Bringing together visual analytics and probabilistic programming languages

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Zusammenfassung

A probabilistic programming language (PPL) provides methods to represent a probabilistic model by using the full power of a general purpose programming language. Thereby it is possible to specify complex models with a comparatively low amount of code. With Uber, Microsoft and DARPA focusing research efforts towards this area, PPLs are likely to play an important role in science and industry in the near future. However in most cases, models built by PPLs lack appropriate ways to be properly visualized, although visualization is an important first step in detecting errors and assessing the overall fitness of a model. This could be resolved by the software Lumen, developed by Philipp Lucas, which provides several visualization methods for statistical models. PPLs are not yet supported by Lumen, and the goal of the master thesis at hand is to change that by implementing an interface between Lumen and a chosen PPL, so that exploring PPL models by visual analytics becomes possible. The thesis will be divided into two main parts, the first part being an overview about how PPLs work and what existing PPLs there are available. Out of these, the most appropriate one will be chosen for the task. The second, more practical part will then document the actual implementation of the interface.

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1 Road Map

- 1. Getting started
 - set up Master thesis document
 - Probabilistic Programming Languages
 - play at least with: PyMC3, Stan
 - read the docs, wiki, ...
 - download the libraries
 - reproduce the getting started tutorials
 - --> understand the ideas and how to use it, get a feeling for it
 - theoretic background: Read Bayesian Data Analysis part I, Chapter 1,2 and part II, chapter 6,7
 - Lumen
 - install locally and play with
 - understand main idea of Lumen and what we want to do with it
 - Start filling up your MA thesis document
 - understand and write down in MA thesis the "why & what"
 - describe the background of the work, e.g. summarize PPLs
 - give a short presentation
 - what have you done and learned
 - what do you plan to do?
 - why is it relevant?
 - how do you plan to measure the success?
- 2. First connection of PPLs to Lumen
 - Start from small, very simple and specific example. Generalize later.
 - Choose PPL to work with
 - work out requirements on PPL
 - work out preferred features of PPL
 - choose a PPL based on these requrements and preferences
 - design wrapper of PPL with Lumen
 - work out requirement and interface
 - identify necessary work on Lumen
 - identify necessary work
 - Connect chosen specific example with lumen
 - Continue to work on your master thesis document!
- 3. Improve, generalize and clean up the connection of your PPL to Lumen

2 Introduction

3 The Bayesian Approach

"Probabilistic programming is an emerging paradigm in statistical learning, of which Bayesian modeling is an important sub-discipline."[1]

In the field of statistics, one deals usually with the following questions:

- What kind of model should I choose?
- Which features should I include in my model?
- What are the most likely values for my model parameters?
- What is the most likely outcome for additional observations under certain conditions?
- How certain am I that my estimated parameter values are correct?

The latter three of these questions can be answered by using a Bayesian approach to statistics. Given a model, Bayesian statistics is about computing a joint probability distribution over all parameters and outcomes, so that a probability density for each possible value of parameters and outcomes can be obtained. Prior knowledge of the problem as well as observed data affect the form of the joint probability distribution. Using marginalization, conditional probability distributions can then be calculated for the desired outcomes. These conditional probability distributions are also called posteriors and can be used for parameter estimation, prediction and assessing the uncertainty of estimates. This chapter will outline how Bayesian statistics are applied to answer the abovementioned questions.

3.1 Rules of Probabilistic Inference

When working with Bayesian models, it will be necessary to transform conditional, marginal and joint distributions into one another. There are three rules of probabilistic inference which achieve this: The chain rule, the total probability rule, and the Bayes' rule. The following explanations are taken from [2].

3.1.1 Chain rule

The chain rule is used to calculate a joint probability distribution of several variables from local conditional probability distributions of these variables:

$$P(X_1, X_2, ... X_n) = P(X_1)P(X_2|X_1)P(X_3|X_1, X_2)...P(X_n|X_1, X_2, ... X_{n-1}))$$
(1)

3.1.2 Total probability rule

The total probability rule calculates the probability distribution over a subset of variables, also called a marginal distribution, by summing out all the other variables, that is by summing the probability distributions for each combination of values of these variables:

$$P(\boldsymbol{X}|\boldsymbol{Z}) = \sum_{\boldsymbol{y}} P(\boldsymbol{X}, \boldsymbol{Y} = \boldsymbol{y}|\boldsymbol{Z})$$
 (2)

3.1.3 Bayes' rule

Bayes' rule calculates the probability of a cause, given an effect, by using the prior probability of the cause and the probability of the effect, given the cause.

$$P(X|Y) = (P(Y|X) * P(X))/P(Y)$$
(3)

3.2 Difference to frequentist approach

Classical approach: Evaluate the procedure used to estimate the parameters over the distribution of possible outcome values conditional on the true unknown parameters.

