Stochastic Variational Inference with Gradient Linearization

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Abstract

Variational inference has experienced a recent surge in popularity owing to stochastic approaches, which have yielded practical tools for a wide range of model classes. A key benefit is that stochastic variational inference obviates the tedious process of deriving analytical expressions for closed-form variable updates. Instead, one simply needs to derive the gradient of the log-posterior, which is often much easier. Yet for certain model classes, the log-posterior itself is difficult to optimize using standard gradient techniques. One such example are random field models, where optimization based on gradient linearization has proven popular, since it speeds up convergence significantly and can avoid poor local optima. In this paper we propose stochastic variational inference with gradient linearization (SVIGL). It is similarly convenient as standard stochastic variational inference – all that is required is a local linearization of the energy gradient. Its benefit over stochastic variational inference with conventional gradient methods is a clear improvement in convergence speed, while yielding comparable or even better variational approximations in terms of KL divergence. We demonstrate the benefits of SVIGL in three applications: Optical flow estimation, Poisson-Gaussian denoising, and 3D surface reconstruction.

1. Introduction

Computer vision algorithms increasingly become building blocks in ever more complex systems, prompting for ways of assessing the reliability of each component. Probability distributions allow for a natural way of quantifying predictive uncertainty. Here, variational inference (VI, see [43] for an extensive introduction) is one of the main computational workhorses. Stochastic approaches to variational inference [17, 21, 31, 33] have recently rejuvenated the interest in this family of approximate inference methods. Part of their popularity stems from their making variational inference applicable to large-scale models, thus enabling practical systems [40]. Another benefit, which should not be underestimated, is that they allow to apply variational inference in a black-box fashion [31, 40], since it is no longer required to carry out tedious and moreover model-specific derivations of the update equations. This allows practitioners to apply variational inference to new model classes very quickly. The only required model specifics are gradients of the log-posterior w.r.t. its unknowns, which are typically much easier to derive than variational update equations. Moreover, automatic differentiation [4] can be used to further reduce manual intervention.

While this makes stochastic variational inference techniques attractive from the user’s perspective, there are some caveats. In this paper we specifically focus on the limitations of gradient-based optimization techniques in the context of certain highly nonlinear model classes. One such category are random field models [6], which often arise in dense prediction tasks in vision. Let us take optical flow [8, 32] as an illustrative example. The data model is highly multimodal and the prior frequently relies on non-convex potentials, which complicate inference [5]. Gradient-based optimization is severely challenged by the multi-modal energy function. Hence, approaches based on energy minimization [8, 32, 42] often rely on a optimization technique called gradient linearization [29], which proceeds by iteratively linearizing the gradient at the current estimate and then solving the resulting system of linear equations to obtain the next iterate. Our starting point is the following question: If gradient linearization is beneficial for maximum a-posteriori (MAP) estimation in certain model classes, would not stochastic variational inference benefit similarly?

In this paper, we derive stochastic variational inference with gradient linearization (SVIGL) – a general optimization algorithm for stochastic variational inference that only hinges on the availability of linearized gradients of the underlying energy function. In each iteration, SVIGL linearizes a stochastic gradient estimate of the Kullback-Leibler (KL) divergence and solves for the root of the linearization. We show that each step of this procedure optimizes a sound objective. Furthermore, we make interesting experimental findings for challenging models from optical flow estimation and Poisson-Gaussian denoising. First, we observe that SVIGL leads to faster convergence of the variational objective function than gradient-based stochas-
tive variational inference (SVI) with the strong optimizers Adam [20] and stochastic gradient descent (SGD). Second, we show that SVIGL is more robust w.r.t. its optimization parameters than standard gradient-based approaches. Finally, SVIGL enables re-purposing existing well-proven energy minimization schemes and implementations to obtain uncertainty estimates while maintaining, or even improving, application performance. Figure 1 shows exemplary flow fields and uncertainty predictions of SVIGL. As expected intuitively, high uncertainty values coincide with errors in the estimated flow field, e.g. near motion discontinuities. Finally, we show that SVIGL benefits problems beyond dense prediction by employing it for 3D surface reconstruction.

2. Related Work

Variational inference. For Bayesian networks, VI w.r.t. the exclusive Kullback-Leibler divergence KL(q∥p) has usually been restricted to certain model classes. The parametric form of the approximating distribution q is chosen such that update equations for the variational parameters of q are analytically tractable. Here, conjugate-exponential models [47] are very common as they often arise in the context of topic modeling, e.g. in the LDA model [7, 38].

