

Pigeons.jl: Distributed sampling from intractable distributions

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ABSTRACT

We introduce a software package, Pigeons.jl, that provides a way to leverage distributed computation to obtain samples from complicated probability distributions, such as multimodal posteriors arising in Bayesian inference and high-dimensional distributions in statistical mechanics. Pigeons.jl provides simple APIs to perform such computations single-threaded, multi-threaded, and/or distributed over thousands of MPI-communicating machines. In addition, Pigeons.jl guarantees a property that we call strong parallelism invariance: the output for a given seed is identical irrespective of the number of threads and processes, which is crucial for scientific reproducibility and software validation. We describe the key features of Pigeons.jl and the approach taken to implement a distributed and randomized algorithm that satisfies strong parallelism invariance.

Keywords

distributed computation, Bayesian inference, parallelism invariance, multi-threading, Message Passing Interface, Markov chain Monte Carlo

1. Introduction

In many scientific application domains, the ability to obtain samples from a distribution is crucial. For instance, sampling methods have been used to discover magnetic polarization in the black hole of galaxy M87 [2] and to image the Sagittarius A* black hole [1]. They have also been used to model the evolution of single-cell cancer genomes [22], infer plasma dynamics inside nuclear fusion reactors [13], and to identify gerrymandering in Georgia's 2021 congressional districting plan [31]. Similarly, evaluating high-dimensional integrals or sums over complicated combinatorial spaces are related tasks that can also be solved with sampling via Markov chain Monte Carlo (MCMC) methods. However, such calculations can often be bottlenecks in the scientific process, with simulations that can last days or even weeks to finish.

Pigeons.jl¹ enables users to sample efficiently from highdimensional and complex distributions and solve integration problems by implementing state-of-the-art sampling algorithms [27, 26] that leverage distributed computation. Its simple API allows users to perform such computations single-threaded, multi-threaded, and/or distributed over thousands of MPI-communicating machines. Further, it comes with guarantees on strong parallelism invariance wherein the output for a given seed is *identical* irrespective of the number of threads or processes. Such a level of reproducibility is rare in distributed software but of great use for the purposes of debugging in the context of sampling algorithms, which produce stochastic output. Specifically, Pigeons.jl is designed to be suitable and yield reproducible output for:

- (1) one machine running on one thread;
- (2) one machine running on several threads;
- (3) several machines running, each using one thread, and
- (4) several machines running, each using several threads.

1.1 Problem formulation

We describe the class of problems that can be approached using Pigeons.jl. Let $\pi(x)$ denote a probability density 2 called the *target*. In many problems, e.g. in Bayesian statistics, the density π is typically known only up to a normalizing constant,

$$\pi(x) = \frac{\gamma(x)}{Z}, \qquad Z = \int \gamma(x) \, \mathrm{d}x,$$
 (1)

where γ can be evaluated pointwise but Z is typically unknown. Pigeons.jl takes as input the function γ .

The output of Pigeons.jl can be used for two main tasks:

- (1) Approximating integrals of the form $\int f(x)\pi(x) \, \mathrm{d}x$. For example, the choice f(x)=x computes the mean and $f(x)=I[x\in A]$ computes the probability of A under π , where $I[\cdot]$ denotes the indicator function.
- (2) Approximating the value of the normalization constant Z. For example, in Bayesian statistics, this corresponds to the marginal likelihood, which can be used for model selection.

Its implementation shines compared to traditional sampling packages in the following scenarios:

—When the target density π is challenging due to a complex structure (e.g., high-dimensional, multi-modal, etc.).

¹The source code for Pigeons.jl can be found at https://github.com/ Julia-Tempering/Pigeons.jl. Pigeons.jl v0.2.0, used in the example scripts below, is currently compatible with Julia 1.8+.

²This density may also be a probability mass function (discrete variables). We also allow for a combination of discrete/continuous variables.

- —When the user needs not only $\int f(x)\pi(x) dx$ but also the normalization constant Z. Many existing tools focus on the former and struggle or fail to do the latter.
- —When the target distribution π is defined over a non-standard space, e.g. a combinatorial object such as a phylogenetic tree.