Bayesian approach: Estimate the parameters conditioned on the outcomes. [3] Bayesian models are about finding a full probability model for a given problem, whereas conventional models only deal with estimation of the most likely parameters. The model parameters β themselves are also considered as random variables which depend on hyperparameters α .

3.3 Choosing an appropriate prior distribution

There are two interpretations of prior distributions. The *population* interpretation, where the prior distribution is thought of as a population of possible parameters, from where the current parameter is drawn. This, as far as I understand, requires the range of possible values to be known, e.g from past experience. On the other hand, the *state of knowledge* interpretation looks at the prior distribution as an expression of the user's knowledge or uncertainty, so that the assumption is plausible, that the real parameter value is taken from a random realization of the prior distribution.

When we have lots of knowledge about the parameter already, it makes sense to choose an *i*nformative prior, which has a big influence on the posterior distribution. On the other hand, if one does not want prior beliefs to affect the outcome of an analysis, one should not choose informative priors, even when lots of knowledge is available. E.g. when testing a hypothesis, it is not wise to include information in the prior that supports the hypothesis for fairness reasons(?? look that up again on p. 56.). If there is no sufficient data to estimate a prior, it is desirable to choose a prior that is *n*oninformative, meaning that it will contribute very little to the posterior distribution ('let the data speak for itself'). A

common noninformative prior, motivated by Laplace's principle of insufficient reason, is the uniform distribution. However, the uniform distribution is not ideal since it is dependent on the parametrization: Applying a uniform distribution to p(x) leads to a non-uniform distribution when looking at $p(x^2)$ and the other way round [3]. Jeffrey's approach to find noninformative prior distributions negates this problem by choosing a prior that is invariant to the parameterization (see https://eventuallyalmosteverywhere.wordpress.com/2013/05/10/bayesian-inference-and-the-jeffreys-prior/). Besides informative and noninformative prior distributions there is also the weakly informative prior distribution. This kind of prior does affect the posterior distribution in terms of regularization (e.g. it prevents extreme outliers), but it does not contain any further special knowledge about the parameter. A normal distribution with high variance is often used as weakly informative prior.

The parameters of the prior distributions are called hyperparameters. The property that prior and posterior distribution are of the parametric form (e.g., both are a beta distribution) (for a given likelihood distribution), is called conjugacy. Conjugate prior distributions have the advantages of being computationally convenient and being intertpretable as additional data.

A prior distribution is called *proper* if it does not depend on data and sums to 1 [3]. Proper distributions can be normalized. The uniform prior for example is *improper*, since it can't be normalized.

3.4 Sampling distribution

The sampling distribution, also called the likelihood of the data.

Often it is about which distribution class should be chosen for the prior and the likelihood. Standard, convenient distributions for single-parameter models: normal, binomial, Poisson, exponential. Those can also be combined to represent more complex distributions. For different classes of sampling distributions there are corresponding conjugate prior distributions which lead in turn to posterior distributions of the same form. [3]

distributions can be chosen for mathematical convenience. One could estimate hyperparameters of the likelihood distribution from the data in some cases. This is a bit of a circular reasoning, but apparently it is appropriate for [3].

3.5 Posterior distribution

The posterior distribution is a compromise between the prior distribution and the sample distribution, with the prior distribution becoming less important as the sample size increases [3]. posterior distributions can be described numerically by mean, median, modes. The uncertainty can be described by quantiles. The highest posterior density region is also a possibility, it is the area that for example contains 95% of the posterior probability density, like quantiles, but has the additional constraint that the density on each point inside the area has to be bigger than the density on any point outside the area. It does not have to

be one single connected area [3].

For the posterior distribution, a normal distribution is often assumed.

3.6 Drawing inference

According to [4], the likelihood of a new datapoint can be calculated by integrating the product of the prior likelihood and conditional probability of β over the space of β , as shown in equation 4. This formula can also be derived using the chain rule (I think).

Making predictions for a new data points requires both the probability distribution of the new data point, given a parameter value, as well as the probability distribution of the parameter, given the old data points.

If it is not possible to perform calculations directly from the posterior distribution (e.g. if there is no closed form but only a discrete approximation of the posterior distribution), one can simulate data points from the posterior distribution instead and perform calculations on them [3].

$$p(x_{new}|\boldsymbol{x},\alpha) = \int p(x_{new}|\beta) * p(\beta|\boldsymbol{x},\alpha)d\beta$$
 (4)

3.7 Evaluating models

Since "all models are wrong", it is necessary, after a Bayesian model has been found, to capture the extent to which the model fails to describe reality. In Bayesian analysis, both the prior and the likelihood distribution can be sources of error.

