right) - \log a - \log b, \end{aligned}$$

where $\phi^{(0)}$ is the digamma function and $\gamma \approx 0.577$ is the Euler-Mascheroni constant. The digamma function and its gradient, the trigamma function $\phi^{(1)}(x)$, can be represented as infinite series which converge rapidly and thus can be used effectively in numerical applications. They are included as standard functions in common auto-differentiation frameworks.

2.1. Continuous distributions with bounded interval support. Among distributions with bounded interval support, the KS uniquely satisfies desiderata (i)–(iii) in Section 1. It supports the reparameterization trick through its closed-form, differentiable inverse CDF, providing efficient sampling and low-variance gradients. The KS supports four distinct prototypical shapes — bell, U, increasing, and decreasing (Figure 1, right) — providing expressivity for diverse modeling tasks. Its log-pdf, CDF, and inverse CDF, along with their gradients, are composed only of affine transformations, exponentials, and logarithms, and can be parameterized directly in terms of unconstrained logarithmic values; see Section 3.3. This enables straightforward implementation with minimal dependencies and keeps most computation in log-space,

Property / Distributions.	\mathcal{CB}	$\tanh_{\mathcal{N}}$	Beta	KS
Expressiveness	low	high	high	high
Gradient Reparam.	explicit	explicit	implicit	explicit
Contains Uniform	✓	✗	✓	✓
Closed-form CDF	✓	✗	✗	✓
Closed-form inverse CDF	✓	✗	✗	✓
Numerical Issues	mild	high	low	low
Complex Functions	\tanh^{-1}	$\log(1 - \tanh^2(x))$	β, I	None
Parameterization	\mathbb{R}	\mathbb{R}^2	\mathbb{R}_+^2	\mathbb{R}^2
Analytical Moments	✓	✗	✓	✓
Closed-form KL	Exp. Family	$\tanh_{\mathcal{N}}$	Exp. Family	Beta
Entropy \mathcal{H}	✓	✗	✓	✓

TABLE 1

Comparison of bounded interval-supported distribution families.

enhancing stability and accuracy. The unconstrained logarithmic parameterization makes it well-suited for NNs, eliminating the need for positivity-enforcing link functions. Additionally, the KS has differentiable, closed-form expressions for moments, median, differential entropy $\mathcal{H}(\text{KS})$, and the Kullback-Leibler (KL) divergence to the Beta distribution, facilitating efficient incorporation of prior information.

We briefly introduce workhorse bounded-interval supported distribution families, namely the the Continuous Bernoulli, the Beta, and the tanh-squashed-Gaussian. The Continuous Bernoulli (\mathcal{CB}) [18] arises in deep learning for modeling continuous $[0, 1]$ -valued pixel intensities in natural images. It provides a normalized probabilistic counterpart to the commonly used binary cross-entropy loss, with density $p(x; \lambda) = C(\lambda)\lambda^x(1-\lambda)^{1-x}$, $x \in [0, 1]$, $\lambda \in (0, 1)$, where $C(\lambda) = \{2 \text{ if } \lambda = \frac{1}{2}, \text{ else } \frac{2 \tanh^{-1}(1-2\lambda)}{1-2\lambda}\}$ is the normalizing constant. The Beta distribution is a flexible two-parameter family, widely used for modeling probabilities and proportions. Its density, parameterized by $a, b > 0$, is given by: $p(x; a, b) = B(a, b)^{-1}x^{a-1}(1-x)^{b-1}$, $x \in (0, 1)$, where $B(a, b)$ is the Beta function. The tanh-squashed-Gaussian ($\tanh_{\mathcal{N}}$) maps Gaussian samples through the tanh function to produce outputs in $[-1, 1]$: $y = \tanh(z)$, $z \sim \mathcal{N}(\mu, \sigma^2)$. It is widely used in reinforcement learning over continuous bounded action spaces [8] due to its support for the reparameterization trick.

Table 1 compares these bounded-interval supported distribution families across important properties for latent variable modeling. *Expressiveness* measures the variety of prototypical shapes a distribution can represent. All distributions except \mathcal{CB} exhibit four prototypical shapes; \mathcal{CB} is limited to two. *Contains uniform* refers to the ability to represent the uniform distribution, critical for modeling complete uncertainty. All distributions except $\tanh_{\mathcal{N}}$ can express the uniform. *Closed-form CDF* indicates whether a closed-form CDF is available. Only \mathcal{CB} and KS provide such expressions. Similarly, *closed-form inverse CDF* indicates the availability of a closed-form inverse CDF, with only \mathcal{CB} and KS satisfying this criterion. *Numerical issues* capture challenges in stable evaluation. For example, \mathcal{CB} requires a Taylor expansion to handle singularities as $\lambda \rightarrow 0.5$. The $\tanh_{\mathcal{N}}$ distribution requires log-pdf clipping and parameter regularization to maintain stability, as appears in various implementations [8]. *Complex functions* highlight reliance on non-affine, non-logarithmic, or non-exponential operations. The $\tanh_{\mathcal{N}}$ involves computing $\log(1 - \tanh^2(x))$, which is numerically

unstable [2]. The Beta distribution relies on the Beta function and the regularized incomplete Beta function in its log-pdf and CDF, respectively, both requiring numerical approximations. In contrast, KS avoids such complexity in our novel parameterization (Section 3.3), computing a^{-1} as $\exp(-\log a)$ to sidestep division. In contrast, our novel parameterization of the KS distribution avoids complex functions; note a^{-1} is computed via $\exp(-\log a)$, avoiding division. *Parameterization* examines whether a distribution can be effectively expressed with unconstrained parameters. Both \mathcal{CB} (via $\log \lambda \in \mathbb{R}$) and $\tanh_{\mathcal{N}}$ (via $(\mu, \log \sigma) \in \mathbb{R}^2$) support unconstrained parameterization. We introduce the first unconstrained parameterization for KS in Section 3.3, using $(\log a, \log b) \in \mathbb{R}^2$. The Beta distribution, due to its dependence on the Beta function, resists effective unconstrained parameterization. *Closed-form KL functions* refer to analytical KL divergence expressions. The \mathcal{CB} and Beta distributions, as members of the exponential family, admit closed-form KL expressions with other exponential family members. The KS also has closed-form KL expressions with Beta family members, while $\tanh_{\mathcal{N}}$ is restricted to closed-form KL expressions within its own family. *Entropy* considers the availability of closed-form expressions for differential entropy. This property is present for all distributions except $\tanh_{\mathcal{N}}$.

2.2. Latent variable modeling with stochastic variational inference (SVI).

The primary method for fitting large-scale latent variable models is SVI [9]. Consider a model $p_{\theta}(\mathbf{x}) = \int p_{\theta}(\mathbf{x}|\mathbf{z})p(\mathbf{z})d\mathbf{z}$, where $\mathbf{x} \in \mathbb{R}^M$ is the observation, $\mathbf{z} \in \mathbb{R}^D$ is a vector-valued latent variable, $p_{\theta}(\mathbf{x}|\mathbf{z})$ is the likelihood function with parameters θ , and $p(\mathbf{z})$ is the prior distribution. Except for a few special cases, maximum likelihood learning in such models is intractable because of the difficulty of the integrals involved. Variational inference [10] provides a tractable alternative by introducing a variational posterior distribution $q_{\phi}(\mathbf{z})$ and maximizing a lower bound on the marginal log-likelihood called the ELBO:

$$(2.1) \quad \mathcal{L}(\mathbf{x}, \theta, \phi) = \mathbb{E}_{q_{\phi}(\mathbf{z})} [\log p_{\theta}(\mathbf{x}|\mathbf{z})] - D_{\text{KL}}(q_{\phi}(\mathbf{z}) \| p(\mathbf{z})) \leq \log p_{\theta}(\mathbf{x}).$$

Training models with modern SVI [14, 22] involves gradient-based optimization of this bound w.r.t. both the model parameters θ and the variational parameters ϕ . The first term in (2.1) encourages the model to assign high likelihood to the data, but its exact evaluation and gradients are typically intractable and so the expectation is often approximated with samples from $q_{\phi}(\mathbf{z})$. The KL divergence term incorporates prior information by penalizing deviations of the variational posterior from the prior $p(\mathbf{z})$. Closed-form expressions of $D_{\text{KL}}(q_{\phi}(\mathbf{z}) \| p(\mathbf{z}))$ allow efficient encoding of prior information; otherwise, sample-based approximations are required. In the common setting of i.i.d. data with per-datapoint latent variables, amortized inference introduces a shared NN, parameterized by ‘inference parameters’ ϕ , to map observations to variational parameters, approximating their individual posteriors as $q_{\phi}(\mathbf{z}|\mathbf{x})$. Modifying the ELBO by scaling the KL term with a parameter $\beta_{\text{KL}} > 0$ is often necessary to balance the trade-off between data likelihood and prior regularization [1]. We denote the sample-based approximation of this modified ELBO as $\hat{\mathcal{L}}_{\beta_{\text{KL}}}$.