In other application areas, e.g. in computer vision, Markov random field (MRF) models are more common. Traditionally, VI has only been applied to specific model classes with closed-form updates, e.g. [11, 23, 24, 27, 36]. Miskin and MacKay [27] pioneered the use of VI for Bayesian blind deconvolution, but made the restrictive assumption that the prior is fully factorized. Levin et al. [23] use a mixture of Gaussian prior on the image derivatives. However, this more powerful prior comes at the cost of additionally maintaining a variational approximation of all mixture components. Krähenbühl and Koltun [22] consider fully-connected conditional random fields (CRF) with Gaussian edge potentials. In this special case mean-field inference can be done efficiently through filtering. Schelten and Roth [36] apply VI to high-order random fields.

In all of the previously mentioned works the variational inference algorithm is closely tied to the probabilistic model at hand and oftentimes requires tedious derivations of analytical update equations. In this paper, we aim to make VI more practical as the only interaction with the probabilistic model is through the linearized gradient of its log probability density function, thus allowing for easy variational inference for a rich class of graphical models.

Stochastic variational optimization. Recently, it was shown that the KL divergence is amenable to stochastic optimization if the approximating distribution q can be re-parameterized in terms of a base distribution that does not or only weakly depend on the parameters of q [21, 33, 35]. While SVI was originally proposed for learning deep latent variable models, such as variational auto-encoders, it is also applicable more generally to graphical models. Re-parameterization allows for deriving efficient stochastic estimators of the gradient of the KL divergence [21, 28]. Only the unnormalized log-density and its gradient w.r.t. the hidden variables are required, thus enabling black-box VI [19, 31, 40]. Note that by stochastic variational inference we do not just refer to the method of Hoffman et al. [17], which, in contrast, requires the true posterior to be from the conjugate-exponential family. Instead, we use the term more generally to describe VI using stochastic optimization.

Having access to a gradient estimator, stochastic algorithms [34] are employed to do the actual optimization. Nowadays, one of the default choices is Adam [20], but other approaches are in use as well, e.g. RMSprop [39], AdaGrad [13], or L-BFGS-SGVI [14]. These algorithms each implement a gradient descent method that is tuned with the recent history of gradient evaluations. In contrast, we assume that we observe a linearization of the gradient and use the information contained therein to modify the direction of
the parameter updates. This can be seen as a gradient descent with a special preconditioner [29], see supplemental.

**Applications of uncertainties.** Aside from being a popular inference tool in many areas of computer vision, e.g. [22, 23], VI yields an assessment of the uncertainty, which can be exploited to post-process point estimates, e.g. with the fast bilateral solver [3]. When used as input for higher-level tasks, optical flow uncertainties allow to discard unreliable estimates and avoid error propagation [45], e.g. in image segmentation [30] or tracking [46]. Uncertainties in image restoration can be beneficial in video restoration, where estimates are fused over several frames [12].

**3. Preliminaries**

**Variational inference** [43] generally aims to approximate an intractable distribution $p$ with a tractable distribution $q$. Since our applications are based on CRFs, we will specifically look at finding approximations to a posterior distribution $p(x \mid y)$. Note, however, that our approach can be applied to marginal and joint distributions as well. We assume that $p$ is a density function over continuous variables, and can be expressed as a Gibbs distribution with its energy function $E(x, y)$ and partition function $Z(y)$ as

$$p(x \mid y) = \frac{1}{Z(y)} \exp \left\{ -E(x, y) \right\}. \quad (1)$$

To ease notation, we assume the temperature parameter to be subsumed into $E(x, y)$, which we furthermore assume to be differentiable. The approximating distribution $q$ is chosen to be a member of some parameterized family of distributions with parameter $\theta$, usually from the exponential family [43]. To determine $q$, variational inference then aims to find variational parameters $\theta$ that minimize the exclusive Kullback-Leibler divergence KL $(q || p)$, i.e.

$$\hat{\theta} = \arg\min_{\theta} \text{KL} (q || p) \quad (2a)$$

$$= \arg\min_{\theta} -E_{q(x; \theta)}[\log p(x \mid y)] + E_{q(x; \theta)}[\log q(x; \theta)] \quad (2b)$$

$$= \arg\min_{\theta} -E_{q(x; \theta)}[\log p(x \mid y)] - H(q), \quad (2c)$$

where $H(q) = H(q(x; \theta))$ denotes the entropy of $q$.