Different models can lead to different posterior inferences, even if both models have a good fit.

model checking: How good does the model fit the data? Sensitivitiy analysis: Comparison of different models with good fit

This is necessary after a joint probability density and a posterior density are calculated. How good are the prior and the likelihood model? *Sensitivity analysis* deals with the question, how much the posterior is changed when one chooses a different (but also reasonable) model for the likelihood or the prior, or other assumptions affecting the posterior.

In theory: Setting up a 'super-model' as joint-distribution which includes all possible realities. Posterior of this super-model then automatically incorporates the sensitivity analysis. In practice this is conceptually and computationally infeasible and also is still dependent on the assumptions made when creating the super-model being correct.

One way is just to compare model predictions with the actual outcomes (this is referred to as external validation). If for some reason it is not possible to get the acutal outcomes (for example if they lie in the future), one needs to approximate external validation with the available data.

Posterior predictive checking: Use global summaries of the data distribution and the predictive distribution to evaluate the model: Draw random samples

from the predictive distribution, compare e.g. the smallest of these values with the smallest value in the data...

model checking by calculating test quantities. test quantities are scalar summaries of parameters that are used to compare data to simulations. test quantities are similar to test statistics in the classical approach. Besides on the data, test quantities can also depend on the parameters.

Graphical comparison of simulated and data histograms (!! maybe here point out usefulness of Lumen)

tail-area-probability: p-value p-value in classical statistics: Probability, that, given a parameter θ , a test statistic for replicated data is greater or equal the test statistic for the observed data, as shown in equation 5:

$$p_C = Pr(T(y^{rep}) >= T(y)|\theta) \tag{5}$$

[3]

The Bayesian p-value, on the other hand, is not conditional on a fixed θ , since in Bayesian statistics parameters are drawn from a distribution the same way as outcome variables. Instead, the observed data y is the fixed quantity for the Bayesian p-value, as shown in equation 6

$$p_B = Pr(T(y^{rep}, \theta)) > = T(y, \theta)|y)$$
(6)

4 Probabilistic Programming

stochastic data types (-> probability distributions) make it easier to perform bayesian data analysis

4.1 What are Probabilistic Programming Languages

Modelle spezifizieren/beschreiben

Random Variables can be represented as objects [5]

effizienter in der Beschreibung von Modellen als herkömmliche Programmiersprachen [6]

unifying general purpose programming with probabilistic modeling [7]

A probabilistic reasoning system uses prior knowledge in the form of a probabilistic model to answer a certain query. The particular properties of the query as well as the prior knowledge are given to an inference algorithm which returns an answer in the form of probabilities. Example is shown in figure 1. Probabilistic Programming is the implementation of a probabilistic reasoning system by using a programming language.

Traditional means for representing models are not always sufficient for probabilistic models. Therefore, probabilistic programming languages were introduced to be able to represent models with the full power of a programming language (http://www.probabilistic-programming.org/wiki/Home).

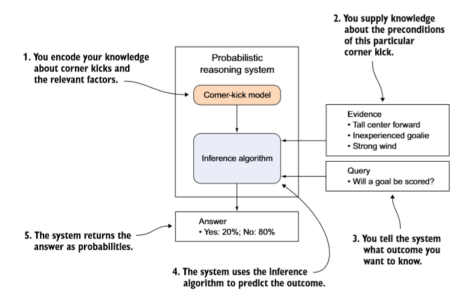


Abbildung 1: General workflow example of a probabilistic reasoning system

4.2 Difference to conventional Programming Languages

When estimating parameters, not only the most likely value for the parameters is given, but also their uncertainty.

5 Lumen

5.1 Functionality

compare to the plotting methods of PyMC3

5.2 Requirements for a PPL

possible criterium: variety of distributions that can be described?

6 Comparing Different Probabilistic Programming Languages

- stan for python: https://pystan.readthedocs.io/en/latest/
- \bullet pymc3: https://docs.pymc.io/notebooks/getting_started.html#Case-study-2:-Coal-mining-disasters
- edward: http://edwardlib.org/getting-started
- pyro: http://pyro.ai/

6.1 Stan for python

Stan is an open-source program written in C++ that is designed for Bayesian inference on user-specified models. It consists of the following components:

- A modeling language that allows users to specify custom models
- An inference engine which uses Hamilton Monte Carlo methods for sampling to get an approximation for the posterior distribution
- An L-BFGS optimizer for finding local optima
- Procedures for automatic differentiation which are able to compute gradients required by the sampler and the optimizer
- Routines to monitor the convergence of parallel chains and compute inferences and effective sample sizes
- Wrappers for Python, R, Julia and other languages

The code for describing a model by using the aforementioned modeling language is divided into six blocks: data, transformed data, parameters, transformed parameters, model, and generated quantities: In data and transformed data, the structure of the input data, along with any constraints, is given. The difference between the former and the latter is that variables in transformed data are functions of other data variables, whereas variables in data are not. For example if in data, a variable x is listed, then in transformed data one can specify a variable

 $y=x^2$. In parameters and transformed parameters, the parameters of the models are described, whereat, as in the data blocks, the latter contains variables that are functions of other parameters. In model, the model structure in the form of prior and likelihood distributions is given. Finally, the block generated quantities can be used to perform simulations and make predictions. In ??, an example from [8] is shown, where the model $y=a_1e^{-b_1x}+a_2e^{-b_2x}$ is fit to data, using the PyStan interface of Stan for Python. Since not all of the aforementioned blocks are mandatory, only four of them are used in the example.