2.3. Gradient reparameterization: explicit and implicit. A distribution $q_{\phi}(\mathbf{z})$ is said to be *explicitly* reparameterizable, or amenable to the ‘reparameterization trick’, if it can be expressed as a deterministic, differentiable transformation $\mathbf{z} = g(\epsilon, \phi)$ of a base distribution $\epsilon \sim p(\epsilon)$. This base distribution is typically simple, such as Uniform or standard Normal, enabling fast sample generation by first sampling from the base and then applying g . This enables the use of backpropagation to estimate

gradients of the form [cf. (2.1)]

$$(2.2) \quad \nabla_{\phi} \mathbb{E}_{q_{\phi}(\mathbf{z})}[f(\mathbf{z})] = \mathbb{E}_{p(\epsilon)}[\nabla_{\phi} f(g(\epsilon, \phi))] = \mathbb{E}_{p(\epsilon)}[\nabla_{\mathbf{z}} f(\mathbf{z})|_{\mathbf{z}=g(\epsilon, \phi)} \nabla_{\phi} g(\epsilon, \phi)],$$

an expectation with form encompassing the ELBO. Explicit reparameterization is compatible with distributions in the location-scale family (e.g., Gaussian, Laplace, Cauchy), distributions with tractable inverse CDFs (e.g., exponential, KS, \mathcal{CB}), or those expressible as deterministic transformations of such distributions (e.g., $\tanh_{\mathcal{N}}$). When explicit reparameterization is not available, implicit reparameterization [6] is commonly used for distributions with numerically tractable CDFs, such as truncated, mixture, Gamma, Beta, Dirichlet, or von Mises distributions. This method expresses the parameter gradient through the sample $\nabla_{\phi} \mathbf{z}$ as a function only of the CDF gradients, not its inverse. Such CDF gradients are either found analytically (if feasible) or more commonly using numerical methods, e.g., forward mode auto-differentiation on CDF estimates, as in the Gamma and Beta distributions. Without explicit reparameterization, sampling and gradient computations tend to be slower and more complex, and produce higher-variance estimates of (2.2), reducing learning efficiency and stability [14, 11].

3. Stabilizing the Kumaraswamy. The KS distribution’s utility relies on stable computation of its log-pdf, CDF, inverse CDF, and their gradients. In the standard parameterization, these functions contain instabilities from hidden $\log(1 - \exp(x))$ terms. We address this by introducing an unconstrained logarithmic parameterization that isolates these unstable terms, enabling their straightforward replacement with the stable `log1mexp` function. Finally, we show why naive stabilization techniques, such as parameter clipping, fail in high-dimensional applications.

3.1. Identifying the instability: $\log(1 - \exp(x))$. Naive computation of $\log(1 - \exp(x))$ for $x < 0$ leads to significant numerical errors as x approaches 0 (Figure 2, red). These errors grow so large that they can cause *numerical instability*, i.e., an irrecoverable error such as `-inf`. These errors result from *catastrophic cancellation*, which occurs when subtracting nearly equal numbers — here, $1 - \exp(x)$. As $x \rightarrow 0$, $\exp(x) \approx 1$, so $1 - \exp(x)$ results in the cancellation of leading significant bits, leaving only a few less significant, less accurate bits to represent the result. This causes large relative errors in $1 - \exp(x)$, which are amplified when input to the logarithm as its magnitude grows sharply near zero. If the cancellation is complete, $1 - \exp(x)$ underflows to 0 and the logarithm returns `-inf`, as seen in Figure 2 (red) when $\log_2 |x| < -24$.

3.2. Numerical building blocks for accurate $\log(1 - \exp(x))$ computation. When $|x| \ll 1$, both $\log(1 + x)$ and $\exp(x) - 1$ can suffer from severe cancellation: the former between 1 and x , the latter between $\exp(x)$ and -1 . In both cases, a simple solution for accurate computation is to use a few terms of the Taylor series, as

$$\begin{aligned} \text{log1p}(x) &:= \log(1 + x) = x - \frac{x^2}{2} + \frac{x^3}{3} - \dots, \quad \text{for } |x| < 1, \\ \text{expm1}(x) &:= \exp(x) - 1 = x + \frac{x^2}{2!} + \frac{x^3}{3!} + \dots, \quad \text{for } |x| < 1, \end{aligned}$$

where $n!$ denotes the factorial. These functions form the basis for two common methods to compute $\log(1 - \exp(x))$: `log(-expm1(x))` and `log1p(-exp(x))`. [19] showed neither method provides sufficient accuracy across the domain. However, each

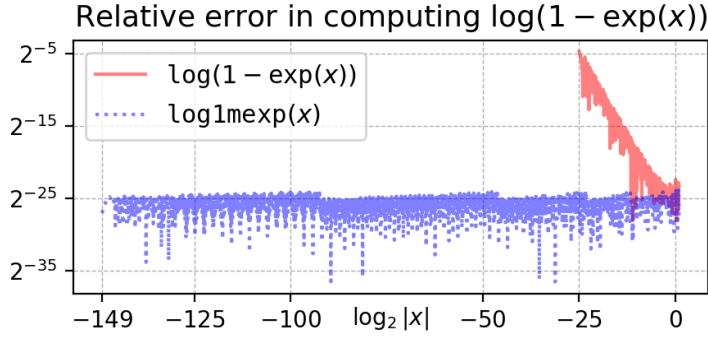


FIG. 2. Naive computation of $\log(1 - \exp(x))$ (red) becomes unstable as $x \rightarrow 0$ due to catastrophic cancellation, while $\log1mexp(x)$ (blue) ensures accurate computation.

approach is accurate in complementary regions, leading to

$$(3.1) \quad \log1mexp(x) := \begin{cases} \log(-\expm1(x)) & -\log 2 \leq x < 0 \\ \log1p(-\exp(x)) & x < -\log 2, \end{cases}$$

which computes $\log(1 - \exp(x))$ accurately throughout single precision, shown in Figure 2 (blue).