**Gradient linearization.** We now take a step back and first look at MAP estimation for the energy $E(x, y)$ in Eq. (1), i.e. the problem of finding

$$\hat{x} = \arg\max_x \log p(x \mid y) = \arg\min_x E(x, y). \quad (3)$$

Assuming that $E$ is differentiable, we could now apply a standard gradient method, but this may lead to slow convergence. On the other hand, second-order methods may be difficult to apply as the Hessian can be tedious to obtain and/or too dense. For many large-scale prediction problems in computer vision, e.g. estimating optical flow [8, 32], denoising [41], or deblurring [42], this has been addressed through iterative gradient linearization (GL). In this procedure, given a current estimate $x^{(t)}$, the gradient of the energy function $E$ w.r.t. $x$ is linearized around $x^{(t)}$ as

$$\nabla_x E(x, y) \approx \bar{\nabla}_x E(x; x^{(t)}) = A_x(x^{(t)})x + b_x(x^{(t)}). \quad (4)$$

For notational brevity, we omit $y$ here and in the following. Note that the linearized gradient $\bar{\nabla}_x E(x; x^{(t)})$ is exact at $x = x^{(t)}$. To obtain the next iterate $x^{(t+1)}$, we set $\bar{\nabla}_x E$ to zero and solve the resulting linear system of equations

$$x^{(t+1)} = -A_x^{-1}(x^{(t)})b_x(x^{(t)}) \quad (5)$$

using an exact or approximate standard solver. Like in any iterative optimization, an initial guess $x^{(0)}$ is required.

Iterative GL is also known by various other names. Nikolova and Chan [29] showed it to be equivalent to multiplicative half-quadratic minimization [16] for Gaussian likelihoods. Moreover, it is closely related to iteratively reweighted least squares through their equivalence to half-quadratic approaches [18]. Finally, GL can be seen as preconditioned gradient descent using $A_x^{-1}$ as preconditioner [29], c.f. supplemental. In comparison to Newton’s method no second-order derivatives are required – a benefit that is shared with other quasi-Newton methods, such as the popular L-BFGS [10]. However, every regular gradient step couples variables only within a local spatial neighborhood. In contrast, one iteration of GL (Eq. 5) causes a joint update of all variables leading to faster convergence in highly non-linear objectives (see Fig. 2 for an example).

**4. Stochastic Variational Inference with Gradient Linearization (SVIGL)**

We now aim to leverage the advantages of GL in the context of stochastic variational inference. To that end, we assume access to a linearized gradient, given by $A_x$ and $b_x$ in Eq. (4). By applying the re-parameterization trick [21, 33], we can rewrite the KL divergence of Eq. (2) as

$$\hat{\theta} = \arg\min_{\theta} -E_{z \sim G} \left[ \log p(x(z) \mid y) \right] - H(q), \quad (6)$$

where $x(z) \equiv x(z; \theta)$, and $z$ is distributed following a base distribution $G$ independent of $\theta$. In the following, we approximate the full expectation over $z$ with a finite set of samples $\mathcal{Z} = \{z_i\}$. Using the approximation to the true gradient given by $A_x$ and $b_x$, we can then easily derive a stochastic approximation of the gradient of the KL diver-
To illustrate the use of Gaussian mean field inference. To do this, we consider the individual
respect to \( \mu \) distributed, \( \bar{q} \equiv \approx \frac{1}{|Z|} \sum_{z_i \in Z} \theta \equiv \frac{1}{|Z|} \sum_{z_i \in Z} \left( A_x(x(z_i)) x(z_i) + b_x(x(z_i)) \right) \cdot \nabla_\theta x(z_i) - \frac{1}{|Z|} \sum_{z_i \in Z} \mu \cdot \nabla_\theta H(q)
\end{equation}
\begin{equation}
\equiv \nabla_\theta \text{KL}(q \parallel p).
\end{equation}

Gaussian mean field inference. To illustrate the use of this approximation, we now apply the common naive mean-field framework [11, 21, 23] and assume that the variational distribution \( q \) factorizes along all elements of \( x = (x_l) \), for \( l = 1, \ldots, L \). Moreover, \( q \) is modeled as an uncorrelated Gaussian distribution with \( \theta = \{ \mu, \sigma \} \):
\begin{equation}
q(x) = \prod_{l=1}^L \mathcal{N}(x_l | \mu_l, \sigma_l^2).
\end{equation}
Following [21], \( z \) is thus chosen to be standard normally distributed, \( \text{i.e.} \, z \sim \mathcal{N}(0, I) \), and we set \( x(z) = z \cdot \sigma + \mu \) with element-wise operations.

