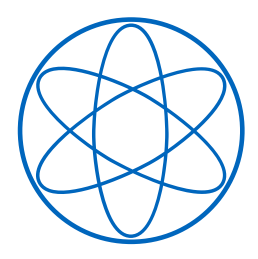


# Development and Benchmarking of a Bayesian Inference Pipeline for LHC Physics



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#### Abstract

In High Energy Physics statistical models are an important component to verify the compatibility of collision events with theoretical predictions. A large portion of these models can be sufficiently described by the statistical framework HistFactory where parameter inference is mainly carried out in a frequentist manner. In contrast, this thesis aims to perform Bayesian inference on the model parameters and derives a formalism to obtain prior distributions from auxiliary measurements. In addition a complete Bayesian inference pipeline is implemented in Python which uses HistFactory models and the Julia implementation of BAT (Bayesian Analysis Toolkit) to carry out parameter inference. The posterior distributions are generated by two MCMC methods – Metropolis-Hastings and Hamiltonian Monte Carlo (HMC) sampling, the latter taking advantage of the automatic differentiation enabled by jax. All required steps for interfacing Julia code in Python (and vice versa) are elaborated in this work for both sampling methods. Furthermore theoretical foundations of MCMC sampling, Bayesian inference and HistFactory are presented. Finally the Python-Julia pipeline is benchmarked against a pure-Julia Bayesian inference implementation for different scenarios and the results are discussed.

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# Abbreviations

Autograd auto differentiation.

**BAT** Bayesian Analysis Toolkit.

**GC** garbage collector.

**HEP** High Energy Physics.

**HF** HistFactory.

**HMC** Hamiltonian Monte Carlo.

**LHC** Large Hadron Collider.

llh log likelihood.

MC Monte Carlo.

MCMC Markov chain Monte Carlo.

MH Metropolis-Hastings.

**NP** nuisance parameter.

**pdf** probability density function.

**POI** parameter of interest.

**XLA** accelerated linear algebra.

vi Abbreviations

# Chapter 1

# Introduction

The experimental evidence of the Higgs boson [1] in 2012 was a breakthrough in particle physics and changed our view on the Standard Model of particle physics. The Higgs boson was discovered in proton-proton collisions with the ATLAS [2] detector at the Large Hadron Collider (LHC) in CERN. The basic building block for detecting the Higgs boson however, is a statistical model which has predicted the Higgs boson mass by hypothesis testing. In case of the Higgs boson, the statistical model was created by HistFactory (HF), a tool for binned statistical analysis which is widely used in LHC physics. In the very most cases, statistical inference in LHC physics is carried out from a frequentist perspective, by performing hypothesis based on the statistical model.

In contrast, this work aims to use Bayesian inference with HistFactory (HF) models for parameter inference. There are three main contributions for this thesis.

- First it derives a formalism to obtain prior distributions from auxiliary measurements
- Second it develops a Python-Julia pipeline which uses the Bayesian Analysis Toolkit (BAT) for parameter inference on HF models and generates posterior distributions by Metropolis-Hastings (MH) and Hamiltonian Monte Carlo (HMC)
- Third it provides run-time benchmarks for the implemented Bayesian inference pipeline

# Chapter 2

# Mathematical Preliminaries

# 2.1 Frequentist versus Bayesian Inference

In statistical inference there are 2 main approaches, the *frequentist* and *Bayesian* inference. Both methods make use of a statistical model  $p(x | \theta)$  to describe observations

$$x \sim p\left(x \mid \theta\right) \tag{2.1}$$

with parameters  $\theta$ .

#### 2.1.1 Frequentist Inference

The frequentist probability interpretation p(x) counts the relative frequency of x occurring during some repeatable process

$$p(x) = \lim_{N \to \infty} \frac{n_x}{N} \tag{2.2}$$

Frequentist inference is *purely* based on the *statistical model* of the observable data  $p(x | \theta)$ , i.e. the likelihood. Parameter inference is performed on the likelihood function  $\mathcal{L}(\theta)$  which is defined for *fixed* data x and only depends on the parameter  $\theta$ .

# 2.1.2 Bayesian Inference

The Bayesian probability interpretation is based on a *prior* believe p(x) that the event x is happening. Parameter inference is obtained by applying Bayes' theorem

$$p(\theta \mid x) = \frac{p(x \mid \theta) p(\theta)}{p(x)}$$
(2.3)

while incorporating prior believes  $p(\theta)$  on the parameters  $\theta$ . In (2.3)  $p(x | \theta)$  is called *likelihood*,  $p(\theta)$  prior, p(x) the evidence and  $p(\theta | x)$  the posterior.

# 2.2 MCMC Sampling

The key difficulty for computing the posterior  $p(\theta|x)$  in Bayes' rule (2.3) is dividing by the evidence p(x). There are only a few special cases where the posterior density can be computed analytically. In most cases p(x) is a high-dimensional, complex density where sampling with standard Monte Carlo (MC) methods is intractable [3].

Instead Markov chain Monte Carlo (MCMC) methods are used to sample from p(x). These methods create a reversible Markov chain that has as an equilibrium distribution which matches the posterior distribution.

#### 2.2.1 Markov Chains

A Markov chain generates a correlated sequence of states  $x_1, \ldots, x_n$ . Each step in the sequence is drawn from a transition operator  $T(x' \leftarrow x)$ , which represents the probability of moving from state x to state x' [3]. According to the Markov property, the transition probability only depends on the current state x. In order to achieve an equilibrium distribution, the Markov chain must fullfil the property of detailed balance

$$T(x' \leftarrow x)p(x) = T(x \leftarrow x')p(x') \qquad \forall x, x' \tag{2.4}$$

This means that transitions under T at equilibrium have the same probability "forwards"  $x \to x'$  and "backwards"  $x' \to x$ . When a Markov chain satisfies (2.4), it is called reversible, as it is statistically indistinguishable whether the chain runs forwards or backwards in time.

#### 2.2.2 Metropolis-Hastings

The Metropolis-Hastings (MH) algorithm [4] is the original MCMC algorithm to generate a set of random numbers that have the properties of a Markov chain and converge towards a target distribution. The MH procedure is summarized in Algorithm 1.

Metropolis-Hastings is initialized by an initial state x and the number of iterations nsteps. For each step we sample proposals for the next state x' from the proposal distribution  $q(x' \leftarrow x)$ . Next, we calculate the acceptance probability a (line 4) which is used to decide whether to accept or reject the proposal. We accept the proposal, if a is greater than a uniformly sampled random number r and update the value of x. Otherwise the proposal is rejected.

It is easy to show that MH satisfies detailed balance in (2.4) [3].

#### **Algorithm 1** Metropolis-Hastings [3]

- 1: Input: initial state x, nsteps
- 2: for  $n = 1 \dots nsteps do$
- 3: Propose  $x' \sim q(x' \leftarrow x)$
- 4: Compute the acceptance probability  $a = \min \left( 1, \frac{p(x')}{p(x)} \frac{q(x' \leftarrow x)}{q(x \leftarrow x')} \right)$
- 5: Accept or reject
  - (a) Draw  $r \sim \text{Uniform}(0, 1)$
  - (b) **if** $(r \le a)$  **then** accept the new state, set  $x \leftarrow x'$  else reject the proposal
- 6: end for

Setting x at the end of each iteration is considered as sample from the density p(x).

In general, MH is valid for any proposal distribution q. "Ideal" proposals would ensure a rapid exploration of the distribution of interest p. There are sophisticated approaches that are based on the observed data [3]. However, q is often chosen to be symmetric and centered around x, e.g. a Normal distribution. Then the acceptance probability (line 4) simplifies to

$$a = \min\left(1, \frac{p(x')}{p(x)}\right) \tag{2.5}$$

MH has a low  $acceptance\ ratio^1$  as the proposals basically perform a  $random\ walk$  in the parameter space.

#### 2.2.3 Hamiltonian Monte Carlo

HMC sampling produces distant proposals for the Metropolis algorithm while avoiding the slow exploration of the parameter space by random-walk proposals [5]. It employs *Hamiltonian dynamics* and the likelihood gradient for an enhanced exploration of the parameter space. Therefore, HMC can only be applied for differential likelihoods.

# Hamiltonian dynamics

Hamiltonian dynamics operates on a 2d dimensional phase space with position vector  $\mathbf{q} \in \mathbb{R}^d$  and momentum vector  $\mathbf{p} \in \mathbb{R}^d$ . The system is fully described by the Hamiltonian

$$\mathcal{H}(q, p) = U(q) + V(p) \tag{2.6}$$

<sup>&</sup>lt;sup>1</sup>the ratio of proposed samples that are accepted

where U(q) is the potential energy and V(p) the kinetic energy of the system. The change of the generalized coordinates q and p over time is determined by the partial derivatives of the Hamiltonian

$$\frac{\mathrm{d}q_i}{\mathrm{d}t} = \frac{\partial \mathcal{H}}{\partial p_i} \qquad \frac{\mathrm{d}p_i}{\mathrm{d}t} = -\frac{\partial \mathcal{H}}{\partial q_i} \qquad \text{for } i = 1, ..., d \qquad (2.7)$$

The kinetic energy in (2.6) is defined as

$$V(\mathbf{p}) = \frac{1}{2} \mathbf{p}^T \mathbf{M}^{-1} \mathbf{p}$$
 (2.8)

where M is a symmetric, positive-definite mass matrix that is typically diagonal. Applying Equations (2.7) to the kinetic energy in (2.8) and U(q) yields

$$\frac{\mathrm{d}q_i}{\mathrm{d}t} = \left[ \mathbf{M}^{-1} \mathbf{p} \right]_i \qquad \frac{\mathrm{d}p_i}{\mathrm{d}t} = -\frac{\partial U(\mathbf{q})}{\partial q_i}$$
 (2.9)

# Probability and the Hamiltonian - Canonical distributions

When using Hamiltonian dynamics to sample from a distribution, the density function needs to be translated to a potential energy function. A solution for this requirement is the *canonical distribution* from statistical mechanics. Given a energy function E(x) for a state x, the canonical distribution has a probability density function (pdf) of [5]

$$p(x) = \frac{1}{Z} \exp\left(-E(x)/T\right) \tag{2.10}$$

where T is the temperature of the system and Z a normalization constant. The Hamiltonian in (2.6) describes the energy for the joint state of position q and momentum p. With (2.10) we obtain the *joint* distribution for q and p as

$$p(\boldsymbol{q}, \boldsymbol{p}) = \frac{1}{Z} \exp\left(-\mathcal{H}(\boldsymbol{q}, \boldsymbol{p})/T\right) = \frac{1}{Z} \exp\left(-U(\boldsymbol{q})/T\right) \exp\left(-K(\boldsymbol{p})/T\right)$$
(2.11)

For the definition of  $\mathcal{H}(q, p)$  in (2.6), the densities for q and p are distributed by *independent* canonical distributions with energies U(q) and K(p).