3.3. A stable Kumaraswamy. The direct implementation of the KS's log-pdf and inverse CDF — as found in all core auto-differentiation libraries — produces numerical instabilities. Here, we introduce a novel parameterization in terms of unconstrained logarithmic parameter values, which isolates and makes explicit the unstable terms

$$\begin{aligned} w_{b^{-1}}(u) &= \log(1 - u^{b^{-1}}) = \log(1 - \exp(b^{-1} \log u)) \\ w_a(x) &= \log(1 - x^a) = \log(1 - \exp(a \log x)), \end{aligned}$$

eliminates the need for positivity-enforcing link functions, and whose expressions involve only affine, exponential, and logarithmic transformations. This allows the log-pdf and its gradients to be expressed as

$$(3.2) \quad \log f(x) = \log a + \log b + (a - 1) \log x + (b - 1) w_a(x)$$

$$(3.3) \quad \nabla_{\log x} \log f(x) = (a - 1) - (b - 1) \cdot \exp(a \log x - w_a(x)) + \log a$$

$$(3.4) \quad \nabla_{\log a} \log f(x) = 1 + a \log x \cdot \{1 - (b - 1) \cdot \exp(a \log x - w_a(x))\}$$

$$(3.5) \quad \nabla_{\log b} \log f(x) = 1 + b \cdot w_a(x).$$

Likewise for the CDF

$$(3.6) \quad F(x) = 1 - (1 - x^a)^b = 1 - \exp(b \cdot w_a(x))$$

$$(3.7) \quad \nabla_x F(x) = \exp(\log a + \log b + (a - 1) \cdot \log x + (b - 1) \cdot w_a(x))$$

$$(3.8) \quad \nabla_{\log a} F(x) = \exp(\log a + \log b + a \cdot \log x + (b - 1) \cdot w_a(x)) \cdot \log x$$

$$(3.9) \quad \nabla_{\log b} F(x) = \exp(\log b + b \cdot w_a(x)) \cdot (-w_a(x)),$$

and the inverse CDF

$$(3.10) \quad F^{-1}(u) = (1 - u^{b^{-1}})^{a^{-1}} = \exp(a^{-1} w_{b^{-1}}(u))$$

$$(3.11) \quad \nabla_{\log a} F^{-1}(u) = \exp(-\log a + a^{-1} w_{b^{-1}}(u)) \cdot (-w_{b^{-1}}(u))$$

$$(3.12) \quad \nabla_{\log b} F^{-1}(u) = \exp(-\log a - \log b + b^{-1} \log u + (a^{-1} - 1) w_{b^{-1}}(u)) \cdot \log u.$$

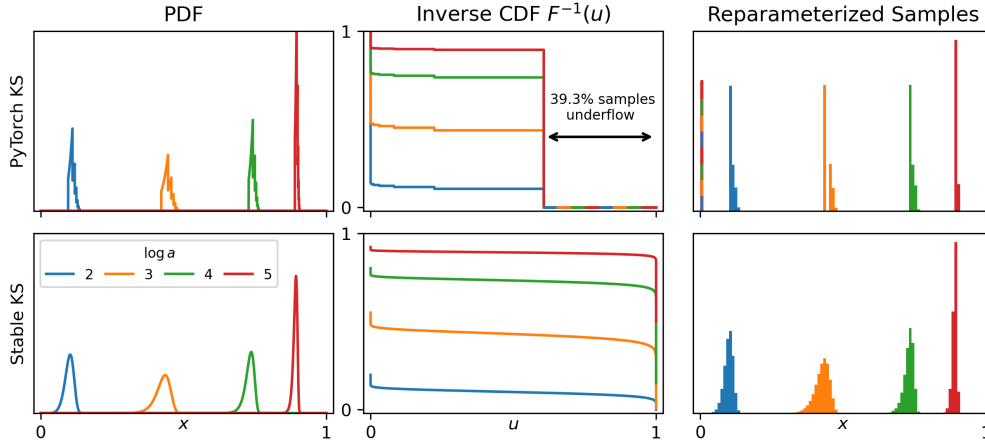


FIG. 3. Stabilizing $\log(1 - \exp(x))$ terms eliminates numerical instabilities in the KS log-pdf and inverse CDF. We compare the unstable PyTorch KS implementation (top row) and our stable KS (bottom row) for realistic KS distributions ($\log_2 b = 24$, varying a). Catastrophic cancellation in the $\log(1 - \exp(x))$ terms in the PyTorch KS causes jagged curves and inverse CDF underflow beyond $u \approx 1 - 39.3$, resulting in a point mass of ≈ 39.3 at $x = 0$ in the sampling distribution. Our stable KS removes the instability by using `log1mexp`.

This parameterization’s algebraic form allows direct replacement of the dominant unstable terms, substituting $w_{b^{-1}}(u)$ with $\text{log1mexp}(b^{-1} \log u)$ and $w_a(x)$ with $\text{log1mexp}(a \log x)$. Access to $\log a$ and $\log b$ avoids errors from unnecessary transitions in-and-out of log-space. We also avoid the error prone expressions produced in backpropogation’s direct application of the chain rule, e.g.,

$$\begin{aligned} \nabla_{\log b} F^{-1} &= \frac{1}{a} \cdot \exp\left(\frac{1}{a} \log\left(1 - \exp\left(\frac{1}{b} \log u\right)\right)\right) \\ &\quad \cdot \left(1 - \exp\left(\frac{1}{b} \log u\right)\right)^{-1} \cdot \exp\left(\frac{1}{b} \log u\right) \cdot \log u \cdot \frac{-1}{b^2} \cdot b \end{aligned}$$

and (3.12) are equivalent expressions for $\nabla_{\log b} F^{-1}$, but their computed values can differ greatly for extreme parameter values. Desirable KS distributions can obtain such problematic extreme parameter values, e.g., the KS distributions in Figure 3 have $b \approx 10^6$. See Section 3.4 for further discussion on how instability in the unmodified KS can worsen with increasing evidence.

Figure 3 compares the PDF, inverse CDF, and histograms of reparameterized samples for KS distributions which are typical to real-world modeling scenarios. The PyTorch implementation (top row) shows jaggedness in both the PDF and inverse CDF, caused by catastrophic cancellation in the unstable terms $w_a(x)$ and $w_{b^{-1}}(u)$. Additionally, the PyTorch inverse CDF underflows beyond $u \approx 1 - 39.3$: here, $w_{b^{-1}}(u) = -\infty$, and $F^{-1}(u) = \exp(a^{-1} \cdot -\infty) = 0$. This underflow results in a point mass at $x = 0$ (a point outside of the KS support) with probability ≈ 39.3 in each of the reparameterized sampling distributions, and produces infinite gradients via $\nabla_{\log a} F^{-1} = \infty$ [cf. (3.11)]. This infinite gradient triggers a cascade: infinite parameter values after the optimizer step and NaN activations in the next forward pass, which is what users ultimately observe when training fails.

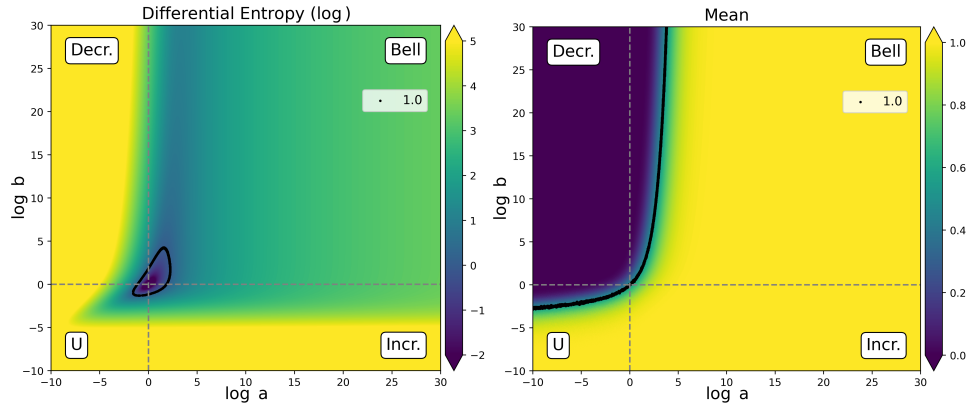


FIG. 4. Differential entropy and mean of Kumaraswamy distributions across a wide range of parameter values. Low-entropy distributions are concentrated near the origin, where $\log a = \log b = 1$ corresponds to the uniform distribution. Distributions with a log differential entropy of 1 and mean of 0.5 are marked in black. Small changes in parameter magnitudes rapidly yield extremely high-entropy distributions — essentially delta functions at 0 or 1 — except in the narrow region around the black curve representing distributions with mean 0.5.