1.2 What is of interest to the general Julia developer?

Ensuring code correctness at the intersection of randomized, parallel, and distributed algorithms is a challenge. To address this challenge, we designed Pigeons.jl based on a principle that we refer to as strong parallelism invariance (SPI). Namely, the output of Pigeons.jl is completely invariant to the number of machines and/or threads. Without explicitly keeping SPI in mind during software construction, the (random) output of the algorithm is only guaranteed to have the same distribution. This is a much weaker guarantee that, in particular, makes debugging difficult. However, with our notion of SPI we make debugging and software validation considerably easier. This is because the developer can first focus on the fully serial randomized algorithm, and then use it as an easyto-compare gold-standard reference for parallel/distributed implementations. This strategy is used extensively in Pigeons. il to ensure correctness. In contrast, testing equality in distribution, while possible (e.g., see [12]), incurs additional false negatives due to statistical error.

The general Julia developer will be interested in:

- —The main causes of a violation of strong parallelism invariance that we have identified (Section 4)—some of which are specific to Julia—and how we address them in Pigeons.jl.
- —The SplittableRandoms.jl³ package that was developed by our team to achieve strong parallelism invariance in Pigeons.jl (Section 4.2).

2. Examples

In this section we present a set of minimal examples that demonstrate how to use Pigeons.jl for sampling. For further reading, we also direct readers to our growing list of examples, which can be found at https://github.com/Julia-Tempering/Pigeons.jl/tree/main/examples. We begin by installing the latest official release of Pigeons.jl:

2.1 Targets

To use Pigeons.jl, we must specify a target distribution, given by γ in Eq. (1). Numerous possible types of target distributions are supported, including custom probability densities (specified up to a normalizing constant) written in Julia. We also allow to interface with models written in common probabilistic programming languages, including:

- —Turing.jl [11] models (TuringLogPotential)
- -Stan [9] models (StanLogPotential)
- —Comrade.jl⁴ models for black hole imaging (ComradeLogPotential)

—Non-Julian models with foreign-language Markov chain Monte Carlo (MCMC) code (e.g. Blang [4] code for phylogenetic inference over combinatorial spaces)

Additional targets are currently being accommodated and will be introduced to Pigeons.jl in the near future.

In what follows, we demonstrate how to use Pigeons with a Julia Turing model applied to a non-identifiable "coinflip" data set, modified from the example at https://turing.ml/v0.22/docs/using-turing/quick-start. The Bayesian model can be formulated as

$$p_1, p_2 \stackrel{iid}{\sim} U(0, 1)$$

$$Y \mid p_1, p_2 \sim \text{Binomial}(n, p_1 p_2).$$

$$(2)$$

The random variable Y is the number of heads observed on n coin flips where the probability of heads is p_1p_2 . This model is non-identifiable, meaning that it is not possible to distinguish the effects of the two different parameters p_1 and p_2 . As a consequence, the target distribution exhibits a complicated structure, as displayed in Fig. 1. The density of interest corresponding to this model is

$$\pi(p_1, p_2) = \gamma(p_1, p_2)/Z,$$

where

$$\gamma(p_1, p_2) = \binom{n}{y} (p_1 p_2)^y (1 - p_1 p_2)^{n-y} I[p_1, p_2 \in [0, 1]]$$
(3)
$$Z = \int_0^1 \int_0^1 \gamma(p_1, p_2) dp_1 dp_2.$$
(4)

The distribution π is also known as the *posterior distribution* in Bayesian statistics.

Suppose that we perform n=100,000 coin tosses and observe y=50,000 heads. We would like to obtain samples from our posterior, π , having collected this data. We begin by installing Turing

```
Pkg.add("Turing")
```

and then defining our Turing model and storing it in the variable model:

```
using Turing
@model function coinflip(n, y)
   p1 ~ Uniform(0.0, 1.0)
   p2 ~ Uniform(0.0, 1.0)
   y ~ Binomial(n, p1 * p2)
   return y
end
model = coinflip(100000, 50000)
```

From here, it is straightforward to sample from the density given by Eq. (3) up to a normalizing constant. We use non-reversible parallel tempering [27] (PT), Pigeons.jl's state-of-the-art sampling algorithm, to sample from the target distribution. PT comes with several tuning parameters and [27] describe how to select these parameters effectively, which Pigeons.jl implements under the hood. We also specify that we would like to store the obtained samples in memory to be able to produce trace-plots, as well as some basic online summary statistics of the target distribution and useful diagnostic output by specifying record = [traces, online, round_trip, Pigeons.timing_extrema, Pigeons.allocation_extrema]. It is also possible to leave the