# **HMC Algorithm**

In Bayesian statistics, we define the potential energy function as [5]

$$U(q) = -\log \left[ p(q \mid x) \ p(q) \right]$$
 (2.12)

where p(q|x) is the likelihood and p(q) the prior density. As the canonical distributions of q and p in (2.11) are independent, we can choose the distribution of the momentum variables p independent of q. A practical choice for the momentum variables

$$p \sim \text{Normal}(0, \Sigma)$$
 (2.13)

is a multivariate zero-mean Gaussian distribution [5].

The HMC sampling is summarized in Algorithm 2

```
Algorithm 2 Hamiltonian Monte Carlo
```

```
1: Input: initial state q_0, nsteps, \Sigma, L, \epsilon
 2: for i = 1 \dots nsteps do
           Draw p \sim \text{Normal}(0, \Sigma)
 3:
           (q^{(0)}, p^{(0)}) = (q_{i-1}, p)
 4:
           for j = 1 \dots L do
 5:
                                                                                                       \triangleright apply L leapfrog steps
                p^{(j-\frac{1}{2})} = p^{(j-1)} - \frac{\epsilon}{2} \nabla U(q^{(j-1)})
 6:
                q^{(j)} = q^{(j-1)} + \epsilon \Sigma p^{(j-\frac{1}{2})}
 7:
                p^{(j)} = p^{(j-\frac{1}{2})} - \frac{\epsilon}{2} \nabla U(q^{(j)})
 8:
           end for
 9:
           (q^*, p^*) = (q^{(L)}, p^{(L)})
10:
           Draw r \sim \text{Uniform}(0, 1)
                                                                                              ▶ Metropolis acceptance step
11:
           a = \min \left(1, \exp\left(-\mathcal{H}(q^*, p^*) + \mathcal{H}(q, p)\right)\right)
12:
           if (r \leq a) then
13:
                (q_i, p_i) = (q^*, p^*)
                                                                                                                                 \triangleright accept
14:
           else
15:
                 (q_i, p_i) = (q_{i-1}, p_{i-1})
                                                                                                                                  \triangleright reject
16:
           end if
18: end for
19: return \{q_i, p_i\}_{i=0}^{nsteps}
```

HMC is initialized with an initial value  $q_0$ , the number of nsteps, a covariance matrix  $\Sigma$ , the number of leapfrog steps L as well as the leapfrog step size  $\epsilon$ . In every step the momentum variable p is drawn from the multivariate Normal distribution. Next, the position and momentum variables q and p are integrated in time by a leapfrog integrator for L steps with step size  $\epsilon$ . The differentials for the numerical integration are derived from (2.9).

The integrated values at step L represent the proposals  $(q^*, p^*)$  which are evaluated in a Metropolis acceptance step similar to Algorithm 1. Here the majority of proposals are accepted since the energy in the system can only be dissipated due to numerical integration errors.

Therefore, HMC has a much higher acceptance rate than MH.

# Chapter 3

# **HistFactory**

HF is a tool for binned statistical analysis which is widely used in LHC physics to measure the consistency of collision events with theoretical predictions. It has been employed for the discovery of the Higgs Boson [1] and is used in searches for new physics [6] by research groups around the planet. The relationship between theoretical predictions and collision events is formalized as statistical model  $p(x | \theta)$ . It describes the probability of observing data x given the model parameters  $\theta$ . Typically, statistical models in High Energy Physics (HEP) are complex with many parameters. HF enables a standardized way to build parametrized probability density functions and infer parameter properties from it.

# 3.1 Formalism

Statistical models in HistFactory describe simultaneous measurements of disjoint channels c (binned distributions as subspace of all collision events), where we observe the event counts n. In a particle detector several physical processes produce events in the selected channels. Hence, the total number of expected events<sup>1</sup> is the sum of all involved processes, the so called samples. This sample rates underlie variations and can be modified by the parameters  $\theta$ . It is distinguished between free parameters  $\eta$  (e.g. the luminosity) and constrained parameters  $\chi$  that account for systematic uncertainties. In a frequentist setting, these constrained terms can be viewed as auxiliary measurements a which result together with the channel events a in the observations a (a). Equation (3.1) illustrates this parametrization

$$p(\boldsymbol{x} \mid \boldsymbol{\theta}) = p(\boldsymbol{n}, \boldsymbol{a} \mid \boldsymbol{\eta}, \boldsymbol{\chi}) \tag{3.1}$$

<sup>&</sup>lt;sup>1</sup>this is often denoted as *event rate* since it used as input parameter to a Poisson distribution

The statistical model in HF consists of two parts – the *main model* of simultaneous measurements over multiple channels, and a *constrained term* that takes into account auxiliary measurements [7].

$$p(\boldsymbol{n}, \boldsymbol{a} \mid \boldsymbol{\eta}, \boldsymbol{\chi}) = \prod_{\substack{c \in \text{channels } b \in \text{bins}_c \\ \text{main model}}} \text{Pois}(n_{cb} \mid \nu_{cb}(\boldsymbol{\eta}, \boldsymbol{\chi})) \qquad \prod_{\substack{\chi \in \boldsymbol{\chi} \\ \text{constraint terms} \\ \text{for "auxiliary measurements"}}} c_{\chi}(a_{\chi} \mid \boldsymbol{\chi})$$
(3.2)

For each bin b and channel c, the total event rate  $\nu_{cb}$  is is the sum over all involved sample rates  $\nu_{scb}$ .

$$u_{cb}(\boldsymbol{\eta}, \boldsymbol{\chi}) = \sum_{s \, \in \, \mathrm{samples}} 
u_{scb}(\boldsymbol{\eta}, \boldsymbol{\chi})$$

The sample event rates  $\nu_{scb}$  are determined by a nominal rate  $\nu_{scb}^0$  and a set of multiplicative and additive rate modifiers  $\kappa(\theta)$  and  $\Delta(\theta)$ , which are controlled by the model parameters  $\theta = (\eta, \chi)$  [7].

$$\nu_{cb}(\boldsymbol{\eta}, \boldsymbol{\chi}) = \sum_{s \in \text{ samples}} \underbrace{\left(\prod_{\kappa \in \boldsymbol{\kappa}} \kappa_{scb}(\boldsymbol{\eta}, \boldsymbol{\chi})\right)}_{\text{multiplicative modifiers}} \left(\nu_{scb}^{0}(\boldsymbol{\eta}, \boldsymbol{\chi}) + \underbrace{\sum_{\Delta \in \boldsymbol{\Delta}} \Delta_{scb}(\boldsymbol{\eta}, \boldsymbol{\chi})}_{\text{additive modifiers}}\right)$$
(3.3)

The available modifiers in HistFactory are summarized in Table 3.1. Each modifier is defined for bin b, sample s and channel c and is controlled by at least one parameter  $\theta \in \{\gamma, \alpha, \lambda, \mu\}$ . By convention, bin-wise modifiers are denoted by  $\gamma$  and interpolation parameters with  $\alpha$ . In contrast, luminosity  $\lambda$  and scale factors  $\mu$  affect all bins equally.

**Table 3.1:** HistFactory modifiers and constraints [7].

Description	Modification	Constraint Term $c_\chi$	Input
Uncorrelated Shape shapesys	$\kappa_{scb}(\gamma_b) = \gamma_b$	$\prod_{b} \operatorname{Pois} \left( r_{b} = \sigma_{b}^{-2}     \rho_{b} = \sigma_{b}^{-2} \gamma_{b} \right)$	$\sigma_b$
Correlated Shape histosys	$\Delta_{scb}(\alpha) = f_p(\alpha \mid \Delta_{scb,\alpha=\pm 1})$	Normal $(a = 0 \mid \alpha, \sigma = 1)$	$\Delta_{scb,\alpha=\pm 1}$
Normalization Uncert.	$ \kappa_{scb}(\alpha) = g_p\left(\alpha   \kappa_{scb,\alpha\pm 1}\right) $	Normal $(a = 0 \mid \alpha, \sigma = 1)$	$\kappa_{scb, \alpha\pm 1}$
MC Stat. Uncertainty staterror	$\kappa_{scb}(\gamma_b) = \gamma_b$	$\prod_{b} \operatorname{Normal}\left(a_{\gamma_{b}} = 1   \gamma_{b}, \delta_{b}\right)$	$\delta_b^2 = \sum_s \delta_{sb}^2$
Luminosity lumi	$ \kappa_{scb}(\lambda) = \lambda $	Normal $(l = \lambda_0 \mid \lambda, \sigma_{\lambda})$	$\lambda_0,\sigma_\lambda$
Normalization normfactor	$\kappa_{scb}(\mu_b) = \mu_b$		
Data-driven Shape shapefactor	$\kappa_{scb}(\gamma_b) = \gamma_b$		

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The first five entries in Table 3.1 are constrained modifiers, defined by a constraint term  $c_{\chi}$  (right part in (3.2)) and an input parameter.