For the case of a fully-factorized Gaussian \( q \), it is now possible to express \( \nabla_\theta \text{KL}(q \parallel p) \) again in the form of a linearized gradient. To do this, we consider the individual parameter gradients w.r.t. \( \mu \) and \( \sigma \). For the gradient with respect to \( \mu \), we exploit that the entropy of a Gaussian distribution does not depend on its mean. Hence, we arrive at
\begin{equation}
\nabla_\mu \text{KL}(q \parallel p) = \frac{1}{|Z|} \sum_{z_i \in Z} \left( A_x(x(z_i)) x(z_i) + b_x(x(z_i)) \right) \cdot \nabla_\mu x(z_i)
\end{equation}
\begin{equation}
= \frac{1}{|Z|} \sum_{z_i \in Z} A_x(x(z_i)) (z_i \cdot \sigma + \mu) + b_x(x(z_i))
\end{equation}
\begin{equation}
= \left[ \frac{1}{|Z|} \sum_{z_i \in Z} A_x(x(z_i)) \right] \mu
\end{equation}
\begin{equation}
+ \left[ \frac{1}{|Z|} \sum_{z_i \in Z} A_x(x(z_i)) \right] \sigma
\end{equation}
\begin{equation}
+ \left[ \frac{1}{|Z|} \sum_{z_i \in Z} b_x(x(z_i)) \right]
\equiv A_{\mu, \mu}(\theta) \mu + A_{\mu, \sigma}(\theta) \sigma + b_\mu(\theta),
\end{equation}
where \( D(z_i) \) denotes a diagonal matrix comprised of the elements of \( z_i \). The gradient w.r.t. \( \sigma \) involves the derivative of the Gaussian entropy, \( \text{i.e.} \, \nabla_\sigma H(q) = \nabla_\sigma (\log \sigma + \text{const}) \), which can be linearized in several ways. We opt for using the element-wise second-order Taylor expansion of the logarithm around the current estimate: \( \sigma^{(t)} \):
\begin{equation}
\log \sigma \approx \log \sigma^{(t)} + \frac{1}{\sigma^{(t)}} (\sigma - \sigma^{(t)}) - \frac{1}{(\sigma^{(t)})^2} (\sigma - \sigma^{(t)})^2
\end{equation}
\begin{equation}
= \frac{1}{\sigma^{(t)}} \sigma - \frac{1}{(\sigma^{(t)})^2} (\sigma - \sigma^{(t)})^2 + \text{const.}
\end{equation}
With that we can derive our stochastic approximation to the linearized gradient of the KL divergence w.r.t. \( \sigma \) as
\begin{equation}
\nabla_\sigma \text{KL}(q \parallel p) = \frac{1}{|Z|} \sum_{z_i \in Z} \left( A_x(x(z_i)) x(z_i) + b_x(x(z_i)) \right) \cdot \nabla_\sigma x(z_i)
\end{equation}
\begin{equation}
- \frac{3}{\sigma^{(t)}} + \frac{2}{(\sigma^{(t)})^2} \sigma
\end{equation}
\begin{equation}
= \left[ \frac{1}{|Z|} \sum_{z_i \in Z} D(z_i) A_x(x(z_i)) \right] \mu
\end{equation}
\begin{equation}
+ \left[ \frac{1}{|Z|} \sum_{z_i \in Z} D(z_i) A_x(x(z_i)) \right] \sigma
+ \left[ \frac{1}{|Z|} \sum_{z_i \in Z} b_x(x(z_i)) \right] - \frac{3}{\sigma^{(t)}}
\end{equation}
\begin{equation}
\equiv A_{\sigma, \mu}(\theta) \mu + A_{\sigma, \sigma}(\theta) \sigma + b_\sigma(\theta).
\end{equation}

From Eqs. (9d) and (11d), we now obtain an approximate linearized gradient of the KL divergence in Eq. (2) with respect to \( \mu \) and \( \sigma \). Following the GL procedure, the optimization proceeds by solving the linear system of equations
\begin{equation}
\theta^{(t+1)} = -A_\theta \left( \theta^{(t)} \right)^{-1} b_\theta \left( \theta^{(t)} \right)
\end{equation}
with
\begin{equation}
A_\theta(\theta) = \begin{bmatrix} A_{\mu, \mu}(\theta) & A_{\mu, \sigma}(\theta) \\ A_{\sigma, \mu}(\theta) & A_{\sigma, \sigma}(\theta) \end{bmatrix}, \quad b_\theta(\theta) = \begin{bmatrix} b_\mu(\theta) \\ b_\sigma(\theta) \end{bmatrix}.
\end{equation}

Note that we can treat the underlying energy \( E \) as a black box. The only interaction with \( E \) is through its linearized gradient. Algorithm 1 summarizes our approach.

Discussion. Each gradient iteration in Eq. (12) can be interpreted as fitting a quadratic function to the Monte Carlo
approximation of the KL divergence (Eq. 7b), such that the quadratic approximation and the KL divergence agree on their first-order derivatives at \( \theta^{(t)} \). This alone does not guarantee that the extremum \( \theta^{(t+1)} \) of the quadratic function is actually a minimum of the approximation. Hence, we now show that the Hessian of the quadratic approximation \( A_\theta \) is positive semi-definite, thus ensuring that \( \theta^{(t+1)} \) minimizes the approximated KL divergence.