 $^{^3 {\}tt https://github.com/Julia-Tempering/SplittableRandoms.jl}$

⁴https://github.com/ptiede/Comrade.jl

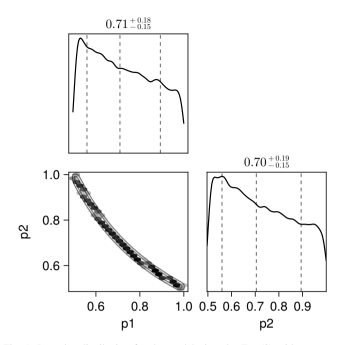


Fig. 1: Posterior distribution for the model given by Eq. (2) with n=100,000 coin flips and y=50,000 observed heads, estimated using 2^{17} samples from Pigeons.jl. We present the pairwise plot for p_1 and p_2 , as well as the estimated densities of the marginal of the posterior for each of the two parameters. Note that because the model is non-identifiable, as we collect more data the posterior distribution concentrates around the curve $p_1p_2=0.5$, instead of a single point, assuming that the true probability of observing heads is 0.5.

record argument empty and reasonable defaults will be selected for the user. The code below runs Pigeons.jl on one machine with one thread. We use the default values for most settings, however we explain later how one can obtain improved performance by setting arguments more carefully (see Section 2.4).

```
using Pigeons
pt = pigeons(
   target = TuringLogPotential(model),
   record = [
        traces, online, round_trip,
        Pigeons.timing_extrema,
        Pigeons.allocation_extrema])
```

Note that to convert the Turing model into an appropriate Pigeons.jl target for sampling, we pass the model as an argument to TuringLogPotential(). Once we have stored the PT output in the variable pt we can access the results, as described in the following section. The standard output after running the above code chunk is displayed in Fig. 2 and explained in the next section. For purposes of comparison, we also run a traditional (single-chain Markov chain Monte Carlo) method.

2.1.1 Other targets. As mentioned previously, it is also possible to specify targets with custom probability densities, as well as Stan and Turing models. Additionally, suppose we have some code implementing vanilla MCMC, written in an arbitrary "foreign" language such as C++, Python, R, Java, etc. Surprisingly, it is very

simple to bridge such code with Pigeons.jl. The only requirement on the foreign language is that it supports reading the standard input and writing to the standard output, as described in our online documentation.

2.2 Outputs

Pigeons.jl provides many useful types of output, such as: plots of samples from the distribution, estimates of normalization constants, summary statistics of the target distribution, and various other diagnostics. We describe several examples of possible output below.

2.2.1 Standard output. An example of the standard output provided by Pigeons.jl is displayed in Fig. 2. Each row of the table in the output indicates a new tuning round in parallel tempering, with the #scans column indicating the number of scans/samples in that tuning round. During these tuning rounds, Pigeons.jl searches for optimal values of certain PT tuning parameters. Other outputs include:

- —restarts: a higher number is better. Informally, PT performs well at sampling from high-dimensional and/or multi-modal distributions by pushing samples from an easy-to-sample distribution (the reference) to the more difficult target distribution. A tempered restart happens when a sample from the reference successfully percolates to the target. (See the subsequent sections for a more detailed description of parallel tempering.) When the reference supports i.i.d. sampling, tempered restarts can enable large jumps in the state space.
- —Λ: the global communication barrier, as described in [27], which measures the inherent difficulty of the sampling problem. A rule of thumb to configure the number of PT chains is also given by [27], where they suggest that stable performance should be achieved when the number of chains is set to roughly 2Λ. See Section 2.2.6 for more information.
- —time and allc: the time (in seconds) and number of allocations (in bytes) used in each round.
- $-\log(Z_1/Z_0)$: an estimate of the logarithm of the ratio of normalization constants between the target and the reference. In many cases, $Z_0=1$.
- — $\min(\alpha)$ and $\max(\alpha)$: minimum and average swap acceptance rates during the communication phase across the PT chains. See Section 3 for a description of PT communication.

2.2.2 *Plots.* It is straightforward to obtain plots of samples from the target distribution, such as trace-plots, pairwise plots, and density plots of the marginals.