3.4. Counter intuitive stability properties of the unstable Kumaraswamy.

When using the unstable KS to model latent variables with SVI, the instability of the KS distribution can *paradoxically worsen as evidence increases*. Here, evidence refers to observed data that sharpens the posterior distribution and reduces uncertainty. Representing sharper, high-entropy bell-shaped KS distributions — indicative of reduced posterior uncertainty — requires extremely large b values. Figure 4 illustrates this: a bell-shaped KS distribution with mean 0.5 and differential entropy $\mathcal{H} \approx \exp(2)$ necessitates $\log b = 24$, and thus $b = \exp(24)$ in the unstable KS implementation which lacks logarithmic parameterization; see Figure 3 for examples of such moderate entropy distributions with $\log b = 24$. SVI will leverage the inverse CDF and its gradient expressions (3.10)–(3.12), which critically depend on b through the term $w_{b-1}(u) = \log(1 - \exp(b^{-1} \log u))$. Large b values will act to worsen instability by driving $\exp(b^{-1} \log u)$ closer to 1, increasing the risk of catastrophic cancellation. We believe this counter-intuitive behavior likely frustrated modelers, but is no longer an issue in the stabilized KS.

As an illustrative example, consider modeling the latent probability of heads in a Bernoulli coin-flipping experiment using a KS distribution as the variational posterior, where the true probability of heads is 0.5. With a uniform prior and a small number of observed flips, the posterior is well-approximated by a mild-entropy, bell-shaped KS distribution, characterized by low-magnitude parameters $a, b > 1$. In this regime, b^{-1} remains sufficiently far from zero, minimizing the risk of catastrophic cancellation in the term $1 - \exp(b^{-1} \log u)$, as $\exp(b^{-1} \log u)$ stays safely away from 1. However, as the number of observed flips increases, the posterior sharpens to reflect reduced uncertainty, demanding larger values of b to represent the corresponding higher entropy KS distribution. This drives $\exp(b^{-1} \log u)$ closer to 1, increasing the risk of catastrophic cancellation and numerical instability.

3.5. The inadequacy of parameter clipping in large-scale settings. Numerical instability in the KS is inherently stochastic, and in high-dimensional settings, the compounded probability of failure across multiple variables makes program failure

almost certain. As the program goes unstable if any single KS goes unstable, the overall instability probability can be modeled as the probability of at least one failure in D independent Bernoulli trials: $1 - p(\text{KS stable})^D$, for D KS latent variables. In practical large-scale settings, e.g., recommendation systems with 10^7 users and $D = 10^7$ recommendation items, the probability of instability approaches 1 across all reasonable parameter clipping values, rendering clipping an ineffective stabilization strategy.

Quantitative illustration. Consider the stochastic instability arising from the term $\log(1 - \exp(b^{-1} \log u))$, where catastrophic cancellation occurs if $1 - \exp(b^{-1} \log u)$ becomes too small. To avoid logarithmic domain errors in single precision, we enforce $-b^{-1} \log u > 2^{-24}$ (Figure 2). We aim to select a b_{\max} to satisfy this constraint: a larger b_{\max} expands the variational family allowing improved posterior approximation, but worsens stability. Consider the moderate entropy KS distributions in Figure 3 which use $b = 2^{24}$. Using $b_{\max} = 2^{24}$, only $u < 0.6321$ satisfies the stability condition, i.e., $p(\text{KS stable}) \approx 0.6321$ per sample. With $D = 10^7$, the overall probability of instability becomes $1 - 0.6321^{10^7} \approx 1$. Now consider aggressively restricting $b_{\max} = 2^4$, as done in [20]. Now $u < 0.9999$ satisfies the stability condition. Even then, introducing $D = 10^7$ variables, we still have the overall probability of instability is $1 - 0.9999^{10^7} \approx 1$. Thus, even extreme clipping fails to stabilize KS distributions at scale. Further, this analysis considers only a single posterior sample. In practice, training with SVI requires $T \sim 10^3$ optimization steps, each requiring posterior samples for gradient estimation. This compounds the instability probability to $1 - p(\text{KS stable})^{TD}$, making clipping ineffective in realistic large-scale scenarios.

4. Experiments and New Variational Architectures. Using the well established Variational Auto-Encoder (VAE) framework on MNIST and CIFAR-10 datasets, we show that the stabilized KS enables reliable training as both a variational posterior [Eqns. (3.10)–(3.12)] and likelihood function [Eqns. (3.2)–(3.5)]. We then introduce two new deep variational architectures that leverage bounded interval-supported latent variables: the Variational Bandit Encoder (VBE) for improving exploration-exploitation trade-offs in contextual multi-armed bandits (Section 4.2), and the Variational Edge Encoder (VEE) for enhancing uncertainty quantification in link-prediction with graph neural networks (Section 4.3). These novel architectures tend to be most performant when using the KS as their variational posterior. Across the experimental domains, our stable KS tends to be more performant and easier to use than alternative variational distributions supported on bounded intervals. For instance, $\tanh_{\mathcal{N}}$ models require log-pdf clipping for stability, while Beta models show significant performance variability based on the chosen positivity-enforcing link function and often fail to converge, e.g., on CIFAR-10 in Section 4.1. Finally, our new variational models are fast: the VBEs in Section 4.2 are $8 - 22\times$ faster than the state-of-the-art baseline.

Remark 4.1. **Across all three experimental settings, models using the unstable KS produce NaN errors in training and are therefore excluded.** Prior work using the KS in low-dimensional latent variable models [20, 21, 24] similarly find NaN errors, and rely on parameter clipping to avoid instability. See Section 3.5 for why this is approach does not work in large-scale settings. Our stabilization approach directly resolves these numerical issues, enabling stable training at scale.

4.1. Image variational auto-encoders. The VAE [14] is a generative latent variable model trained using amortized variational inference. Both the variational posterior $q_{\phi}(\mathbf{z}|\mathbf{x})$ and the conditional likelihood $p_{\theta}(\mathbf{z}|\mathbf{x})$ are parameterized using NNs,

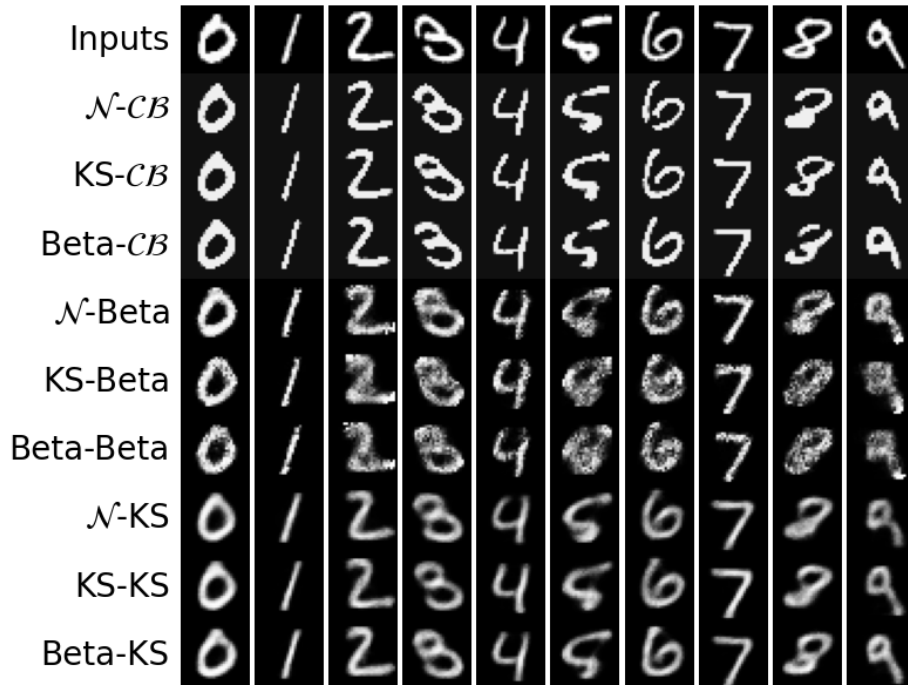


FIG. 5. MNIST test digit VAE reconstructions.

known as the encoder $e_\phi(\mathbf{x}) : \mathbb{R}^M \mapsto \mathbb{R}^D$ and decoder $d_\theta(\mathbf{z}) : \mathbb{R}^D \mapsto \mathbb{R}^M$, respectively. VAEs typically use the standard Normal distribution as the prior and a factorized Normal as the variational posterior. The use of alternative variational distributions allows incorporating different prior assumptions about the latent factors of the data, such as bounded support or periodicity [6].