- Uncorrelated Shape modifiers affect each bin individually and hence the sample shape. They are applied to model uncorrelated background, with rate uncertainties  $\delta_b$  for each bin.  $\sigma_b = \delta_b/\nu_b$  is the relative uncertainty of the expected total event rate  $\nu_b$ , and serves as input for the constraint term.
- Correlated Shape modifiers are additive to the nominal sampling rate  $\nu_{scb}^0(\eta, \chi)$  and are controlled by one nuisance parameter (NP). They use the interpolation function  $f_p$  to interpolate between sample distribution shapes represented by a downward variation 10 ( $\alpha = -1$ ) and an upward variation hi ( $\alpha = +1$ ).

The remaining modifiers directly affect the sample rates and are therefore *shape-invariant*.

- The Normalization Uncertainty modifies the sample rate by a factor  $\kappa_{scb}(\alpha)$  which is constructed as interpolation  $g_p$  between an upward "hi" and downward value "lo", for instance "data": {"hi":1.1, "lo":0.9}
- MC Statistical Uncertainty. Many sample counts are derived from Monte Carlo (MC) datasets which necessarily carry uncertainties. These uncertainties are modelled by a set of bin-wise scale factors  $\gamma_b$  while each bin is constraint by a Normal distribution centered at *one*.
- Luminosity. Sample rates that are derived from theory predictions are scaled to the integrated luminosity of the observed data. As this value is derived from measurements that itself carry uncertainties, the luminosity uncertainty  $\sigma_{\lambda}$  needs to be specified.
- Unconstrained Normalization modifiers scale the sample event rate by a free parameter μ. Such parameter are frequently the parameter of interest (POI) of a given measurement.
- Data-driven Shape modifiers are unconstrained, bin-wise multiplicative parameters. They are introduced to support data-driven estimation of sample rates (e.g. multijet backgrounds).

# 3.2 Workspaces

Statistical models in HF are described in plain-text JSON format. This scheme fully specifies the model structure as well as necessary constrained data in a single document and hence is implementation independent. This JSON files represent a *workspace* which consists of *channels*, *measurements* and *observations*.

- Channels are certain regions in the space of collision events, each described by a statistical model. The regions are chosen to be disjoint and typically contain signal regions (SR) and control regions (CR) for a certain particle decay of interest.
  - The statistical model for a channel is constructed by a list of *samples* (models of involved physical processes) which consists of the predicted event rates and a set of modifiers (see Table 3.1).
- Measurements are a small subset of all model parameters the parameter of interest (POI). The measurement scheme in the JSON file can be configured with the initial value, interval bounds or auxdata for the parameter.
- Observations are the actual observed events for each bin in all channels.

For a detailed description of the HF JSON format, the reader is referred to the pyhf documentation [7].

Models in this thesis. Since real world examples are too complex to benchmark in acceptable time, this work mainly uses "toy" workspaces with a single channel. All model specifications utilized in this work are listed in Appendix A.

# 3.3 HistFactory Implementations

Currently, HistFactory is available in three different programming languages

- a C++ implementation,
- a Python version pyhf [8], and
- a Julia implementation LiteHF [9]

In chapter 5, this work employs pyhf and LiteHF for run-time benchmarks. To verify the equivalence of pyhf and LiteHF, the log posterior measure is evaluated for both implementations. The log likelihood in Python for a parameter vector  $\theta$  is computed by the logpdf of the main\_model, i.e.

```
def llh(param: np.ndarray) -> float
    """pyhf log likelihood from the main_model."""
    return model.main_model.logpdf(main_data, param)
```

In LiteHF one can access the log likelihood function by

For visualization purposes, the verification step is illustrated for the 2D 2\_bin\_corr model in section A.2. The prior vector is chosen to be

$$p(\boldsymbol{\theta}) = \begin{bmatrix} \text{Uniform}(0, 5) \\ \text{Normal}(0, 1) \end{bmatrix}$$
(3.4)

and the log posterior measure is evaluated for  $10^5$  points  $x \sim p(\theta)$  with equal initial seed. The samples coincidence for both implementations up to numerical precision. In Figure 3.1 1000 randomly chosen data points are picked out and illustrated for pyhf (blue) and LiteHF (orange).

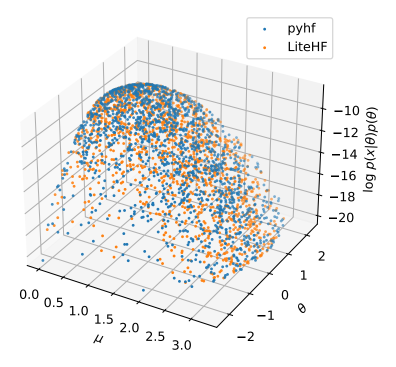


Figure 3.1: Verify the equivalence of the pyhf and LiteHF implementation by evaluating the log posterior measure  $\log \left( p\left( \boldsymbol{x} \,|\, \boldsymbol{\theta} \right) p(\boldsymbol{\theta}) \right)$  at 1000 random points  $\boldsymbol{x} \sim p(\boldsymbol{\theta})$  for the 2\_bin\_corr model (see section A.2). The 2D prior vector is set to  $p(\mu, \boldsymbol{\theta}) = \left[ \text{Uniform}(0, 5), \, \text{Normal}(0, 1) \right]^T$ .

# Chapter 4

# Bayesian Inference with HistFactory

While the majority of statistical results in LHC physics are obtained by a frequentist setting this chapter formalizes a Bayesian approach for parameter inference with HistFactory. First, it introduces the Bayesian Analysis Toolkit (BAT) which is used for MCMC sampling. Second, it derives a formalism to obtain prior distributions from HF models. Third, it discusses implementation details of batty – a Python-to-Julia interface to execute BAT functions in Python. The chapter ends by a review of the implemented Bayesian inference pipeline for the HF models in Appendix A.

# 4.1 BAT

The BAT [10] is a multi-purpose software for Bayesian statistical inference, written in Julia [11]. The software design of BAT is modular and code can be written in a clear and easy fashion. At the same time, it achieves a high run-time performance, which is comparable to C or C++ implementations. Likelihoods, priors and posteriors are all expressed as densities, represented by the type AbstractDensity<sup>1</sup>. To operate on densities, BAT offers function like bat\_sample(), bat\_findmode() or bat\_integrate().

Currently, BAT provides two MCMC sampling algorithms, Metropolis-Hastings (MH) and Hamiltonian Monte Carlo (HMC) which are both applied in this work. Both algorithms have been discussed in section 2.2. For further details the reader is referred to [10].

<sup>&</sup>lt;sup>1</sup>Even density-like objects like log likelihood (llh) functions or histograms are automatically converted to subtypes of AbstractDensity

# 4.2 Priors for HistFactory Models

This section derives the procedure of generating priors for HF models. As discussed in section 3.1, statistical models in HistFactory consist of a main model and constrained terms. Bayesian inference for the parameters  $\theta = (\eta, \chi)$  only uses the likelihood of the main model in (3.2)

$$p_{\text{main}}(\boldsymbol{n} | \boldsymbol{\theta}) \equiv p_{\text{main}}(\boldsymbol{x} | \boldsymbol{\theta}) = \prod_{c \in \text{channels } b \in \text{bins}_c} \text{Pois}(n_{cb} | \nu_{cb}(\boldsymbol{\eta}, \boldsymbol{\chi}))$$
 (4.1)

while the event rate  $\nu_{cb}(\eta, \chi)$  is still computed according to (3.3). Hence, auxiliary measurements  $\boldsymbol{a}$  (see section 3.1) do not appear in the main model  $p_{\text{main}}(\boldsymbol{x} \mid \boldsymbol{\theta})$  and  $\boldsymbol{x}$  corresponds to the observed events  $\boldsymbol{n}$ .

In contrast, priors make use of the auxiliary measurements  $\boldsymbol{a}$  and are derived from the constraint terms  $c_{\chi}(a_{\chi}|\chi)$  in Table 3.1. To formalize this step further, we rewrite Bayes' rule:

$$p(\boldsymbol{\theta} \mid \boldsymbol{x}) = \frac{p_{\min}(\boldsymbol{x} \mid \boldsymbol{\theta}) p(\boldsymbol{\theta})}{p(\boldsymbol{x})}$$

$$\equiv \frac{p_{\min}(\boldsymbol{x} \mid \boldsymbol{\theta}) p(\boldsymbol{\theta} \mid \boldsymbol{a})}{p(\boldsymbol{x})}$$

$$= \frac{p_{\min}(\boldsymbol{x} \mid \boldsymbol{\theta}) p(\boldsymbol{\theta} \mid \boldsymbol{a})}{p(\boldsymbol{x})} \frac{p(\boldsymbol{a} \mid \boldsymbol{\theta}) p_{\text{ur}}(\boldsymbol{\theta})}{p(\boldsymbol{a})}$$

$$= \frac{p_{\min}(\boldsymbol{x} \mid \boldsymbol{\theta})}{p(\boldsymbol{x})} \frac{p(\boldsymbol{a} \mid \boldsymbol{\theta}) p_{\text{ur}}(\boldsymbol{\theta})}{p(\boldsymbol{a})}$$

The prior  $p(\theta \mid a)$  can be interpreted as *posterior* computed from the likelihood of auxiliary measurements  $p(a \mid \theta)$ , an ur-prior  $p_{ur}(\theta)$  and a normalization constant. The last line illustrates the frequentist approach of incorporating auxiliary data – the *green* highlighted part is a product of likelihoods and corresponds to the statistical model in (3.2).

However, as discussed in section 2.2, computing posteriors

$$p(\boldsymbol{\theta} \mid \boldsymbol{a}) = \frac{p(\boldsymbol{a} \mid \boldsymbol{\theta}) p_{\text{ur}}(\boldsymbol{\theta})}{p(\boldsymbol{a})}$$
(4.2)

analytically is only possible for special choices of likelihood and prior. One of these special cases are *conjugate priors*: If the posterior distribution  $p(\boldsymbol{\theta} \mid \boldsymbol{a})$  is in the same distribution family as the prior distribution  $p_{\rm ur}(\boldsymbol{\theta})$ , then prior and posterior are called *conjugate distributions* and  $p_{\rm ur}(\boldsymbol{\theta})$  is called *conjugate prior* for the likelihood  $p(\boldsymbol{a}|\boldsymbol{\theta})$ . This method is applied in this work and elaborated in the next section.