To obtain posterior densities and trace-plots, we first make sure that we have the third-party MCMCChains.jl⁵, StatsPlots.jl⁶, and PlotlyJS.jl⁷ packages installed via

```
Pkg.add("MCMCChains", "StatsPlots", "PlotlyJS")
```

With the pt output object from before for our non-identifiable coinflip model, we can run the following:

```
using MCMCChains, StatsPlots, PlotlyJS
```

 $^{^5 {\}tt https://github.com/TuringLang/MCMCChains.jl}$

 $^{^6 {\}tt https://github.com/JuliaPlots/StatsPlots.jl}$

⁷https://github.com/JuliaPlots/PlotlyJS.jl

#scans	restarts	Λ	time(s)	allc(B)	$log(Z_1/Z_0)$	$\min(\alpha)$	$mean(\alpha)$
2	0	1.04	0.383	3.48 e+0 7	-4.24 e+0 3	0	0.885
4	0	4.06	0.00287	1.79 e+0 6	-16.3	4.63 e-06	0.549
8	0	3.49	0.00622	3.55 e+06	-12.1	0.215	0.612
16	0	2.68	0.0161	7 .46 e+0 6	-10.2	0.518	0.703
32	0	4.29	0.0353	1.37 e+07	-11.8	0.222	0.524
64	3	3.17	0.0699	2.86 e+07	-11.5	0.529	0.648
128	8	3.56	0.139	5.53 e+07	-11.5	0.523	0.605
256	12	3.38	0.241	1.1 e+08	-11.6	0.526	0.625
512	37	3.48	0.473	2.22 e+08	-12	0.527	0.614
1.02e+03	77	3.55	0.895	4.46 e+08	-11.8	0.571	0.605

Fig. 2: Standard output provided by Pigeons.jl. Rows indicate tuning rounds of the PT algorithm with an exponentially increasing number of PT iterations (#scans). Columns indicate various useful diagnostics, such as the number of allocations per round, time (in seconds), and estimates of the log of the normalization constants. The output is described in greater detail in Section 2.2.6 and has been modified to exclude columns that are not described in the paper.

```
plotlyjs()
samples = Chains(
    sample_array(pt), variable_names(pt))
my_plot = StatsPlots.plot(samples)
display(my_plot)
```

The output of the above code chunk is an interactive plot that can be zoomed in or out and exported as an HTML webpage. A modified static version of the output for the first parameter, p_1 , is displayed in the top panel of Fig. 3, along with a comparison to the output from a single-chain algorithm in the bottom panel of the same figure.

To obtain pair plots, we add the PairPlots.jl⁸ and CairoMakie.jl⁹ packages:

```
Pkg.add("PairPlots", "CairoMakie")
```

and then run

```
using PairPlots, CairoMakie
my_plot = PairPlots.pairplot(samples)
display(my_plot)
```

The output of the above code chunk with an increased number of samples is displayed in Fig. 1. 10

2.2.3 Estimate of normalization constant. The (typically unknown) constant Z in Eq. (1) is referred to as the normalization constant. In many applications, it is useful to approximate this constant. For example, in Bayesian statistics, this corresponds to the marginal likelihood and can be used for model selection.

As a side-product of PT, we automatically obtain an approximation to the natural logarithm of the normalization constant. This is done automatically using the stepping stone estimator [30]. The estimate can be accessed using

```
stepping_stone(pt)
```

In the case of the normalization constant given by Eq. (4) for n=100,000 and y=50,000, we can exactly obtain its value as $\log(Z)\approx -11.8794$ using a computer algebra system. Note that this is very close to the output provided in Fig. 2.

2.2.4 Online statistics. Having specified the use of the online recorder in our call to pigeons(), we can output some basic summary statistics of the marginals of our target distribution. For instance, it is straightforward to estimate the mean and variance of each of the marginals of the target with

```
using Statistics
mean(pt); var(pt)
```

Other constant-memory statistic accumulators are made available in the OnlineStats.jl [10] package. To add additional constant-memory statistic accumulators, we can register them via Pigeons.register_online_type(), as described in our online documentation. For instance, we can also compute constant-memory estimates of extrema of our distribution.

2.2.5 Off-memory processing. When either the dimensionality of the model or the number of samples is large, the obtained samples may not fit in memory. In some cases it may be necessary to store samples to disk if our statistics of interest cannot be calculated online and with constant-memory (see Section 2.2.4). We show here how to save samples to disk when Pigeons.jl is run on a single machine. A similar interface can be used over MPI.