Experimental setup and metrics. Inspired by [18], we train VAEs with fully factorized priors and variational posteriors on MNIST and CIFAR-10 without pixel binarization, using an unmodified ELBO ($\beta_{\text{KL}} = 1$). We adopt the most effective likelihoods from their work (Beta and CB), identical latent dimension D (MNIST: $D = 20$, CIFAR-10: $D = 50$), and the same standard NN architectures, which are detailed in Appendix A, along with the training hyperparameters. For each variational posterior factor, we choose the canonical prior: $\mathcal{N}_{(0,1)}$ for \mathcal{N} , and $U_{(0,1)}$ for KS and Beta. We evaluate the models using a single sample approximation of the test ELBO. To assess usefulness of the learned latent representations, we encode test data \mathbf{x}_n , compute the mean $\mathbb{E}[q_\phi(\mathbf{z}_n|\mathbf{x}_n)]$, and classify the test labels using a 15-nearest neighbor classifier; the classifier accuracy (%) is denoted $\mathcal{K}(\phi)$. For subjective evaluation, we display the mean decoded likelihood of a single sample from the encoded posterior of random test digits in Figure 5.

Discussion of results. The sole purpose of this experiment is to provide evidence toward the stabilization of the KS. Notably, stable KS VAEs maintain numerical stability while all VAEs with the unstable KS produce unstable training. VAEs with Beta-distributed variational posteriors often do not converge; indeed, [6] reported strong performance on binarized MNIST using a softplus link function, but did not present results on CIFAR-10, nor could we find other works that did. We suspect this is due to similar instability issues, with the higher gradient variance of the Beta’s implicit

TABLE 2
VAE on MNIST and CIFAR-10.

Prior	$q_\phi(\mathbf{z} \mathbf{x})$	$p_\theta(\mathbf{x} \mathbf{z})$	MNIST		CIFAR-10	
			ELBO	$\mathcal{K}(\phi)$	ELBO	$\mathcal{K}(\phi)$
$\mathcal{N}_{(0,1)}$	\mathcal{N}	\mathcal{CB}	1825 ± 98	97.3	1167 ± 901	37.9
$\mathbf{U}_{(0,1)}$	KS	\mathcal{CB}	1818 ± 104	97.4	1172 ± 908	41.5
$\mathbf{U}_{(0,1)}$	Beta	\mathcal{CB}	1821 ± 98	97.5	1167 ± 907	40.3
$\mathcal{N}_{(0,1)}$	\mathcal{N}	Beta	4073 ± 5701	92.1	3566 ± 1203	48.5
$\mathbf{U}_{(0,1)}$	KS	Beta	4061 ± 1932	91.3	3483 ± 1133	50.1
$\mathbf{U}_{(0,1)}$	Beta	Beta	4082 ± 1522	90.1	N/A	N/A
$\mathcal{N}_{(0,1)}$	\mathcal{N}	KS	3328 ± 989	96.4	1720 ± 884	47.1
$\mathbf{U}_{(0,1)}$	KS	KS	3355 ± 512	96.8	1738 ± 877	48.8
$\mathbf{U}_{(0,1)}$	Beta	KS	3348 ± 515	97.1	N/A	N/A

reparameterization a likely explanation. In an attempt to overcome this instability in Beta VAEs we report the best metrics across softplus or exp link functions in Table 2. When neither converges, we report N/A. The results in Table 2 show that across datasets, VAEs with KS-distributed variational posteriors consistently produce useful latent spaces, evidenced by strong $\mathcal{K}(\phi)$, and yield reconstructions with high ELBOs and visual quality.

When paired with any variational posterior, a KS likelihood yields stronger MNIST reconstructions than Beta likelihoods: compare rows *-Beta to *-KS in Table 5. As in [18], we find \mathcal{CB} likelihoods produce the most subjectively performant VAEs on MNIST, unsurprising as \mathcal{CB} was introduced specifically for the approximately binary MNIST pixel data.

4.2. Contextual Bernoulli multi-armed bandits. The contextual Bernoulli multi-armed bandit (MAB) problem involves a decision maker who, at each time step $t = 1, \dots, T$, selects one arm from a finite set of K options. Each arm has an associated context $\mathbf{x}_k \in \mathbb{R}^d$, and pulling an arm yields a binary reward $r_k \sim \text{Ber}(v_k)$, where $v_k \in [0, 1]$ is the unknown mean reward. MABs originate by analogy to casino slot machines, where each machine (arm) has a different payout rate, and the challenge lies in deciding which arms to pull in order to maximize total winnings while learning about their payout rates, a situation called the exploration-exploitation dilemma. MABs have found applications in modern recommendation systems [17], clinical trials design [28], and mobile health [25]. Thompson Sampling (TS) is a simple, empirically effective [3], and scalable [13] arm selection heuristic. It selects the arm corresponding to the highest value drawn from the posterior distributions over the latent z_k 's. This approach naturally balances exploration and exploitation: the uncertainty in the posteriors promote exploration, while concentration of probability mass on large mean rewards drive exploitation.

Variational Bandit Encoder (VBE): VAE \cap TS. Our novel VBE posits a fully factorized KS variational posterior $\prod_k q_\phi(z_k|\mathbf{x}_k)$, prior $p(\mathbf{z}) = U_{(0,1)}^K$, and a Bernoulli reward likelihood $p(r|v_k)$ for each arm. Similar to VAEs, we employ amortized inference using a shared NN encoder $e_\phi(\mathbf{x}_k)$, which defines a reparameterizable variational distribution $q_\phi(z_k|\mathbf{x}_k)$. However, unlike VAEs, VBES omit the decoder; samples $\tilde{z}_k \sim q_\phi(z_k|\mathbf{x}_k)$ directly parameterize the reward likelihood. The arm selection at step t follows TS: $a = \arg\max_k \{\tilde{z}_k\}$. We then draw reward $r \sim \text{Ber}(v_a)$ and record it in

Algorithm 4.1 Variational Bandit Encoder (VBE)**Require:** $\{\mathbf{x}_k\}_{k=1}^K, \{v_k\}_{k=1}^K, \eta, \beta_{\text{KL}}$

- 1: Variational posterior $q \leftarrow \text{KS}$
- 2: Replay buffer $\mathcal{D} \leftarrow \emptyset$
- 3: **for** $t = 1 \dots T$ **do**
- 4: Encode: $(a_k, b_k) = e_\phi(\mathbf{x}_k)$
- 5: Sample: $\tilde{z}_k \sim q(z_k; a_k, b_k)$
- 6: TS: $a = \operatorname{argmax}_k \{\tilde{z}_k\}$
- 7: Reward: $r \sim \text{Ber}(v_a)$
- 8: $\mathcal{D} \leftarrow \mathcal{D} \cup \{(\mathbf{x}_a, a, r)\}$
- 9: Construct $\hat{\mathcal{L}}_{\beta_{\text{KL}}, t}$ as in (4.1)
- 10: $\phi \leftarrow \phi + \eta \nabla_\phi \hat{\mathcal{L}}_{\beta_{\text{KL}}, t}$
- 11: **end for**

the replay buffer $\mathcal{D} \leftarrow \mathcal{D} \cup \{(\mathbf{x}_a, a, r)\}$. We construct a sample approximation of the modified ELBO over the subset of arms $\mathcal{K}_t \subset \{1, \dots, K\}$ that have been pulled by time t as

$$(4.1) \quad \hat{\mathcal{L}}_{\beta_{\text{KL}}, t}(\mathcal{D}, \phi) = \sum_{(\mathbf{x}_a, a, r) \in \mathcal{D}} \log p(r|\tilde{z}_a) + \beta_{\text{KL}} \sum_{k \in \mathcal{K}_t} \mathcal{H}[q_\phi(z_k|\mathbf{x}_k)].$$

The second term promotes exploration by penalizing overconfidence with the exploration effect proportional to β_{KL} . We maximize (4.1) w.r.t. ϕ via gradient ascent, enabled by the reparameterizable KS. VBE execution is summarized in Algorithm 4.1.