**Proposition 1.** \( A_\theta(\theta^{(t)}) \) is positive semi-definite, i.e. \( \theta^T A_\theta(\theta^{(t)}) \theta \geq 0, \forall \theta, \theta^{(t)} \in \mathbb{R}^{2L}, \) if the matrix \( A_x(\mathbf{x}(z)) \) of the energy GL is positive semi-definite for all \( z \).

*Proof.* Let us first assume that we just draw a single sample \( z \). To simplify notation let \( A_x \equiv A_x(\mathbf{x}(z)) \) and \( A_\theta \equiv A_\theta(\theta^{(t)}) \). Now, for \( \theta = [\mu, \sigma]^T \) we have that

\[
\theta^T A_\theta \theta \equiv \mu^T A_\mu, \mu + \sigma^T A_{\mu}, \mu + \mu^T A_{\mu}, \sigma + \sigma^T A_{\mu}, \sigma \equiv (14a)
\]

\[
= \mu^T A_\mu, \mu + \sigma^T D(z)^T A_x, \mu + \mu^T A_x, D(z) \sigma + \sigma^T D(z)^T \left( A_x + \mathbf{D} \left( \frac{2}{(\sigma^2)} \right)^2 \right) D(z) \sigma \equiv (14b)
\]

\[
= (\mu + \mathbf{D}(z) \sigma)^T A_x (\mu + \mathbf{D}(z) \sigma) + \mathbf{D}(z) \sigma^T \mathbf{D} \left( \frac{2}{(\sigma^2)} \right)^2 (\mathbf{D}(z) \sigma) \geq 0, \quad (14c)
\]

where we inserted the definition of the individual matrices (Eqs. 9d and 11d). For the last step, we used our assumption that \( A_x \) is positive semi-definite. The case of multiple samples \( z_i \) can be shown analogously by expanding each of the four terms in Eq. (14a) into a sum.

To put the above proposition into perspective, we now give two mild conditions on the energy function such that the corresponding matrix \( A_x \) is positive semi-definite.

**Proposition 2.** An energy function can be linearized with a positive semi-definite matrix \( A_x \) if it is composed of a sum of energy terms \( \rho_i(\mathbf{w}_i) \) that fulfill the following conditions:

1. Each penalty function \( \rho_i(\cdot) \) is symmetric and \( \rho_i'(\mathbf{w}_i) \geq 0 \) for all \( \mathbf{w}_i \geq 0 \). (⋆)
2. Each penalty function \( \rho_i(\cdot) \) is applied element-wise on \( \mathbf{w}_i \), which is of the form \( \mathbf{w}_i = \mathbf{K}_i \mathbf{x} + \mathbf{g}_i(y) \), with filter matrix \( \mathbf{K}_i \) and function \( \mathbf{g}_i \), not depending on \( \mathbf{x} \). (***)

*Proof.* See supplemental material.

The above assumptions of Proposition 2 are not very restrictive but met by many MRF/CRF potentials [6], including the smoothness term used in optical flow and Poisson-Gaussian denoising, as well as the data term of our flow energy, c.f. Sec. 5.1 and 5.2. Moreover, positive semi-definiteness of \( A_x \) can also be shown for more complex energy formulations such as the data term of Poisson-Gaussian denoising used in our experiments.

**Implementation details.** Solving the linear system of equations of Eq. (12) exactly is too costly for many large-scale problems, which may involve millions of variables. Hence, we consistently apply 100 iterations of successive over-relaxation [48] with a relaxation factor of 1.95 and the current estimate \( \theta^{(t)} \) as initialization. We also experimented with a conjugate gradient optimizer, but found convergence to be too slow, probably due to the need of an effective preconditioner. One limitation of our method is that we cannot guarantee that \( \sigma \) stays positive after each optimization step. Therefore, we replace each new iterate \( \sigma^{(t+1)} \) with its absolute value. In practice, however, we found that usually the entropy term is enough to force \( \sigma \) to stay positive. Since the gradient of the KL divergence cannot be expressed conveniently as linear in \( \log \sigma \), we do not use the usual trick of optimizing for \( \log \sigma \) to directly enforce positivity of \( \sigma \).