First, we make sure that we set checkpoint = true, which saves a snapshot at the end of each round in the directory results/all/<unique directory> and is symlinked to results/latest. Second, we make sure that we use the disk recorder by setting record = [disk], along with possibly any other desired recorders. Accessing the samples from disk can then be achieved in a simple way using the Pigeons.jl function process_sample().

2.2.6 PT diagnostics. We describe how to produce some key parallel tempering diagnostics from [27].

⁸https://github.com/sefffal/PairPlots.jl

⁹https://github.com/JuliaPlots/CairoMakie.jl

¹⁰The code chunk above only works with Julia 1.9.

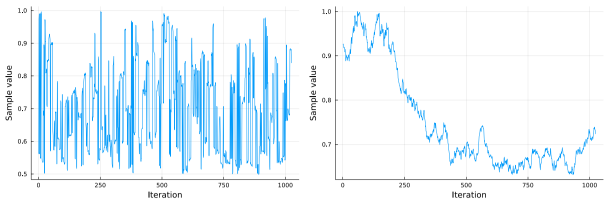


Fig. 3: Trace-plots for the first parameter, p_1 , in the non-identifiable coinflip Turing model. **Left:** Samples from Pigeons.jl using PT with 10 chains. Note that the trace-plot indicates fast mixing/exploration across the state space. **Right:** Single-chain Markov chain Monte Carlo. Note that the trace-plot explores the state space much more slowly when we do not use PT.

The global communication barrier, denoted Λ in Pigeons.jl output, can be used to approximately inform the appropriate number of chains. Based on [27], stable PT performance should be achieved when the number of chains is set to roughly 2Λ . This can be achieved by modifying the n_chains argument in the call to pigeons(). The global communication barrier is shown at each round and can also be accessed with

```
Pigeons.global_barrier(pt)
```

The number of restarts per round can be accessed with

```
n_tempered_restarts(pt)
```

These quantities are also displayed in Fig. 2. Many other useful PT diagnostic statistics and plots can be obtained, as described in our full documentation.

2.3 Parallel and distributed PT

One of the main benefits of Pigeons.jl is that it allows users to easily parallelize and/or distribute their PT sampling efforts. We explain how to run MPI locally on one machine and also how to use MPI when a cluster is available.

2.3.1 Running MPI locally. To run MPI locally on one machine using four MPI processes and one thread per process, use

- 2.3.2 Running MPI on a cluster. Often, MPI is available via a cluster scheduling system. To run MPI over several machines:
- (1) In the cluster login node, follow the Pigeons.jl installation instructions in our online documentation.
- (2) Start Julia in the login node, and perform a one-time setup by calling Pigeons.setup_mpi().

(3) In the Julia REPL running in the login node, run:

```
pigeons(
   target = TuringLogPotential(model),
   n_chains = 1_000,
   on = MPI(n_mpi_processes = 1_000,
        n_threads = 1))
```

The code above will start a distributed PT algorithm with 1,000 chains on 1,000 MPI processes each using one thread. Note that for the above code chunks involving ChildProcess() and MPI() to work, it may be necessary to specify dependencies in their function calls

2.4 Additional options

In the preceding example we only specified the target distribution and let Pigeons.jl decide on default values for most other settings of the inference engine. There are various settings we can change, including: the random seed (seed), the number of PT chains (n_chains), the number of PT tuning rounds/samples (n_rounds), and a variational reference distribution family (variational), among other settings. For instance, we can run

```
pigeons(
    target = TuringLogPotential(model),
    n_rounds = 10,
    n_chains = 10,
    variational = GaussianReference(),
    seed = 2
)
```

which runs PT with the same Turing model target as before and explicitly states that we should use 10 PT tuning rounds with 10 chains (described below). In the above code chunk we also specify that we would like to use a Gaussian variational reference distribution. That is, the reference distribution is chosen from a multivariate Gaussian family that lies as close as possible to the target distribution in order to improve the efficiency of PT. We refer readers to [26] for more details. When only continuous parameters are of interest, we encourage users to consider using variational = GaussianReference() and setting n_chains_variational =

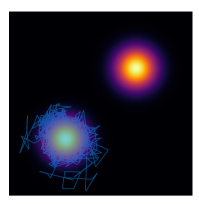


Fig. 4: A simple bimodal distribution from which traditional MCMC methods may struggle to obtain samples. Blue lines display output from 1,000 iterations of a Metropolis-Hastings random walk MCMC algorithm. The sampler in this figure is visibly trapped in one of the two modes.