VBE Modified ELBO Derivation. Let $\mathbf{X} = [\mathbf{x}_1, \dots, \mathbf{x}_K]$ be a matrix where the k -th column corresponds to the context feature \mathbf{x}_k . Assuming independence between arms and within-arm rewards, the data likelihood can be factorized as $p(\mathcal{D}|\mathbf{z}) = \prod_{(\mathbf{x}_a, a, r) \in \mathcal{D}} p(r|z_a)$. We adopt a fully factorized variational posterior of the form $q_\phi(\mathbf{z}|\mathbf{X}) = \prod_{k=1}^K q_\phi(z_k|\mathbf{x}_k)$. Recall that $\mathcal{K}_t \subset \{1, \dots, K\}$ represents the subset of arms that have been pulled, and thus for which we have reward data. The modified ELBO is derived as follows:

$$\begin{aligned} \mathcal{L}_{\beta_{\text{KL}}, t}(\mathcal{D}, \phi) &= \mathbb{E}_{q_\phi(\mathbf{z}|\mathbf{X})} [\log p(\mathcal{D}|\mathbf{z})] - \beta_{\text{KL}} \text{KL}(q_\phi(\mathbf{z}|\mathbf{X}) \| p(\mathbf{z})) \\ &= \mathbb{E}_{q_\phi(\mathbf{z}|\mathbf{X})} [\log p(\mathcal{D}|\mathbf{z})] + \beta_{\text{KL}} \mathcal{H}[q_\phi(\mathbf{z}|\mathbf{X})], \quad p(\mathbf{z}) = U_{(0,1)}^K \\ &= \mathbb{E}_{q_\phi(\mathbf{z}|\mathbf{X})} [\log p(\mathcal{D}|\mathbf{z})] + \beta_{\text{KL}} \sum_{k \in \mathcal{K}_t} \mathcal{H}[q_\phi(z_a|\mathbf{x}_a)] \\ &= \mathbb{E}_{q_\phi(\mathbf{z}|\mathbf{X})} \left[\sum_{(\mathbf{x}_a, a, r) \in \mathcal{D}} \log p(r|z_a) \right] + \beta_{\text{KL}} \sum_{k \in \mathcal{K}_t} \mathcal{H}[q_\phi(z_a|\mathbf{x}_a)] \\ &\approx \sum_{(\mathbf{x}_a, a, r) \in \mathcal{D}} \log p(r|\tilde{z}_a) + \beta_{\text{KL}} \sum_{k \in \mathcal{K}_t} \mathcal{H}[q_\phi(z_a|\mathbf{x}_a)], \quad \tilde{z}_a \sim q_\phi(z_a|\mathbf{x}_a) \end{aligned}$$

where in the final step, we use a single sample approximation of the expectation.

VBE advantages. VBES provide four primary advantages over alternative TS-based Bernoulli MAB approaches, discussed in Section 5.1

- *Scalability and Compatibility.* VBE training consists of a forward pass through a NN, sampling an explicitly reparameterized distribution, and a backward pass for gradient-based updates. This process is scalable and fully compatible with existing gradient-based infrastructure.

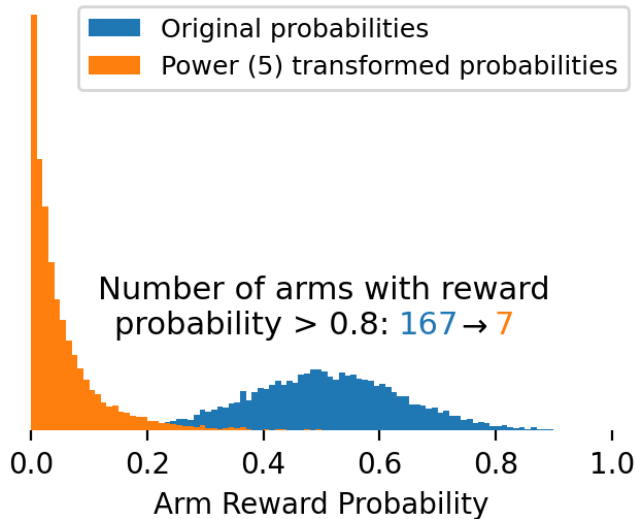


FIG. 6. High arm reward probabilities are reduced via a power 5 exponentiation, challenging agents to explore.

- *Prior Knowledge Incorporation.* When prior knowledge exists on an arm k it can be efficiently encoded as $p(z_k) = \text{Beta}(a_k, b_k)$, replacing $\mathcal{H}[q_\phi(z_k|\mathbf{x}_k)]$ with $-D_{\text{KL}}(q_\phi(z_k|\mathbf{x}_k) \| p(z_k))$.
- *Interpretability and Independence.* Encoding \mathbf{x}_k produces KS distribution parameters, fully encapsulating the model’s beliefs about v_k . This is independent of other arms and past data.
- *Simplicity.* VBEs lack numerous hyperparameters and complex architectural components.

Alternative methods lack some or all of these properties because they do not directly model the mean rewards nor differentiate through mean reward samples; instead, they model the parameters ϕ .

Experimental setup. We construct synthetic data with $K = 10^4$ arms by first sampling a weight vector \mathbf{w} and features $\{\mathbf{x}_k\}_{k=1}^K$ from $\mathcal{N}(\mathbf{0}, \mathbf{I}_5)$. We then compute $\{\mathbf{w}^\top \mathbf{x}_k\}_{k=1}^K$ and apply min-max normalization to produce probabilities (referred to as “Original probabilities” in Figure 6). To introduce non-linearity, we raise these probabilities to the power 5 (shown as “Power (5) transformed probabilities” in Figure 6). Exponentiating the probabilities not only makes the mapping from features to mean rewards more challenging to learn, but it also significantly reduces the number of arms with high probabilities, forcing the agent to explore more. For instance, when raising the probabilities to the power of 5, the number of arms with large probabilities drops from 167 to just 7. We consider $T = 2 \cdot 10^3$ steps.

We evaluate VBEs with either a KS (VBE-KS), Beta (VBE-Beta), or $\tanh_{\mathcal{N}}$ (VBE- $\tanh_{\mathcal{N}}$) all using $\beta_{\text{KL}} = |\mathcal{K}_t|^{-1}$, which makes the second term in (4.1) a mean. VBE- $\tanh_{\mathcal{N}}$ ’s performance is sensitive to the number of samples used in its entropy estimate: we found degraded performance beyond 10 samples. The learning rate is set to $\eta = 10^{-2}$. As a baseline, we use LMC-TS, which employs Langevin Monte Carlo (LMC) to sample posterior parameters of a NN, known for state-of-the-art performance across various tasks [31]. All models use an MLP with 3 hidden layers of width 32. LMC-TS hyperparameters (inverse temperature, LMC steps, weight decay) are set or

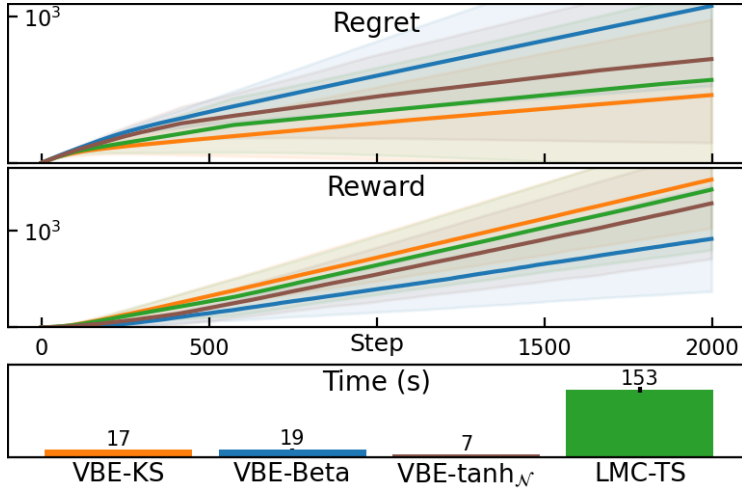


FIG. 7. Synthetic bandit performance over 5 runs. VBE-KS best handles exploration-exploitation trade-offs while being $\sim 10\times$ faster than state-of-the-art Langevin sampling based LMC-TS.

tuned based on the authors’ code. We repeat experiments 5 times on an Apple M2 CPU and report the mean and standard deviation across these runs in Figure 7.