#### 4.2.1 Conjugate Priors

When using conjugate priors, we necessarily incorporate information about the parameter space due to the ur-prior. Nevertheless, to justify this approach we choose a *vague* ur-prior so that the updated prior  $p(\theta \mid a)$  is mainly determined by data likelihood  $p(a \mid \theta)$ .

HistFactory basically uses two types of likelihoods in the constraint terms: On the one hand Poisson constrains for **shapesys** modifiers, on the other hand Normal distributions for the remaining constraint terms (see Table 3.1). Below, conjugate priors for both likelihood types are derived.

#### Poisson distributed Likelihoods

Conjugate priors for Poisson likelihoods are gamma-distributed  $p(\theta) \equiv \text{Gamma}(\alpha, \beta)$ . The pdf for a random variable  $X \sim \text{Gamma}(\alpha, \beta)$  in shape-rate parameterization is given as

$$p(x; \alpha, \beta) = \frac{x^{\alpha - 1} e^{-\beta x} \beta^{\alpha}}{\Gamma(\alpha)} \quad \text{for } x > 0$$
 (4.3)

where  $\alpha, \beta > 0$  are the *shape* and *rate* parameter, and  $\Gamma(\alpha)$  the *gamma function*. The expected value and variance of  $X \sim \text{Gamma}(\alpha, \beta)$  is computed to

$$\mathbb{E}[X] = \frac{\alpha}{\beta}$$
 and  $\operatorname{Var}[X] = \frac{\alpha}{\beta^2}$  (4.4)

The Poisson likelihood of observing N independent and identically distributed events is

$$p(\boldsymbol{x} \mid \theta) = \prod_{i=1}^{N} \operatorname{Pois}(x_i \mid \theta) \quad \text{with} \quad \operatorname{Pois}(x_i \mid \theta) = \frac{\theta^{x_i} e^{-\theta}}{x_i!}$$
 (4.5)

According to Bayes' theorem (2.3) the posterior is calculated to be

$$p(\theta \mid \boldsymbol{x}) = \frac{p(\boldsymbol{x} \mid \theta) p(\theta)}{p(\boldsymbol{x})} \propto p(\boldsymbol{x} \mid \theta) p(\theta)$$
(4.6)

while we neglect the constant evidence term  $p(\mathbf{x})$ . Explicitly computing the posterior using (4.5) and (4.3) for priors  $p(\theta) = \text{Gamma}(\alpha, \beta)$  yields

$$p(\theta \mid \boldsymbol{x}) \propto \prod_{i=1}^{n} \frac{\theta^{x_i} e^{-\theta}}{x_i!} \cdot \frac{\theta^{\alpha - 1} e^{-\beta \theta} \beta^{\alpha}}{\Gamma(\alpha)}$$

$$\propto \theta^{x_1 + x_2 + \dots + x_N} e^{-\theta N} \cdot \theta^{\alpha - 1} e^{-\beta \theta}$$

$$= \theta^{N\bar{x} + \alpha - 1} e^{-(\beta + N)\theta}$$

$$(4.7)$$

In line 2 we skipped the constant terms  $\Gamma(\alpha)$ ,  $x_i!$ ,  $\beta^{\alpha}$  and in line 3 the sample mean

$$\bar{x} = \frac{1}{N} \sum_{i=1}^{N} x_i \tag{4.8}$$

was introduced to simplify notation. Comparing (4.7) with the Gamma pdf in (4.3) and using the argument that the total area under a probability density function must evaluate to *one*, we can conclude that the posterior  $p(\theta \mid x)$  is again a Gamma distribution

$$p(\theta \mid \boldsymbol{x}) = \operatorname{Gamma}(N\bar{x} + \alpha, \beta + N) \tag{4.9}$$

with the updated parameters

$$\alpha' = N\bar{x} + \alpha$$

$$\beta' = \beta + N \tag{4.10}$$

# Normally distributed Likelihoods

Conjugate priors for normally distributed likelihoods are *normally* distributed likewise. To proof this, we consider the likelihood

$$p\left(\boldsymbol{x}\mid\boldsymbol{\theta}\right) = \prod_{i=1}^{N} \frac{1}{\sqrt{2\pi\sigma_{x}^{2}}} \exp\left(-\frac{(x_{i}-\boldsymbol{\theta})^{2}}{2\sigma_{x}^{2}}\right) \propto \exp\left(-\frac{\sum_{i=1}^{N} (x_{i}-\boldsymbol{\theta})^{2}}{2\sigma_{x}^{2}}\right)$$
(4.11)

of N independent observations  $\mathbf{x} = \{x_i\}_{i=1}^N$  with same mean  $\theta$  and variance  $\sigma_x^2$ . During the derivation it is assumed that the likelihood variance  $\sigma_x^2$  is known.

For normally distributed priors

$$p(\theta) = \text{Normal}(\theta_0, \sigma_{\theta}^2)$$
 (4.12)

with mean  $\theta_0$  and variance  $\sigma_{\theta}^2$ , the posterior is again computed according to (4.6). Once more we neglect constant terms and after elementary calculus we obtain the expression

$$p(\theta \mid \boldsymbol{x}) \propto \exp\left(-\frac{\sum_{i=1}^{N} (x_i - \theta)^2}{2\sigma_x^2}\right) \cdot \exp\left(-\frac{(\theta - \theta_0)^2}{2\sigma_\theta^2}\right)$$

$$= \exp\left[\frac{-\sum_{i=1}^{N} x_i^2 + 2\theta N\bar{x} - N\theta^2}{2\sigma_x^2} - \frac{\theta^2 - 2\theta\theta_0 + \theta_0^2}{2\sigma_\theta^2}\right]$$

$$\propto \exp\left[\frac{2\theta N\bar{x} - N\theta^2}{2\sigma_x^2} - \frac{\theta^2 - 2\theta\theta_0}{2\sigma_\theta^2}\right]$$
(4.13)

while we applied the sample mean  $\bar{x}$  (Equation (4.8)) in line 2 to simplify notation. Rewriting (4.13) in powers of  $\theta$  yields

$$p(\theta \mid \boldsymbol{x}) \propto \exp \left[ -\frac{\theta^2}{2} \underbrace{\left( \frac{1}{\sigma_{\theta}^2} + \frac{N}{\sigma_x^2} \right)}_{1/\sigma_{\theta}'^2} + \theta \left( \frac{\theta_0}{\sigma_{\theta}^2} + \frac{N\bar{x}}{\sigma_x^2} \right) \right]$$
(4.14)

Equation (4.14) is proportional to a Normal distribution which can be verified by introducing the variables

$$\sigma_{\theta}^{\prime 2} = \left(\frac{1}{\sigma_{\theta}^2} + \frac{N}{\sigma_x^2}\right)^{-1} \quad \text{and} \quad \theta' = \sigma_{\theta}^{\prime 2} \left(\frac{\theta_0}{\sigma_{\theta}^2} + \frac{N\bar{x}}{\sigma_x^2}\right)$$
(4.15)

By inserting these variables and multiplying (4.14) by a normalization constant

$$p(\theta \mid \boldsymbol{x}) \propto \exp\left[-\frac{\theta^2}{2\sigma_{\theta}'^2} + \frac{\theta\theta'}{\sigma_{\theta}'^2} - \frac{\theta'^2}{2\sigma_{\theta}'^2}\right] = \exp\left[-\frac{(\theta - \theta')^2}{\sigma_{\theta}'^2}\right]$$

we can conclude<sup>2</sup> that the posterior is normally distributed with the parameters in (4.15)

$$p(\theta \mid \boldsymbol{x}) = \text{Normal}(\theta', \sigma_{\theta}'^2)$$

#### 4.2.2 Implementation

After deriving the parameter updates for the conjugate priors, this section describes implementation details. Generating priors from HF models is implemented in the Python package priorhf which was developed during this work.

For all NP, auxiliary data in HF is stored in a member variable auxdata. The parameter order for pyhf models is determined by the parameter map

which contains all required parameter information (auxdata, parameter name, standard deviation sigma (only for staterror and lumi modifiers)) as well as the slice in which the parameter is located.

• For shapesys modifiers (uncorrelated shape), the auxdata entry corresponds to the rate parameter  $r_b = \sigma_b^{-2}$  (Table 3.1, row 1), where  $\sigma_b = \delta_b/\nu_b$  is the relative uncertainty of the expected total event rate  $\nu_b$  [7]. This uncertainties are modeled by a Poisson  $\tilde{\gamma}_b \sim \text{Pois}(r_b)$  with expectation  $\mathbb{E}[\tilde{\gamma}_b] = r_b$ . As initial prior (ur-prior)

<sup>&</sup>lt;sup>2</sup>Again, we use the argument that the area under a pdf must be equal to one.

we choose  $p_{\rm ur}(\tilde{\gamma}_b) = {\rm Gamma}(1,\beta)$ , with  $\alpha = 1$  and compute the rate parameter  $\beta$  of the initial prior so that the expectation of likelihood and initial prior match, i.e.

$$\beta = \frac{\alpha}{r_b} \tag{4.16}$$

Then, we perform the parameter update in (4.10) for N=1,  $\bar{x}=r_b$  and obtain an up-scaled prior

$$p(\tilde{\gamma}_b \mid \boldsymbol{a}) = \operatorname{Gamma}(\alpha', \beta') \text{ with } \mathbb{E}[\tilde{\gamma}_b] = r_b$$

In a final step, we re-scale the density  $p(\tilde{\gamma}_b | \boldsymbol{a})$  so that it has an expected value of 1 by multiplying  $\beta'$  with  $r_b$ . To summarize, the prior distribution for a shapesys parameter  $\gamma_b$  is given by

$$\gamma_b \sim \text{Gamma}(r_b + 1, r_b + 1) \quad \text{with} \quad \mathbb{E}[\gamma_b] = 1$$
 (4.17)

which is derived from an ur-prior  $p_{\rm ur}(\tilde{\gamma}_b) = {\rm Gamma}(\alpha = 1, \beta = 1/r_b)$ .