10, for example, as the number of restarts may substantially increase with these settings.

3. Parallel tempering

Pigeons.jl provides an implementation of distributed parallel tempering (PT) described in [27], which we outline in Algorithm 1. This section gives both a brief overview of PT and some details of its distributed implementation.

In this section we assume a basic understanding of Markov chain Monte Carlo (MCMC) methods. For readers unfamiliar with MCMC, it is important to know that it is a method to obtain approximate samples from a distribution. More specifically, it is a class of algorithms for simulating a Markov chain of states that look like draws from the target distribution π . However, with traditional MCMC methods, the samples may be heavily correlated instead of independent and may fail to sufficiently explore the full space of the distribution (see Fig. 4); PT is a method that aims to address these issues. For a more in-depth review of PT, we refer readers to [26].

3.1 Overview of PT

Suppose that we would like to estimate integrals involving π , such as $\int f(x)\pi(x)\,\mathrm{d}x$. These integrals may be multivariate and even include combinations of continuous and discrete variables (where sums replace integrals in the discrete case). One method is to obtain samples from π to approximate such integrals. Often, the distribution π can be challenging for traditional MCMC methods—such as Metropolis-Hastings, slice sampling, and Hamiltonian Monte Carlo—because of its structure. For example, in a bimodal example such as the one illustrated in Figure 4, traditional methods might remain in one of the two modes for an extremely long period of time.

To resolve this issue, PT constructs a sequence of N distributions, $\pi_1, \pi_2, \ldots, \pi_N$, where π_N is usually equal to π . The distributions are chosen so that it is easy to obtain samples from π_1 with the sampling difficulty increasing as one approaches π_N . An example of such a sequence of distributions, referred to as an *annealing path*, is shown in Figure 5.

We now turn to explain how this path of distributions can be used to enhance sampling from the target distribution, π . PT operates by

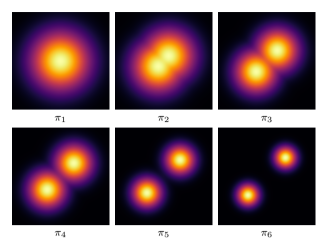


Fig. 5: Heatmaps of six distributions lying on an annealing path from a unimodal reference distribution π_1 , from which it is straightforward to obtain samples, and ending at π_N , which is in this case the bimodal distribution from Figure 4. (Note that in this case the colours between the heatmaps are not directly comparable because the densities of intermediate distributions are not normalized.)

first obtaining samples from each distribution on the path in parallel (referred to as an *exploration phase*). Then, samples between adjacent distributions are swapped (referred to as a *communication phase*). The communication phase in PT is crucial: it allows for the discovery of new regions of the space of the target distribution such as the top-right mode of the distribution presented in Figure 4.

3.2 Local exploration and communication

A struct that is useful for the implementation of PT is the Replica, which we will often refer to in the remainder of this section. A single Replica struct stores a state variable and a chain integer, among other entries. At the beginning of PT, N Replicas are created, one for each distribution on the annealing path, and the chain entries in the N replicas are initialized at $1, 2, \ldots, N$, respectively. For a given Replica, if the chain number is i and the state is x, then this means that the sample corresponding to the i-th distribution in the sequence π_1, \ldots, π_N is currently at location x.

In the local exploration phase, each Replica's state is modified using an MCMC move targeting π_i , where i is given by Replica.chain. The MCMC move can either modify Replica.state in-place, or modify the Replica's state field. This operation is indicated by the local_exploration function in Algorithm 1.

In the communication phase, PT proposes swaps between pairs of replicas. In principle, there are two equivalent ways to do a swap. In the first implementation, the Replicas could exchange their state fields. Alternatively, they could exchange their chain fields. Because we provide distributed implementations, we use the latter as it ensures that the amount of data that needs to be exchanged between two machines during a swap can be made very small (two floating point numbers), resuling in an exchange of O(N) messages of size O(1). Note that this cost does not vary with the dimensionality of the state space, in contrast to the first implementation that would transmit O(N) messages of size O(d), where d is the dimension of the state space, incurring a potentially very high communication cost for large values of d.