5. Experiments

We now demonstrate that SVIGL provides a convenient and efficient way of obtaining accurate variational approximations for popular energy functions of diverse computer vision problems, yielding uncertainty estimates that correlate well with estimation errors. Specifically, we quantitatively evaluate on the tasks of optical flow estimation and Poisson-Gaussian denoising. We compare SVIGL against gradient-based optimization of the KL divergence with SGD as well as the Adam optimizer [20], the default choice in the popular Edward library [40]. To assess the quality of the obtained approximate posterior, we evaluate the KL divergence \( KL(\theta || \rho) \) as well as application specific performance metrics. We always report KL divergences approximated by sampling (c.f. Eq. 6) and up to the unknown, but constant log partition function \( \log Z(y) \).

We conduct several experiments for each application. We begin by evaluating the robustness of Adam (in the con-
text of stochastic variational inference) and SVIGL w.r.t. to their parameters. We first vary the step size $\alpha$ of Adam while using $|Z| = 50$ samples per iteration to approximate the KL divergence gradient. Next, we use the best step size and vary the size of the sample set $|Z|$ for both Adam and SVIGL. For a sample set size of 50, 25, and 12, we set the number of iterations to 100, 200, and 400 for SVIGL and 1000, 2000, and 4000 for Adam, respectively. For SGD, we similarly tune the hyperparameters and find that 4000 iterations with 12 samples and an initial step size of $10^{-6}$, which is cut after each third of iterations by a factor of ten, works best for both applications. We compare the best configurations of SVIGL and SVI with SGD and Adam to a Laplace approximation and MAP estimation baselines. Runtimes refer to an Intel Xeon E5-2650v4, 2.2 GHz, 12 cores. We furthermore show qualitative results for 3D surface reconstruction to demonstrate the benefit of SVIGL for non-vision applications.

5.1. Optical flow

We first apply SVIGL to estimate an optical flow field $x$, describing the motion between images $y = \{I_1, I_2\}$. We use the EpicFlow energy of [32] to induce a Gibbs distribution akin to Eq. (1). Its likelihood encourages the flow to be consistent with the images and is based on a gradient consistency assumption, whereas the prior assumes small flow gradients over a 4-neighborhood, i.e.

$$E(x, y) = \lambda_D \sum_{l=1}^{L} \rho_D \left( \left\| \left( \nabla \tilde{I}_2(x) - \nabla I_1 \right) \right\|_2 \right) + \lambda_S \sum_{j=1}^{J} \sum_{l=1}^{L} \rho_S \left( \left\| (f_j \ast x) \right\|_2 \right).$$

Here, $\nabla I_1$ denotes the spatial derivatives of $I_1$, $\tilde{I}_2(x)$ is the second image warped by $x$, and $f_1, \ldots, f_J$ represent (derivative) filters. Functions $\rho_D$ and $\rho_S$ are robust penalty functions weighted with parameters $\lambda_D, \lambda_S$. Following standard practice, we linearize the likelihood around the current flow.

Setup. As in [45], we initialize our estimates with sparse FlowFields matches [1], densified with the EpicFlow interpolation [32]. Variances are initialized as $\sigma = 10^{-3}$. We use generalized Charbonnier penalties [2] and obtain their parameters as well as the ratio $\lambda_D/\lambda_S$ through Bayesian optimization [37]. To that end, we evaluate the average end-point error (AEPE) of MAP estimates on a subset of Sintel train [9]. The absolute scale of $\lambda_D$ and $\lambda_S$ is subsequently calibrated such that the AEPE of the SVIGL estimates remains comparable to the MAP estimates on the training set.

Results. We conduct experiments on a validation set of 104 images randomly chosen from Sintel training (excluding images used for parameter optimization). We first motivate the use of gradient linearization by comparing the results of MAP estimation performed with up to 200 iterations of L-BFGS to 20 iterations of GL. The results averaged over the validation set are depicted in Fig. 2. We observe a significantly faster minimization of the energy using GL, which highlights its benefits for highly non-linear objectives.

We now compare SVIGL to SVI with Adam. In order to keep the runtime of Adam manageable, we perform the evaluation on manually cropped patches of size $100 \times 100$. In a first setting, we vary the step size $\alpha$ of Adam using 1000 iterations for Adam and 100 iterations for SVIGL. In Fig. 3a, we evaluate the KL divergence plotted against the runtime. SVIGL reduces the KL divergence two orders of magnitude faster than Adam on this challenging energy function. Moreover, the optimization by Adam is highly dependent on the chosen step size; too small or too large a value may equally lead to slow convergence. In contrast, SVIGL does not require the selection of a step size. For the following experiments we fix the step size for Adam to $\alpha = 0.005$. Now, we vary the number of samples and iterations as described above. The results are shown in Fig. 3b. Again, SVIGL attains a significantly better variational approximation than SVI with Adam for all examined settings.