The priors for the 2\_bin\_uncorr model<sup>3</sup> are illustrated in Figure 4.1a). The relative uncertainties  $\sigma_b$  for bin 1 and 2 are 10 % and 20 % (see section A.1). This results in a tighter prior distribution for  $\gamma_1$  and a broader spread for the parameter  $\gamma_2$ .

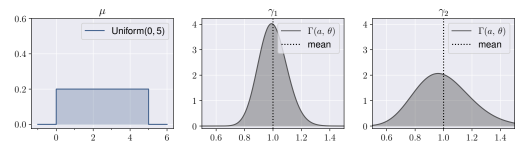
• The modifiers histosys and normsys are constrained by a zero-mean Normal distribution with variance 1 (see Table 3.1). The corresponding values in the auxdata vector are zeros. The initial prior is chosen as zero-mean Normal distribution with variance  $\sigma_{\theta}^2 = 100$ . After the parameter update in (4.15) (with N = 1,  $\bar{x} = \theta_0 = 0$  and  $\sigma_x^2 = 1$ ) we obtain the prior distribution

$$p(\theta \mid \boldsymbol{a}) = \text{Normal}\left(0, \ \sigma_{\theta}^{\prime 2} = \frac{100}{101}\right) \tag{4.18}$$

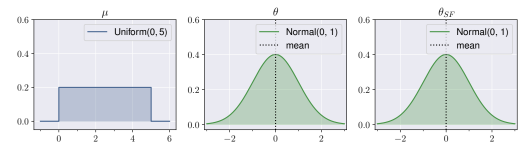
Priors for the 4\_bin model<sup>4</sup> are illustrated in Figure 4.1b) as standard normal distribution for the parameters  $\theta$  and  $\theta_{SF}$ .

<sup>&</sup>lt;sup>3</sup>a 2-bin model with uncorrelated background

<sup>&</sup>lt;sup>4</sup>3D parameter space with normsys and histosys modifiers



(a) Prior distributions for the 2\_bin\_uncorr model (section A.1) parameters



(b) Prior distributions for the 4\_bin model (section A.3) parameters

Figure 4.1: Conjugate priors for the 2\_bin\_uncorr and 4\_bin model. The signal strength parameter  $\mu$  (left) is uniformly distributed for both models. The dashed line represents the mean of the Gamma and Normal distribution. Priors for the 2 dimensional 2\_bin\_corr model (section A.2) are equal to the first two distributions of the 4\_bin model.

• staterror and lumi modifiers are also constrained by a Normal distribution (see Table 3.1). Their mean values are 1 or  $\lambda_0$  and are stored in auxdata. The ur-prior variance  $\sigma_{\theta}^2 = 100$  is again set to 100 and the mean  $\theta_0$  corresponds to the value in auxdata. The likelihood variance  $\sigma_x^2$  is computed from the standard deviation sigma in the parameter map. Once more the parameter update is calculated according to Equation (4.15) with N = 1.

Parameter of interest (POI) (like the signal strength  $\mu$ ) are unconstrained and modeled as uniform distribution from 0 to 5. All prior distributions are implemented as NamedTupleDist [12] of densities from the Distributions.jl package [13]. This step is necessary since priors directly interface to BAT which requires this type. Before converting priors to a NamedTupleDist, priors need to be represented as namedtuple<sup>5</sup> to preserve the parameter order<sup>6</sup>.

 $<sup>^{5} {</sup>m from}$  collections import namedtuple

<sup>&</sup>lt;sup>6</sup>Passing Python dictionaries to the NamedTupleDist constructor interchanges the parameter order even for Python versions  $\geq 3.7$ . This is because when a Python dict is converted to a Julia Dict, the order is not maintained.

This step is illustrated in the snipped below, where names is a list of the parameter names and param a list of Distributions.jl densities.

```
# Makro to convert keys of a dict as parameter string for namedtuple
key_str = lambda d: ' '.join(list(d.keys()))

prior_specs = dict(zip(names, param))
p = namedtuple('Prior', key_str(prior_specs))(**prior_specs)
priors = jl.unshaped(jl.NamedTupleDist(p))
```

In the last line the unshaped [12] function is applied to obtain a *flat* prior vector.

# 4.3 batty – BAT to Python Interface

The complete pipeline for calling BAT functions from Python is implemented in batty, the BAT to Python interface which is developed here [14]. During this work, batty was steadily tested an improved with several contributions. An old version of batty used PyCall.jl [15] to execute Julia functions from Python which had one major drawback: The complete Python environment needed to be compiled each time batty was imported which takes about 2 minutes. The current version of batty employs PythonCall.jl [16] that reduces the total import time to about 10 seconds and speed up inference in batty by a factor of 4.

PythonCall.jl is suited to call Python functions in Julia and vice versa [16]. The corresponding Python package is called juliacall and is used in Python as

```
from juliacall import Main as jl
jl.seval('using BAT, Distributions')
```

The second line imports the Julia modules BAT [10] and Distributions [13] which enables the access to Julia types or functions using the prefix jl.<type/function>.

#### 4.3.1 Likelihoods for pyhf Models

All functionalities for computing pyhf likelihoods are implemented in the Python package pyhf\_llh. For a fast computation of the log likelihood (llh) this work makes use of the jax-backend of pyhf<sup>7</sup>. jax [17] exploits auto differentiation (Autograd) and accelerated linear algebra (XLA) to speed up computations and automatically compute gradients. It is developed by Google and is widely used in Machine Learning frameworks.

 $<sup>^{7}\</sup>mathrm{pyhf}$  can be configured to use different  $tensor\ backends$  (NumPy, PyTorch, TensorFlow and JAX) for all numerical computations.

This work aims to implement MCMC sampling in Python for Metropolis-Hastings (MH) and Hamiltonian Monte Carlo (HMC) sampling. Since HMC requires the likelihood gradient, these values need to be passed from Python to Julia. This is accomplished by using a PyCallDensityWithGrad wrapper that contains pointers to the log likelihood (llh) function logf as well as a pointer to the llh function with gradient valgradlogf, i.e.

```
struct PyCallDensityWithGrad <: BAT.BATDensity
    logf::PythonCall.Py
    valgradlogf::PythonCall.Py
end</pre>
```

In addition, several steps have to be considered to pass gradient from Python to Julia. For details see pybat.jl in [14]. The pyhf likelihood is implemented as functor that returns the llh function for a given HF workspace

To enhance the computation time, we employ the just-in-time compilation (jit) by jax which returns highly optimized machine code for computing the log likelihood efficiently. The second entry for the PyCallDensityWithGrad wrapper is the llh\_with\_grad function

```
def pyhf_llh_with_grad(ws: str) -> Tuple[Callable, Callable]:
    """Returns the Tuple (llh, llh_with_grad) for the pyhf workspace 'ws'"""
    # jit llh and compute grad
    llh = pyhf_llh(ws)
    llh_grad = jit(grad(llh))
    # convert to float and catch out 'nan' values of pyhf
    llh_float = lambda x: -inf if jnp.isnan(llh(x)) else float(llh(x))

def llh_with_grad(param: np.ndarray) -> Tuple[float, np.ndarray]:
    """Convert jax.DeviceArray to (float, np.float64)"""
    return (llh_float(param), np.array(llh_grad(param), dtype=np.float64))

return (llh_float, llh_with_grad)
```

which returns a tuple (llh\_float, llh\_with\_grad) required by PyCallDensityWithGrad. The gradient of the likelihood is computed by the grad() function from jax. To ensure that the return types can be converted with PythonCall.jl, jax.DeviceArray types need to be converted to float types. In addition it is necessary to catch nan values of pyhf and replace them by -inf since bat\_sample() does not work for nan values.

# 4.3.2 pyhf Benchmarks

The effects of using the jax backend instead numpy are discussed below for the HF models 2\_bin\_corr and 2\_bin\_uncorr (Appendix A). The compile time of bat\_sample() for both cases is shown in Figure 4.2 (*left*). Using the jax backend slightly reduces the compile time for both models. In contrast the memory usage during compilation increases by about 10% (Figure 4.2 *right*) for the jax backend.

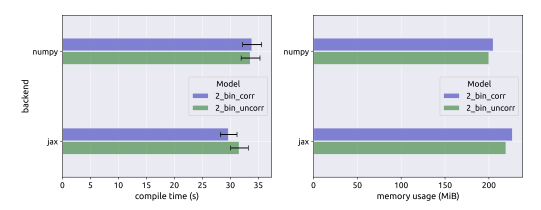


Figure 4.2: Compile time and memory usage while compiling bat\_sample() with jax and numpy backend. Compile times typically fluctuate between  $\pm 5\%$ .

However, more important is the run-time performance. Figure 4.3a) shows the run-time with numpy backend for different MCMC steps in bat\_sample(). The run-time statistics is visualized as box-plot as described in Appendix B. For the numpy setting, the 2\_bin\_corr is about two times slower than the 2\_bin\_uncorr model by performing 1k, 5k or 10k MCMC nsteps. With jax backend, the total run time can be reduced by a factor of 3 to 5 (see Figure 4.3b). In addition the run-time of both models is roughly the same using the jax backend.

Benchmarks for directly evaluating the llh for jax and numpy are given in Figure B.2.

Unless stated differently this work uses the jax backend for evaluating the log likelihood.