Metrics and evaluation. The optimal policy always selects the arm with the highest mean reward r^* . Our objective is to minimize regret, defined as the cumulative difference between the expected reward of the chosen action and the optimal action (accessible in the synthetic setting), i.e., $\sum_{t=1}^T (r^* - r_{a_t})$. VBE-KS achieves lower regret and higher cumulative reward than all baselines. VBE-Beta performs significantly worse than VBE-KS and VBE-tanh $_N$, highlighting the importance of explicit reparameterization. LMC-TS is performant — worse than VBE-KS and better than VBE-tanh $_N$ — but is 8–22 \times slower than VBES: VBES avoid the computational overhead of LMC.

4.3. Variational link prediction with Graph Neural Networks. Graph Neural Networks (GNNs) have become a powerful tool for learning from graph-structured data, with applications in critical areas like drug discovery [32] and finance [29]. A key task is link prediction, where the goal is to infer unobserved edges between nodes. However, real-world deployment of graph learning models is often hindered by a lack of reliable uncertainty estimates and limited capacity to incorporate prior knowledge [30]. To address these challenges, we propose a variational approach where the GNN encodes a KS to model the unobserved probabilities of each network link’s existence, enabling uncertainty quantification and prior knowledge integration with minimal computational overhead.

In a typical link prediction setup, the GNN has access to the features $\mathbf{X} \in \mathbb{R}^{N \times d}$ of all N nodes, but only a subset of positive edges in the training \mathcal{D}_{tr} and validation \mathcal{D}_{val} sets. Edges are labeled as 1 (present) or 0 (absent). The GNN generates edge embeddings through message passing and neighborhood aggregation, outputting probabilities $z_{u,v} \in (0, 1)$ that parameterize a Bernoulli likelihood. The seminal work of [15] proposed Variational Graph Auto-encoders (VGAEs), which posits a Gaussian variational posterior over the final *node* embeddings. When used for link prediction it samples final node embeddings from the variational posterior and decodes them to produce edge probabilities. In contrast, our approach directly

models the probability of an edge using the KS. An advantage of directly modeling edge probabilities is interpretability; deep nodal embeddings are often difficult to interpret, and priors are typically selected for computational tractability rather than their ability to incorporate meaningful prior information. However, the probability of an edge (u, v) existing between two nodes is an interpretable quantity that can often be informed by domain expertise. For example, in gene regulatory networks, epidemiological networks, and social networks experts often have prior knowledge about the likelihood of specific interactions, transmissions, or friendships, respectively, which can be directly incorporated into edge prior $p(z_{(u,v)})$. We believe the limited exploration of variational modeling for edge probabilities is due to the previous lack of an expressive, stable, explicitly reparameterizable bounded-interval distributions.

Variational Edge Encoder (VEE). We propose a fully factorized KS variational posterior $\prod_{(u,v) \in \mathcal{D}_{tr}} q_\phi(z_{u,v} | \mathbf{X}, \mathcal{D}_{tr})$ with a uniform prior $p(\mathbf{z}) = U_{(0,1)}^{|\mathcal{D}_{tr}|}$. The GNN encoder e_ϕ parameterizes a KS distribution for each possible edge (u, v) . The remaining structure is highly similar to VBEs: a single sample $\tilde{z}_{u,v} \sim q_\phi(z_{u,v} | \mathbf{X}, \mathcal{D}_{tr})$ directly parameterizes the Bernoulli likelihood, and we maximize a sample approximation of the modified ELBO

$$(4.2) \quad \hat{\mathcal{L}}_{\beta_{\text{KL}}}((\mathbf{X}, \mathcal{D}_{tr}), \phi) = \sum_{(u,v) \in \mathcal{D}_{tr}} \log p(z_{(u,v)} | \mathbf{X}, \mathcal{D}_{tr}) + \beta_{\text{KL}} \sum_{(u,v) \in \mathcal{D}_{tr}} \mathcal{H}[q_\phi(z_{(u,v)} | \mathbf{X}, \mathcal{D}_{tr})].$$

From their similarity with VBEs, VEEs inherit the same advantages outlined in Section 4.2.

Models, metrics, and datasets. All models use a 2-layer GNN with Graph Convolutional Network (GCN) layers and a hidden/output nodal dimension of 32. In Base-GNN, an MLP decodes the final nodal embeddings into link probabilities. In VEE-KS/Beta/tanh \mathcal{N} an MLP parameterizes the KS/Beta/tanh \mathcal{N} variational distributions; all take $\beta_{\text{KL}} = .05|\mathcal{D}_{tr}|^{-1}$. We use 10 samples in tanh \mathcal{N} 's entropy estimate; more did not produce significant performance differences. We train for 300 epochs, with a learning rate of .01, averaging results over 5 runs with different seeds. The posterior predictive distribution over binary links $p(\mathbf{A} | \mathbf{X}, \mathcal{D}_{tr}) = \int p(\mathbf{A} | \mathbf{Z}) q_\phi(\mathbf{Z} | \mathbf{X}, \mathcal{D}_{tr}) d\mathbf{Z}$ is estimated by using a single sample from each KS/Beta distribution, parameterizing each edge Bernoulli distribution with such samples, followed by sampling binary edges. For Base-GNN we directly sample binary edges from the likelihood. Using 30 posterior predictive samples, we compute the edge-wise posterior predictive mean (pred. mean) and standard deviation (pred. stdv.). We report the Pearson correlation ρ between predictive uncertainty (pred. stdv.) and error (ℓ_1 difference between pred. mean and the true label), as a measure of uncertainty calibration: useful uncertainty estimates should show strong associations between uncertainty and error. Additionally, we compute area under the ROC curve (AUC) using pred. mean as a predictor. Figure 8 shows performance across 3 standard citation networks: Cora, Citeseer, and Pubmed.

Discussion of results. On all datasets and all metrics, VEE-KS outperforms or matches the most performant baselines, providing higher predictive accuracy (AUC) and better uncertainty calibration (higher ρ). Similar to Section 4.2, we find Beta distributed variational posteriors perform significantly worse than those using KS or tanh \mathcal{N} , further underlining the importance of explicit reparameterization. Moreover, models using explicitly reparameterizable latents are faster: on the largest dataset (Pubmed), the average time (ms) per epoch for VEE-KS, VEE-tanh \mathcal{N} , and VEE-Beta was 381 ± 61 , 301 ± 26 , and 447 ± 86 respectively, on an Apple M2 CPU.

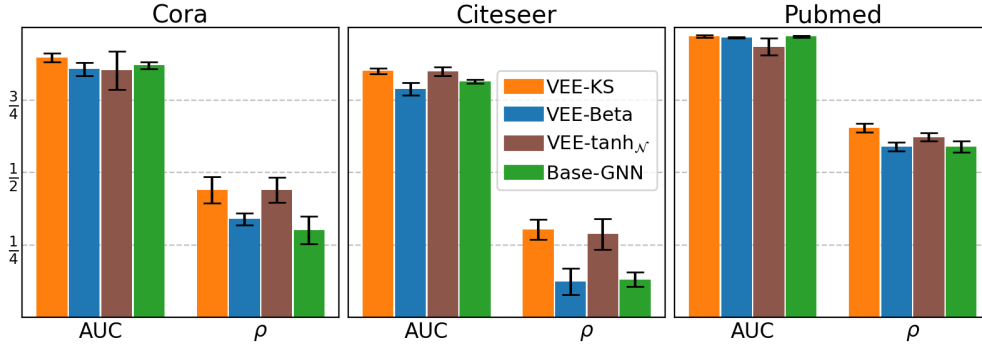


FIG. 8. *VEE-KS produces informative and calibrated edge posterior predictives across graph datasets.*

5. Related Work. This section reviews prior work on TS-based approaches for Bernoulli MABs and the use of the KS distribution as a surrogate for the Beta distribution in latent variable models, highlighting their limitations and connections to our contributions.