We provide a comparison for SGD over the full training set in Table 1. We observe a significantly faster minimization of the energy using GL, which highlights its benefits for highly non-linear objectives. Furthermore, the Laplace approximation requires a significantly lower KL divergence than SVI with Adam or SGD. However, the Laplace approximation is significantly faster than Adam on this challenging energy function. Moreover, the optimization by Adam is highly dependent on the chosen step size; too small or too large a value may equally lead to slow convergence. In contrast, SVIGL does not require the selection of a step size. For the following experiments we fix the step size for Adam to $\alpha = 0.005$. Now, we vary the number of samples and iterations as described above. The results are shown in Fig. 3b. Again, SVIGL attains a significantly better variational approximation than SVI with Adam for all examined settings.

Finally, we evaluate SVIGL on the full-size images of Sintel test. Since SVI with Adam is too slow, we only compare to MAP baselines with 200 iterations of L-BFGS and 20 iterations of GL, respectively. For SVIGL we use 50 samples and also 20 iterations. Both SVIGL and GL yield an AEPE of 5.74 and therefore outperform the L-BFGS baseline with an AEPE of 5.81.

**Interpretation.** The interdependent updates of SVIGL (Eq. (12)) cause information to flow between all variables.

<table>
<thead>
<tr>
<th>Method</th>
<th>KL $\times 10^7$</th>
<th>runtime [s]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initialization</td>
<td>5.13</td>
<td>–</td>
</tr>
<tr>
<td>GL + Laplace</td>
<td>3.83</td>
<td>–</td>
</tr>
<tr>
<td>SVI + SGD</td>
<td>4.45</td>
<td>551</td>
</tr>
<tr>
<td>SVI + Adam</td>
<td>4.24</td>
<td>1148</td>
</tr>
<tr>
<td>SVIGL (ours)</td>
<td>3.78</td>
<td>584</td>
</tr>
</tbody>
</table>

Table 1. Unnormalized KL divergence and average runtime on $100 \times 100$ crops from a Sintel validation set.
while a regular gradient step propagates information in a local spatial neighborhood only. We attribute the observed performance gain of GL and SVIGL over gradient-based methods at least partly to this global update.

**Uncertainty estimates.** Finally, we assess the quality of the per-pixel uncertainty estimates. To this end, we compare SVIGL to update the continuous variables of their energy formulation; see supplemental material for further implementational details. Table 2 shows the metrics introduced in [45], averaged over the full-size images of our validation set. The uncertainty estimates obtained by SVIGL are competitive with the ones of ProbFlowFields. More importantly and unlike [45], the application of SVIGL does not require the tedious derivation of update equations. Example flow fields and the inferred uncertainty maps are shown in Fig. 1.

### 5.2. Poisson-Gaussian denoising

We next apply SVIGL to the problem of removing Poisson-Gaussian noise [15]. Here, it is assumed that image noise comes mainly from two sources that inherently affect any camera sensor. First, the Poissonian arrival process of photons hitting the pixels, and second an additive Gaussian component arising from noise in the electronics of the sensor. The Poisson distribution can be well approximated by a Gaussian [15], giving rise to a Gaussian likelihood with intensity dependent variance, i.e.

\[
y_l \sim N(x_l, \sigma(x_l)^2) \text{ with } \sigma(x_l)^2 = \beta_1 x_l + \beta_2,
\]

where the noise distribution is specified by the parameters \( \beta_1 \) and \( \beta_2 \). We specifically set \( \beta_1 = 0.05 \) and \( \beta_2 = 0.0001 \) in order to simulate strong noise (Poisson rate 20). Combining this likelihood with a 4-connected pairwise MRF with generalized Charbonnier potentials [2] as image prior leads to the energy

\[
E(x, y) = \frac{\lambda_D}{2} \sum_{l=1}^{L} \frac{(x_l - y_l)^2}{\sigma(x_l)^2} + \lambda_S \sum_{j=1}^{J} \sum_{l=1}^{L} \rho_S \left( (f_j * x_l) \right),
\]

(17)

where the \( f_j \) denote horizontal and vertical image derivative filters. The temperature is subsumed by the weights \( \lambda_D, \lambda_S \).

**Setup.** We select the relative importance of \( \lambda_D \) and \( \lambda_S \) as well as the exponent of the robust penalty through Bayesian optimization [37]. To this end, we optimize the peak-signal-to-noise ratio (PSNR) after 20 steps of GL on a set of 100 images from the BSDS training set [26]. We then calibrate the posterior for VI by determining the absolute scale of the weights on the training set. To synthesize noisy images for parameter tuning and testing, we apply Poisson-Gaussian noise to clean ground truth images. Afterwards, we rescale the intensities such that the ground truth lies in \([0, 1]\) and clip the noisy image to that range. For test time inference, we initialize \( \mu \) with the noisy image and \( \sigma \) as \( 10^{-3} \).