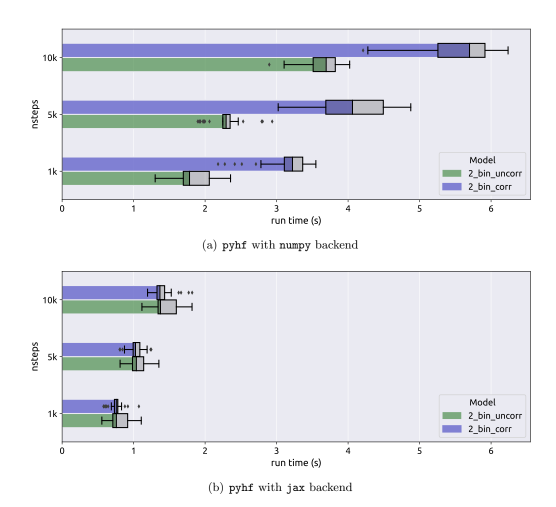
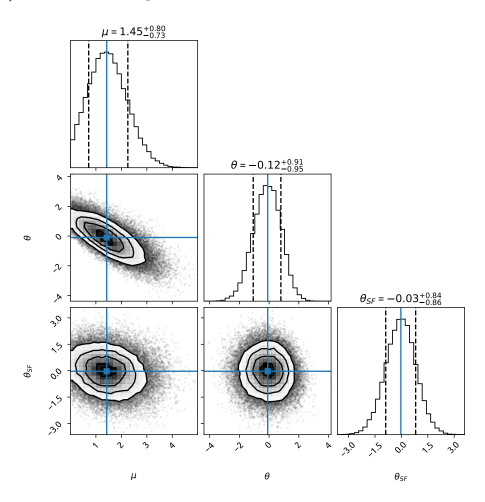


Figure 4.3: Run-time performance of bat\_sample() once using the numpy backend and once with jax backend. The run-time is evaluated for the models 2\_bin\_corr and 2\_bin\_uncorr for different MCMC nsteps.

# 4.4 Bayesian Inference Examples

After setting up the priors and likelihood in section 4.2 and section 4.3, the complete Bayesian inference pipeline is verified for the HF models in Appendix A. As an example, the sampled posterior distribution for the 4-bin model is illustrated in Figure 4.4 as corner plot [18] for 100k MCMC steps.



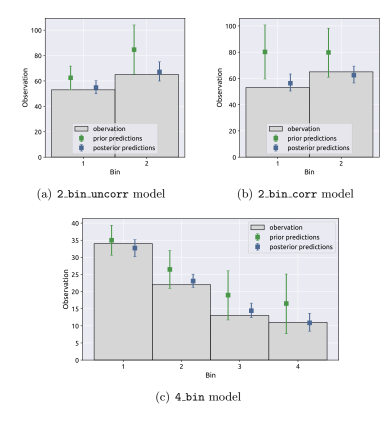
**Figure 4.4:** Corner plot [18] for the 4\_bin Model (section A.3) using 100k HMC steps. The parameter mode is indicated by *blue* lines and the *dashed* lines represent the  $1\sigma$  quantile of the sampled distributions.

A snipped for sampling this posterior in Python is given below

```
likelihood = jl.PyCallDensityWithGrad(llh, llh_grad)
prior = make_prior('path/to/model')
posterior = jl.BAT.PosteriorMeasure(likelihood, prior)
samples = jl.bat_sample(posterior, method).result
```

where method is the MCMC sampling algorithm – Metropolis-Hastings or HMC.

The sampled posteriors are verified by a *posterior predictive check*, which is depicted in Figure 4.5 for all three models. All models sufficiently represent the observations with a reduced variance compared to the prior predictions.



**Figure 4.5:** Prior and posterior predictive checks all models in Appendix A. The square represents the median of the predicted output and the error bars indicate the  $1\sigma$  quantile.

### Chapter 5

# Benchmarking the Python-Julia Pipeline against the Julia Implementation

This chapter compares the run-time performance of batty (section 4.3) with a pure-Julia implementation of Bayesian inference with LiteHF [9]. More specific, the following three implementations are evaluated

Python + BAT (visualized in blue)
 Uses the Python-Julia bridge batty for Bayesian inference on HF models in Python.

• Julia + LiteHF (visualized in green)

The complete inference chain is written and executed in Julia by using LiteHF [9], the Julia implementation of HF.

• Julia + pyhf (visualized in red)

The whole code runs in Julia but uses the Python likelihood from pyhf for inference. Executing Python code from Julia is accomplished by the PythonCall.jl module [16]. The pyhf likelihood is implemented in the pyhf\_llh package that can be embedded in Julia by

```
llh = pyimport("pyhf_llh")
llh_func, llh_with_grad = llh.pyhf_llh_with_grad("path/to/model")
```

Run-times benchmarks are measured for both, MH and HMC based on two HF models.

### 5.1 Benchmark Setup

For each implementation the run-time of bat\_sample() is measured. Therefore, bat\_sample() takes two arguments, the posterior to be sampled and the MCMC method to be used.

```
# Metropolis Hastings
method = BAT.MCMCSampling(mcalg=BAT.MetropolisHastings(), nsteps=nsteps, nchains=2)
# or Hamiltonian MC
method = BAT.MCMCSampling(mcalg=BAT.HamiltonianMC(), nsteps=nsteps, nchains=2)
samples = bat_sample(posterior, method).result
```

The number of chains nchains is set to 2 for all benchmarks. The *complexity* of the MCMC sampling is controlled by the number of nsteps. It is *not* considered whether all chains converge or not.

In order to ensure the reproducibility of benchmarks, the following steps are taken into account

- Code that is executed by bat\_sample() like computing the log likelihood does not use global variables. Accessing global variables inside a function increases the run-time in Julia and Python.
- Since Julia code is pre-compiled before execution, a distinction is made between the *compile time* and *run-time* of bat\_sample(). The proportion of both times is illustrated in Figure 5.1 for the 2\_bin\_corr model for sampling 10k MCMC steps. The compile time accounts for more than 90% of the total execution time in all implementations and has a uncertainty of ±4%. Compiling bat\_sample() in Python takes about twice as long as in Julia (see Figure 5.1 *left*).
- Benchmarks in Julia are obtained using the BenchmarkTools.jl module [19]. To measure "long" run-times sufficiently, the @benchmarkable maken is configured as

```
b = @benchmarkable bat_sample(posterior, mcalg) setup=(mcalg=$method)
sec = bootstrap_sec()
res = run(b, samples=samples, seconds=maximum([sec, 40]))
```

where the required time sec is bootstrapped from previous run-time measurements.

- All run-time benchmarks are written in Python or Julia scripts, since Jupyter Notebooks have lots of overhead.
- The benchmarks are carried out on a Linux computer with a Intel(R) Core(TM) i7-7700HQ CPU.

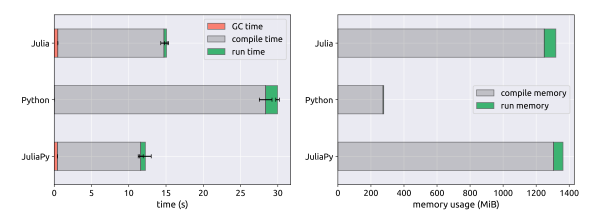


Figure 5.1: Total execution time and memory usage of bat\_sample() for the 2\_bin\_corr model with 10k steps. The red, gray and green portion is the garbage collector (GC) time, the compile time and run time respectively. Compiling bat\_sample() in Python takes about twice as long as in Julia. In contrast, the memory usage during compilation in Python is only one sixth of the memory usage in Julia. (The run-time memory usage in Python can not be measured sufficiently and hence is neglected in the right plot. Moreover the GC time can not be evaluated in Python.)

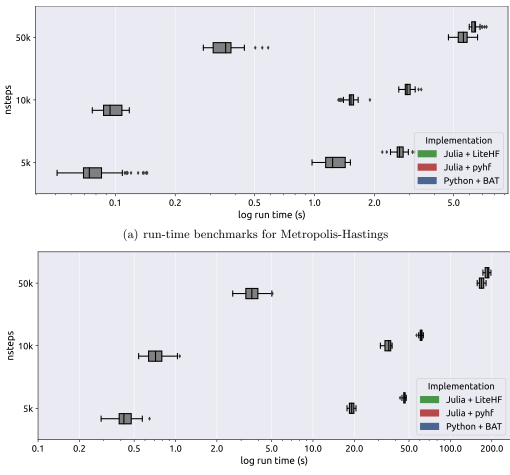
#### 5.2 Benchmarks

#### 5.2.1 2\_bin\_corr Model

First, the run-time of the 2\_bin\_corr model (with a 2D parameter space) is evaluated for different MCMC nsteps. The results are visualized in Figure 5.2a) for Metropolis Hastings and in Figure 5.2b) for Hamiltonian MC. The run-time on the x-axis is plotted logarithmically and the numerical values are summarized Table 5.1. For confirmation, the run-time benchmarks for the 4\_bin model is additionally given in Appendix C.

**Table 5.1:** Summary of run-time benchmarks in Figure 5.2. The values represent the median run-time with  $1\sigma$  standard deviation without outliers for 5k, 10k and 50k nsteps.

		5k	10k	50k	
Method	Implementation	${f run\text{-}time}\ (\mathrm{s})$			
	$\mathrm{Julia} + \mathtt{LiteHF}$	$0.07 \pm 0.01$	$0.09 \pm 0.01$	$0.35 \pm 0.04$	
$\mathbf{MH}$	Julia + pyhf	$1.23 \pm 0.16$	$1.52 \pm 0.10$	$5.59 \pm 0.43$	
	Python + BAT	$2.67 \pm 0.20$	$2.91 \pm 0.15$	$6.39 \pm 0.30$	
	$\mathrm{Julia} + \mathtt{LiteHF}$	$0.42 \pm 0.07$	$0.71 \pm 0.10$	$3.59 \pm 0.50$	
$\mathbf{HMC}$	Julia + pyhf	$19.1 \pm 2.0$	$35.6 \pm 2.3$	$168 \pm 9$	
	Python + BAT	$45.9 \pm 1.3$	$61.2 \pm 1.9$	$186 \pm 8$	



(b) run-time benchmarks for Hamiltonian MC sampling

**Figure 5.2:** Run-time benchmark for the 2\_bin\_corr model for different MCMC nsteps with logarithmic time axis. Uncertainties are visualized as box-plot as illustrated in Figure B.1. The run-time in Julia is averaged over 100 runs, for the other two implementations over 40 runs for MH and 10 runs for HMC sampling.