PyStan also provides basic plotting methods for the posterior distributions of the parameters.

is not able to perform inference on discrete parameters. Discrete data and discrete-data models, however, are possible

computes the log-posterior density

[8]

[9]

für den Code evtl. https://github.com/stephen-hoover/presentations zitieren

6.2 Pymc3

PyMC3 is an open-source probabilistic programming framework for Python. The following explanations are taken from [1]. Specification of Bayesian models in PyMC3 is done by encoding the prior, the sampling and the posterior distributions through three types of random variables: Stochastic, deterministic and observed stochastic ones. Stochastic random variables have values which are in part determined randomly, according to a chosen distribution. Commonly used probability distributions like Normal, Binomial etc. are available for this. Deterministic random variables, on the other hand, are not drawn from a distribution, but are calculated by fixed rules from other variables, for example by taking the sum of two variables. Lastly, there are the observed stochastic random variables which are similar to stochastic random variables, except that they get passed observed data as an argument, that should not be changed by any fitting algorithm. This kind of random variable can be used to represent sampling distributions.

PyMC3 mainly uses simulation techniques to draw inference on posterior distributions. It focuses especially on the No-U-Turn Sampler, a Markov Chain Monte Carlo algorithm, that relies on automated differentiation to get gradient information about continuous posterior distributions. PyMC3 also provides basic methods for plotting posterior distributions.

The code piece in ?? shows a simple example of a Bayesian model, taken from [1]. There, the data X1, X2 and Y is used to fit a regression model. First, prior distributions for the model parameters are set up as stochastic random variables, then the regression model itself is specified by a deterministic random variable and lastly the sampling distribution is described by an observed stochastic random variable to which the observed outcome Y is given as a parameter. Finally, the posterior distribution is simulated by drawing 500 samples from it.

```
# Specify model
example_code = """
        data {
                 // Define input data in this block
                 int N;
                 vector[N] x;
                 vector[N] y;
        }
        parameters {
                 // These are random parameters which we want to estimate
                 vector[2] log_a;
                 ordered[2] log_b;
                 real < lower = 0 > sigma;
        }
        transformed parameters {
                 // Create quantities derived from the parameters.
                 vector < lower = 0 > [2] a;
                 vector < lower = 0 > [2] b;
                 a <- exp(log_a);</pre>
                 b <- exp(log_b);</pre>
        }
        model {
                 // Define your model here
                 vector[N] ypred;
                 ypred <- a[1]*exp(-b[1]*x) + a[2]*exp(-b[2]*x);
                 y ~ lognormal(log(ypred), sigma);
                 log_a \sim normal(0,1);
                 log_b ~ normal(0,1);
        }
0.00
\# Pass data to the model. x and y are the observed data
example_dat = {'x':x,'y':y,'N':len(x)}
# Fit model
sm = pystan.StanModel(model_code=example_code)
fit = sm.sampling(data=example_dat, iter=1000, chains=4)
print(fit)
```

Abbildung 2: Example code of a simple Bayesian model using Stan

```
import pymc3 as pm
basic_model = pm.Model()
with basic_model:
        # describe prior distributions of model parameters. Stochastic variables
        alpha = pm.Normal('alpha', mu=0, sd=10)
        beta = pm.Normal('beta', mu=0, sd=10, shape=2)
        sigma = pm.HalfNormal('sigma', sd=1)
        # specify model for the output parameter. Deterministic variable
        mu = alpha + beta[0]*X1 + beta[1]*X2
        # likelihood of the observations. Observed stochastic variable
        Y_obs = pm.Normal('Y_obs', mu=mu, sd=sigma, observed=Y)
# model fitting by using sampling strategies
with basic_model:
# draw 500 posterior samples
trace = pm.sample(500)
pm.summary(trace)
```

Abbildung 3: Example code of a simple Bayesian model using PyMC3

strictly positive priors are transformed with log transformation, so that they are unconstrained, since that is better for sampling

There is the possibility to create own theano functions in python. Gradient based sampling methods don't work for user-defined functions however, except when a gradient is explicitly added.

Similarly pmc3 allows to define own distributions

- 6.3 Edward
- 6.4 Pyro
- 6.5 BUGS and Jags

mentioned in [8] based on graphical models

6.6 Choose the PPL for the task at hand

7 Practical implementation

8 Fallbeispiele

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9 Literatur

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