**Results.** In Fig. 4 we plot the unnormalized KL divergence against runtime for SVIGL and SVI with Adam, using varying step sizes for Adam and varying sizes of the sample set \( Z \) for both methods. It becomes apparent that the performance of Adam highly depends on these two parameters.
Figure 4. Runtime vs. unnormalized KL divergence for denoising with SVIGL and SVI with Adam with different stepsize parameters $\alpha$ (a) and varying sizes of the sample set $|Z|$ (b). Values averaged over the BSDS test set.

Too small a step size slows down convergence, while setting it too high leads to a KL divergence inferior to the initialization. In contrast, SVIGL does not require setting a step size and converges faster than Adam with the best step size $\alpha = 0.01$. For instance, SVIGL reaches the same KL divergence as Adam in only $1/5$ of the time. When looking at the size of the sample set, we note that smaller sample sets speed up each iteration and hence lead to faster progress of the optimization. However, the solution found by Adam deteriorates after a certain number of iterations with smaller sample set sizes, while SVIGL is not affected by this issue. In summary, SVIGL yields faster convergence while being robust to the setting of nuisance parameters.

The converged solutions are evaluated in Table 3. SVIGL ($|Z| = 50$) not only converges significantly faster than Adam ($\alpha = 0.01$, $|Z| = 50$), but obtains even slightly improved solutions. SGD performs significantly worse than SVIGL and Adam. A Laplace approximation around the mode obtained with 100 iterations of GL provides a poor fit to the denoising posterior since the dependence of the variances $\sigma(x_i)$ on the noise-free intensities $x_i$ results in a skewed distribution. Furthermore, we see that SVIGL obtains a better solution in terms of the standard image quality metrics PSNR and SSIM [44] than the MAP estimation baselines obtained with GL and L-BFGS, e.g. +1.6 dB in PSNR compared to L-BFGS. In the supplemental material we show denoised images obtained by SVIGL along with their uncertainty estimates.

5.3. 3D surface reconstruction

In order to demonstrate that SVIGL is not limited to low-level problems in computer vision, we apply it to the task of reconstructing a smooth point cloud from noisy input data. Specifically, we use the energy of [25] given as

$$E(X, P, C) = \sum_{i=1}^{[X]} \sum_{j=1}^{[P]} \|x_i - p_j\| \cdot h(\|c_i - p_j\|) - \sum_{i=1}^{[X]} \sum_{i'=1}^{[C]} \lambda_i \|x_i - c_{i'}\| \cdot h(\|c_i - c_{i'}\|).$$

Here, $p_j \in P$ denote the noisy input points; the current and the new estimate of the smoothed points are given by $c_i \in C$ and $x_i \in X$, respectively. The contribution of each term is weighted by a Gaussian kernel $h(\cdot)$. Following Lipman et al. [25], we use this energy in a fixed point scheme, i.e.

$$X_{t+1} = \arg \min_{X} E(X, P, X_t),$$

where $X_0$ is an $L_2$ projection of the input points. The supplemental material describes the setup in more detail.

In order to exemplify the use of SVIGL for 3D surface reconstruction, we synthesize a noisy input point cloud of the Stanford bunny by adding noise on the positions of reference points. The noise strength gradually increases from tail to face. Figure 5 shows both the noisy input point cloud as well as the variational approximation from SVIGL with color coded uncertainty $\sigma$. It is apparent that the uncertainty increases with input noise strength, thus reflecting the difficulty of the reconstruction task. Moreover, at points further away from the true surface, the uncertainty is generally higher, c.f. the outliers at the ears.

6. Conclusion

Motivated by the success of gradient linearization techniques for MAP estimation in highly multimodal posteriors, we proposed to combine the benefits of gradient linearization with stochastic variational inference. As a result we obtain SVIGL, an easy-to-use variational inference scheme that only requires access to a gradient linearization of the posterior energy and allows to simply re-purpose well-proven energy minimization schemes. We applied SVIGL to optical flow estimation as well as Poisson-Gaussian denoising and demonstrated its significantly faster convergence compared to standard stochastic variational inference. Moreover, we showed that the optimization accuracy of SVIGL is robust to the choice of parameters. The inferred uncertainty estimates are competitive with state-of-the-art but can be obtained without tedious derivations of update equations. Finally, we demonstrate that SVIGL is not restricted to dense 2D prediction tasks by applying it successfully to the task of 3D surface reconstruction.

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