Metropolis Hastings. The run-time of the pure-Julia implementation (green) is about 20 times faster than the Python-Julia pipeline (blue) for all considered steps. Executing the pyhf likelihood from Julia (red) is about 12 times slower that the pure-Julia implementation. For an increasing number of samples (nsteps) the run-time of Julia + pyhf approaches the execution time of batty (Python + pyhf).

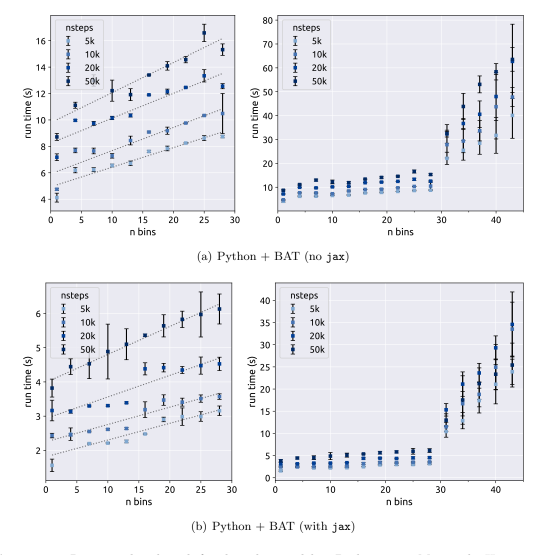
Hamiltonian MC. Executing HMC in Python (Figure 5.2b) is about 100 times slower than the pure-Julia implementation (green) for 5k and 10k steps, and about 50 times slower for 50k steps. Again, calling the pyhf likelihood from Julia (red) outperforms the batty implementation (blue). For a small number of nsteps=[5k, 10k] the Julia version (Julia + pyhf) is about two times faster than batty. For 50k steps both implementations require approximately the same execution time.

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#### 5.2.2 n-Bin-Model

The runtime measurements in this section examine how the dimensionality of the parameter space affects the run-time. The n-bin model is implemented as simplemodel with uncorrelated background according to the  $2\_bin\_uncorr$  model in section A.1. Repeatedly appending bins to the histogram is used as a simple method to increase the dimensionality of the inference problem (n bins corresponds to a (n+1)-dimensional parameter vector).

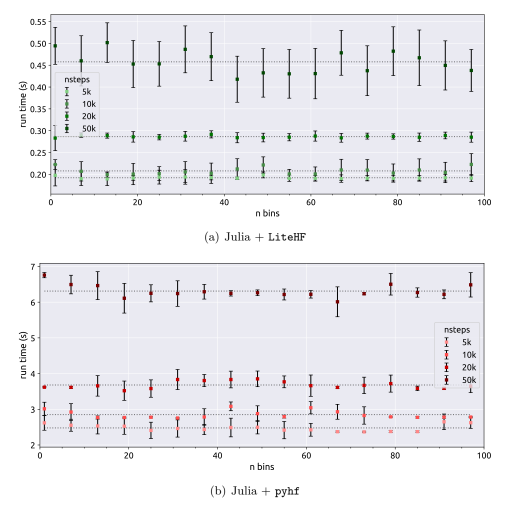
#### Metropolis-Hastings (MH)



**Figure 5.3:** Run-time benchmark for the n-bin model in Python using Metropolis Hastings. A model with n bins represents a (n+1)-dimensional parameter vector. The upper Figure uses a Float64 likelihood with no just-in-time compilation by jax, the likelihood in the lower Figure is jitted and has Float32 precision.

Figure 5.3 shows the run-time of bat\_sample() in batty (Python + BAT) for an increasing number of bins using a jitted and not jitted likelihood<sup>1</sup>. In both plots the run-time slightly increases for models with less than 30 bins (left). For larger models (n > 30 bins), the run-time in Python increases more sharply. The run-time for a 40-bin model already takes about 5 times longer than a 30-bin model.

The Julia benchmarks for the n-bin model are illustrated in Figure 5.4. The pure-Julia version (green) is more than 10 times faster than the the jitted Python implementation in Figure 5.3b). Moreover, run-times stay constant up to a 100 dimensional parameter vector for different number of steps. The same behavior can be observed for calling the pyhf likelihood from Julia (Figure 5.4b).



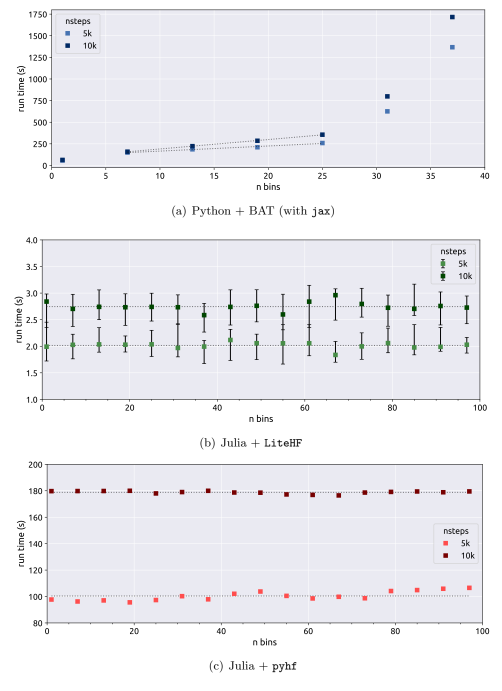
**Figure 5.4:** Run time benchmark for the n-bin model in Julia using Metropolis Hastings. The Julia + LiteHF version (top) is fully implemented in Julia and outperforms the Julia + pyhf implementation (bottom) by a factor of 12.

 $<sup>^{1}</sup>$ The jitted likelihood is about 2.5 times faster for the n-bin model.

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### Hamiltonian Monte Carlo (HMC)

The HMC run-times for the n-bin model are illustrated in Figure 5.5 for 5k and 10k steps.



**Figure 5.5:** Run time benchmark for the n-bin model using HMC for 5k and 10k nsteps. The Julia run-times (top) are averaged over 100 runs, the other two run-times represent a single run.

Similar to MH in Figure 5.3, run-times for the batty implementation (Figure 5.5a) slightly

increase for n-bin models with less than 30 parameters. In this range HMC in batty is about 100 times slower than in pure Julia (green). For n > 30 bins the batty run-time increases significantly, as already noted for Metropolis Hastings in Figure 5.3.

The run-times for the Julia implementations are shown in Figure 5.5b) and c). Again run-times stay approximately constant up to a 100-bin model.

### 5.3 Discussion

In all considered benchmarks Julia + LiteHF clearly outperforms the other two implementations that call the pyhf likelihood. This is, however, to be expected as pure-Julia code is fully compiled to fast machine code while Python code is executed by a high-level interpreter. For the 2\_bin\_corr and 4\_bin model (Appendix C) the run-time performance of batty is similar to the Julia + pyhf implementation for large numbers of MCMC steps. On the other hand for nsteps \leq 10k, calling the pyhf likelihood from Julia is almost twice as fast as for the batty implementation. Again this can be explained by the interpreter overhead of Python which becomes relatively smaller when the run-time of the Julia function bat\_sample() dominates. For sufficiently small parameter spaces we can summarize that Metropolis-Hastings in batty is about 10 times slower than a pure Julia implementation and for HMC batty is even slower by a factor of 100.

With the *n*-bin model the run-time was examined as a function of the parameter dimension. However for both Julia implementations (pure Julia and calling pyhf from Julia) the run-time remains *constant* up to a 100 dimensional parameter space for MH and HMC. The log likelihood for the *n*-bin model appears to be computationally too cheap to measure the influence of the parameter dimension on the run-time in Julia. As a result, other computational load in Julia is predominant which keeps the run-time constant up to 100 bins. In the pure Julia version this is very likely the overhead of the bat\_sample() method. For Julia + pyhf the time for converting Julia types to Python types and vice versa seems to be the time-intensive part.

In Python, on the other hand, the run-time rapidly increases for parameter dimensions greater than 30. Within this thesis task it was not possible to determine the exact reason for this behavior in Python. Still it can be *excluded* that the Float32 precision of jax is an issue, since the Float64 numpy likelihood in Figure 5.3a) shows the same behavior.

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In addition, the run-time of the *n*-bin log likelihood was evaluated for **pyhf** (with and without **jax**) as well as for **LiteHF**. The run-time benchmark is visualized as semi-logarithmic plot in Figure 5.6.

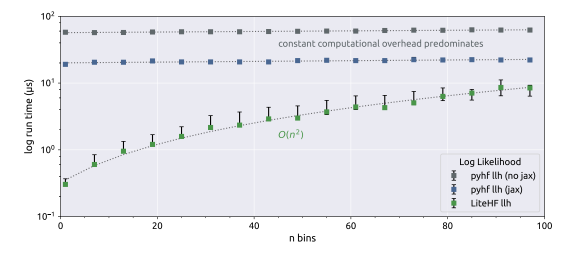


Figure 5.6: Benchmarking the log likelihood of the *n*-bin model up to 100 bins. The pyhf log likelihood is evaluated with and without just-in-time compilation by jax and compared to the LiteHF log likelihood. The run-time is averaged over 10<sup>4</sup> evaluations.

The Float64 likelihood (gray) has a constant run-time of about  $60 \, \mu s$ . Just-in-time compilation (blue) reduces the run-time by a factor of 3. However, the LiteHF log likelihood (green) is still computed more efficiently than the jax pyhf llh and shows an  $\mathcal{O}(n^2)$  dependency.

The log likelihood can therefore also be excluded as the cause of the run-time increase for n > 30. Moreover, batty and Julia use the *same* wrappers and conversions to run the pyhf likelihood in Julia which makes the Python behavior even more inexplicable. Nevertheless this rapid run-time increase is one of the major drawbacks in the current batty implementation which requires further investigation.