5.1. VBEs in context: TS-based Bernoulli MAB approaches. Existing TS-based approaches for Bernoulli MABs assume a prior over model parameters $p(\phi)$, which map contexts to rewards through e_ϕ . At each round, parameters are sampled from the posterior, $\tilde{\phi}_t \sim p(\phi|\mathcal{D})$, and used to compute mean reward posterior samples $\{e_{\tilde{\phi}_t}(\mathbf{x}_k)\}_{k=1}^K$. However, the Bernoulli likelihood often leads to intractable posteriors, making parameter sampling difficult. Common methods use either variational approximations [3, 26, 4], primarily Laplace, or MCMC approaches like Gibbs sampling [5] or LMC [31]. These approaches face several limitations. First, incorporating prior knowledge is challenging since the relationship between a parameter’s value and its effect on rewards is often unclear, except in the simplest models. Second, scalability is a concern: Laplace approximations become inefficient with large context dimensions or model sizes, while MCMC-based methods are compute and memory intensive, requiring long burn-in periods (typically 10^2 iterations) and large machine memory to store the buffer \mathcal{D} . Third, interpreting model beliefs over mean rewards requires drawing numerous posterior samples, adding further computational cost. Finally, these methods often introduce significant complexity through intricate algorithms, architectures, optimization steps, and hyperparameters, particularly MCMC parameters (e.g., burn-in iterations, chain length, LMC inverse temperature/weight decay and their respective schedules). By directly modeling mean rewards with a KS, instead of the parameters ϕ , VBEs offer a simple, scalable, and interpretable approach to Bernoulli MABs.

5.2. Kumaraswamy as a Beta surrogate. A simple approach to overcome the Beta distribution’s lack of explicit reparameterization is to use the KS as a surrogate. This surrogate approach is feasible due to their significant similarities when defined by the same two parameters and the availability of a high-fidelity closed-form approximation of the KL divergence between Beta and KS distributions. [20, 21] use KSs as surrogates for Betas in the Dirichlet Process stick-breaking construction to allow for stochastic latent dimensionality in a VAE. However, both require parameter clipping for numerical stability. In their published code [20] constrains KS parameters $\log a, \log b \in [-2.3, 2.9]$, significantly limiting the expressiveness of latent KS distri-

butions. Also, [21] comments under a *Computational Issues* section that ‘If NaNs are encountered...clipping the parameters of the variational Kumaraswamys usually solve the problem.’ [24] improved upon [20] by resolving the order-dependence issue in approximating a Beta with a KS. Similarly, [23] followed a comparable process using an Indian Buffet Process. Both works maintained numerical stability by restricting the uniform base distribution’s support from the unit interval to a narrower interval, before passing the samples through the inverse CDF producing a distortion of the reparameterized sampling distribution. This work eliminates the need for such distortions, enabling more accurate Beta approximations and simplifying the use of the KS distribution by ensuring numerical stability without additional interventions.

6. Conclusion, Limitations, and Future Work. We identified and resolved key numerical instabilities in the KS distribution, a uniquely attractive option in variational models for bounded latent variables. Our work demonstrates that the stabilized KS can tackle a wide range of large-scale machine learning challenges by powering simple deep variational models. We introduce the Variational Bandit Encoder, which enhances exploration-exploitation trade-offs in contextual Bernoulli MABs, and the Variational Edge Encoder, which improves uncertainty quantification in link prediction using GNNs. Our empirical results show these models are both performant and fast, achieving their best performance with the KS while avoiding the instability and complexity seen in alternatives like the Beta or $\tanh_{\mathcal{N}}$ distributions. These models open avenues for addressing other large-scale challenges, including in recommendation systems, reinforcement learning with continuous bounded action spaces, network analysis, and uncertainty quantification in deep learning, such as modeling per-parameter dropout probabilities using a Concrete distribution [7].

KS generalizations [27] inherit $\log(1 - \exp(x))$ instabilities, which future work can resolve by building on our stabilization technique. A limitation of the current models is their inability to capture multimodal posteriors. Future work could explore KS mixtures or hierarchical latent spaces to bridge this gap. Further, optimizing the β_{KL} parameter with techniques like warm-up schedules could yield further performance gains [1]. Applications of our stable KS distribution to non-parametric models like the Dirichlet Processes follows directly from prior work [21, 24]. Lastly, a theoretical analysis of the VBE, particularly in proving regret bounds and asymptotic results, could extend its applicability to critical areas like clinical trials, where robust decision-making under uncertainty is essential.

Reproducibility Statement. We have made our code publicly available as supplementary material accompanying this submission. Algorithmic details including hyperparameter selections are given in the body, and included in configuration files in the code.

Appendix A. VAE architectural and training choices. The following is almost identical to that used in [18], but provided here for completeness. For both experiments (MNIST and CIFAR-10) we use a learning rate of 0.001, batch size of 500, and optimize with Adam for 200 epochs.

Enforcing positive variational parameters.

- *Gaussian.* When the variational posterior is Normal, the output layer of the encoder uses a softplus nonlinearity for the positive standard deviation.
- *KS.* As we parameterize the KS by unconstrained log values, any required exponentiation occurs internally, so we require no nonlinearity on the output of the encoder.

- *Beta*. The core software libraries do not implement the Beta distribution’s reparameterized sampling with unconstrained log parameter values, so we use an exponential nonlinearity on the output of the encoder to enforce positivity. A softplus nonlinearity was attempted which was found to be less stable likely due to the model seeing very large latent parameter values, which is more stably accessible via an exp.

Enforcing positive likelihood parameters.

- *CB*. When the likelihood is a *CB*, the output of the decoder has a sigmoid nonlinearity to enforce its parameter $\lambda \in (0, 1)$.
- *KS*. As we parameterize the KS by unconstrained log values, any required exponentiation occurs internally, so we require no further transformation on the output of the decoder.
- *Beta*. The core software libraries do not implement the Beta distribution’s log-pdf with unconstrained log parameter values, so we use a softplus nonlinearity on the output of the decoder to enforce positivity. An exponential nonlinearity was attempted which was found to be less stable.

Data augmentation for (0, 1) likelihood functions. The *CB* has support $[0, 1]$ and handles data on the support boundaries without issue. When the likelihood function is a Beta or KS, which have support $(0, 1)$, we clamp pixel intensities to $[\frac{1}{2 \times 255}, 1 - \frac{1}{2 \times 255}]$ to prevent non-finite gradient values.

For all our MNIST experiments we use a latent dimension of $D = 20$, an encoder with two hidden layers with 500 units each, with leaky-ReLU non-linearities, followed by a dropout layer (with parameter 0.9). The decoder also has two hidden layers with 500 units, leaky-ReLU non-linearities and dropout. For all our CIFAR-10 experiments we use a latent dimension of $D = 40$, an encoder with four convolutional layers, followed by two fully connected ones. The convolutions have respectively, 3, 32, 32 and 32 features, kernel size 2, 2, 3 and 3, strides 1, 2, 1, 1 and are followed by leaky-ReLU non-linearities. The fully connected hidden layer has 128 units and a leaky-ReLU non linearity. The decoder has an analogous “reversed” architecture.

Appendix B. Kumaraswamy-Beta KL Divergence. The KL divergence between the Kumaraswamy distribution $q(v)$ with parameters a, b and the Beta distribution $p(v)$ with parameters α, β is given by

$$\mathbb{E}_q \left[\log \frac{q(v)}{p(v)} \right] = \frac{a - \alpha}{a} \left(-\gamma - \Psi(b) - \frac{1}{b} \right) + \log ab + \log \mathcal{B}(\alpha, \beta) - \frac{b - 1}{b} \\ + (\beta - 1)b \sum_{m=1}^{\infty} \frac{1}{m + ab} \mathcal{B} \left(\frac{m}{a}, b \right)$$

where γ is Euler’s constant, $\Psi(\cdot)$ is the Digamma function, and $\mathcal{B}(\cdot)$ is the Beta function. The infinite sum in the KL divergence arises from the Taylor expansion required to represent $\mathbb{E}_q[\log(1 - v_k)]$; it is generally well approximated by the first few terms.

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