### Chapter 6

### Conclusion

This work dealt with the implementation of a Bayesian inference pipeline in Python. During this thesis the Python packages priorhf and pyhf\_llh have been developed and the "BAT to Python interface" batty [14] continuously enhanced.

The first chapters of this work cover theoretical foundations of Bayesian inference, MCMC methods and statistical models in HistFactory (HF). Afterwards, chapter 4 derived a formalism to generate conjugate priors from HF models, which is implemented in the package priorhf. HistFactory likelihoods are computed by pyhf [8] while the overall run-time can be reduced by a factor of 2 when employing the jax [17] backend of pyhf. Here the Python package pyhf\_llh ensures a standardized and yet simple interface to pyhf likelihoods and their gradients which are computed by automatic differentiation via jax. All required conversions to execute BAT functions from Python are implemented in batty and described in section 4.3. The developed Bayesian inference pipeline is verified for three HF models py a posterior predictive check.

Chapter 5 of this work evaluates several run-time benchmarks where the Python-Julia pipeline batty [14] was benchmarked against (i) a pure-Julia Bayesian inference implementation (with LiteHF) and (ii) a Julia implementation that calls the pyhf likelihood. Clearly, the pure Julia implementation outperform batty and (ii) since pure-Julia code is compiled to high-performant machine code while Python code is executed by a high-level interpreter language.

The first benchmark investigates the run-time performance for low-dimensional HF model. Here Metropolis-Hastings (MH) in batty is about 10 times slower than the pure-Julia version. The run-time of Hamiltonian Monte Carlo (HMC) in batty even takes 100 times longer than in a pure Julia implementation.

The second benchmark was carried out for a n-bin model with a (n+1)-dimensional parameter space. For n < 30 bins, the run-time performance of batty is similar to the first benchmark. However for n > 30 bins the run-time of batty significantly increases, while it remains constant for the other two implementations. The reason for this behaviour in Python could not be resolved within this thesis task and requires further investigation.

### Appendix A

# HistFactory Models

### A.1 2\_bin\_uncorr Model

```
• parameter \boldsymbol{\theta} = [\mu, \gamma_1, \gamma_2]
```

• signal model [5.0, 10.0] with normfactor modifier

• bkg model [50.0, 60.0], shapesys modifier with relative uncert. [10%, 20%]

```
"channels": [
  { "name": "singlechannel",
    "samples": [
      { "name": "signal",
        "data": [5.0, 10.0],
        "modifiers": [ { "name": "mu", "type": "normfactor", "data": null} ]
      { "name": "background",
        "data": [50.0, 60.0],
        "modifiers": [
          { "name": "uncorr_bkguncrt",
            "type": "shapesys",
            "data": [5.0, 12.0] }
        ]
    ]
  }
],
"observations": [
  { "name": "singlechannel", "data": [50.0, 60.0] }
],
"measurements": [
  { "name": "Measurement", "config": {"poi": "mu", "parameters": []} }
],
"version": "1.0.0"
```

### A.2 2\_bin\_corr Model

```
• parameter \boldsymbol{\theta} = [\mu, \theta]
   • signal model [12.0, 11.0] with normfactor modifier
   • correlated background [50, 52.0] with histosys modifier
   • observations [53.0, 65.0]
{
    "channels": [
      { "name": "singlechannel",
        "samples": [
          { "name": "signal",
            "data": [12.0,11.0],
            "modifiers": [ { "name": "mu", "type": "normfactor", "data": null } ]
          { "name": "background",
            "data": [ 50.0, 52.0 ],
            "modifiers": [
              { "name": "correlated_bkg_uncertainty",
                "type": "histosys",
                "data": { "hi_data": [45.0, 57.0], "lo_data": [55.0, 47.0] }
            ]
          }
        ]
      }
    "observations": [
      { "name": "singlechannel", "data": [53.0, 65.0] }
   ],
    "measurements": [
        "name": "Measurement",
        "config": {
        "poi": "mu",
        "parameters": []
      }
   ],
    "version": "1.0.0"
```

### A.3 4\_bin Model

- parameter  $\boldsymbol{\theta} = [\mu, \theta, \theta_{SF}]$
- signal model [2, 3, 4, 5] with normfactor modifier
- background model [30, 19, 9, 4] with histosys and normsys modifier
- observations [34, 22, 13, 11]

```
{
  "channels": [
    { "name": "singlechannel",
      "samples": [
        { "name": "signal MC",
          "data": [2, 3, 4, 5],
          "modifiers": [ { "name": "mu", "type": "normfactor", "data": null } ]
        },
          "name": "bkg MC",
          "data": [30, 19, 9, 4],
          "modifiers": [
            { "name": "theta",
              "type": "histosys",
              "data": { "hi_data": [31, 21, 12, 7], "lo_data": [29, 17, 6, 1] }
            { "name": "SF_theta",
              "type": "normsys",
              "data": {"hi": 1.1,"lo": 0.9}
            }
          ]
        }
      ]
    }
  ],
  "observations": [
    { "name": "singlechannel", "data": [34, 22, 13, 11]}
  "measurements": [
    { "name": "Measurement", "config": {"poi": "mu", "parameters": []} }
 ],
  "version": "1.0.0"
}
```

## Appendix B

### Run-time Statistics

While executing code on a PC, the running time of a function varies due to other active processes. A typical run-time statistics for  $10^4$  samples is illustrated in Figure B.1.

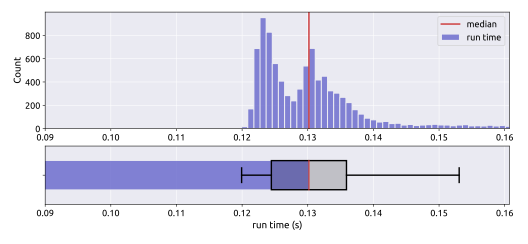


Figure B.1: Run-time statistics for evaluating the log likelihood 10k times.

In order simplify the plot while keeping track of the process statistics, run-times are visualized as bar-plot with an box-plot on top. The final bar value is the median run-time, which corresponds to the red line in the box-plot. The box contains  $50\,\%$  of the data that is closed to the median and the whiskers mark the minimum and maximum value.

### Benchmarking the log likelihood function with numpy and jax backend

A just-in-time compiled likelihood function (by jax) reduces the run-time by a factor of 1.5 to 2.5, compared to the numpy backend.

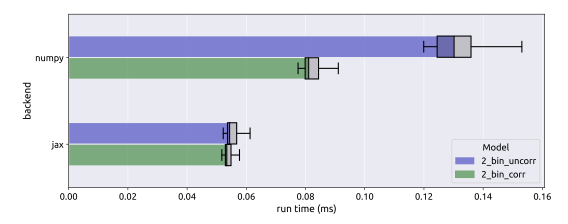


Figure B.2: Benchmarking the log likelihood with numpy and jax backend for two HF models.

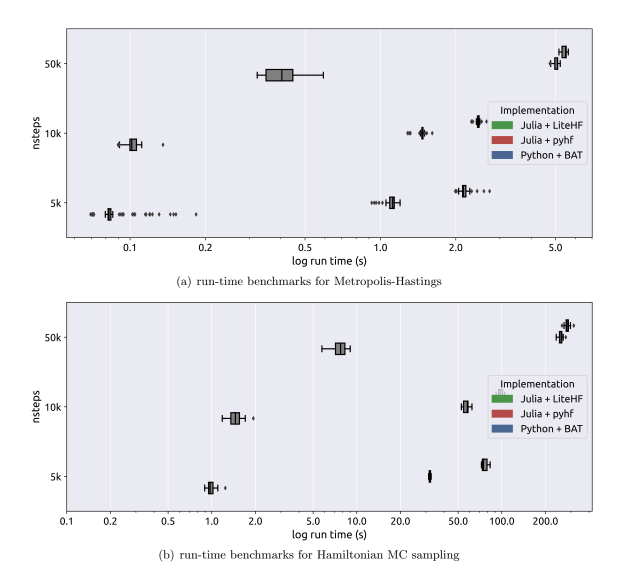
# Appendix C

## Additional Run-time Benchmarks

The run-time benchmarks for the 4-bin model (section A.3) are visualized in Figure C.1 and the numerical values summarized in Table C.1. The run-time performance coincidence with the observations for the 2\_bin\_corr model in Figure 5.2, while HMC need almost twice as long for the 4\_bin model. This can be explained by the increased model complexity. The run-time performance for MH is similar for both models.

**Table C.1:** Summary of the run-time benchmarks in Figure C.1. The values represent the median run-time with  $1\sigma$  standard deviation without outliers for 5k, 10k and 50k nsteps.

		5k	10k	50k
Method	Implementation	$\mathbf{run\text{-}time}\ (\mathrm{s})$		
	$\mathrm{Julia} + \mathtt{LiteHF}$	$0.08 \pm 0.01$	$0.10 \pm 0.00$	$0.40 \pm 0.05$
$\mathbf{MH}$	Julia + pyhf	$1.11 \pm 0.06$	$1.47 \pm 0.05$	$5.09 \pm 0.13$
	Python + BAT	$2.14 \pm 0.12$	$2.46 \pm 0.05$	$5.49 \pm 0.13$
	m Julia + LiteHF	$0.97 \pm 0.06$	$1.45 \pm 0.14$	$7.73 \pm 0.79$
$\mathbf{HMC}$	Julia + pyhf	$31.9 \pm 0.9$	$57.8 \pm 3.0$	$257 \pm 11$
	Python + BAT	$74.7 \pm 4.0$	$97.7 \pm 4.2$	$281 \pm 14$



**Figure C.1:** Run-time benchmark for the 4\_bin model for different MCMC nsteps with logarithmic time axis. Uncertainties are visualized as box-plot as illustrated in Figure B.1. The run-time in Julia is averaged over 100 runs, for the other two implementations over 40 runs for MH and 10 runs for HMC.

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