3.3 Distributed implementation

A distributed implementation of PT is presented in Algorithm 1 and we describe the details of the implementation below. We present the algorithm from the perspective that the number of machines available is equal to the number of Replicas and distributions, N. However, Pigeons.jl also allows for the more general case where the number of machines is not necessarily equal to N.

3.3.1 The PermutedDistributedArray. Recall that our theoretical O(1) message size for communication between two machines is achieved by exchanging chain indices between Replicas instead of their states. One difficulty can be encountered in a distributed implementation, which we illustrate with the following simple example. Suppose we have N=4 distributions, chains, replicas, and machines, and that machine 1 is exploring chain 2 while machine 4 is exploring chain 3. At one point, the Replica at chain 2 (machine 1) might need to exchange chain indices with the Replica that has chain index 3 (machine 4). However, a priori it is not clear how machine 1 should know that it should communicate with machine 4 because it has no knowledge about the chain indices on the other machines.

To resolve this issue, we introduce a special data structure called a PermutedDistributedArray. In the case where the number of replicas is equal to the number of available machines, the construction is quite simple and effectively results in each machine storing one additional integer. Considering the same example above, the solution to the problem is to have machine 1 (chain 2 wanting to communicate with chain 3) communicate with machine 3. Machine 3 stores in its dist_array variable of type PermutedDistributedArray the value 4, which is the machine number that stores chain 3. By updating these PermutedDistributedArray variables at each communication step, we can ensure that each machine j is aware of which machine number currently stores chain j. Therefore, the machines act as keys in a dictionary for the communication permutations. An illustration of communication between four machines is provided in Fig. 6. We note that with a PermutedDistributedArray we make special assumptions on how we access and write to the array elements. Several MPI processes cooperate, with each machine storing data for a slice of this distributed array, and at each time step an index of the array is manipulated by exactly one machine.

3.3.2 MPI implementation details. We use the MPI.jl [7] package to support communication between machines. In our pseudocode in Algorithm 1 we define the MPI-style functions send(), receive!(), and waitall(). The function send() has three arguments in our pseudocode: the first argument is the object to be sent, the second is the machine number to which the object should be sent, and finally the third argument is a unique tag for this specific send/receive request at this time step and on this machine, generated by tag(). The tag() function can be any function that allows one to uniquely identify a tag for a given send/receive request on a given machine at any given time step t. The function receive!() has the same last two arguments as send() except that the first argument specifies the object to which data should be written directly. Finally, waitall(requests) waits for all initialized MPI requests for a given pair of machines to complete.

In Algorithm 1 we also introduce the functions permuted_get() and permuted_set!(). Given the current machine's dist_array and the partner chain with which it should communicate, permuted_get() returns the machine number that holds the partner chain. Given the current machine's dist_array, chain

Algorithm 1 Distributed PT on machine j (one replica per machine)

```
Require: Initial state x_0, sequence of distributions \{\pi_i\}_{i=1}^N, num-
    ber of iterations T, machine number j
 1: \mathtt{chain} \leftarrow j
                                              2: dist_array[j] \leftarrow j
                                            \triangleright chain j is on machine j
 3: for t in 1, 2, ..., T do
         if t is even then
                                  4:
 5:
             P \leftarrow \{i : 1 \le i < N, i \text{ is even}\}
 6:
             P \leftarrow \{i : 1 \le i < N, i \text{ is odd}\}
 7:
 ۸٠
         end if
 9:
         x_t \leftarrow \texttt{local\_exploration}(\pi_{\texttt{chain}}, x_{t-1})
10:
         partner \leftarrow nothing
         if chain \in P then
11:
12:
             \mathtt{partner} \leftarrow \mathtt{chain} + 1
13:
         else if chain - 1 \in P then
14:
            partner \leftarrow chain - 1
         end if
15:
16:
         if partner != nothing and swap is accepted then
             \texttt{to\_machine} \gets \texttt{permuted\_get}(\texttt{dist\_array}, \texttt{partner})
17:
    ▷ retrieve dist_array[partner]
             send_tag \leftarrow tag(t, chain, j)
18:
             send(chain, to_machine, send_tag)
19:
20:
             receive\_tag \leftarrow tag(t, chain, to\_machine)
             receive!(chain, to_machine, receive_tag)
21:
22:
             waitall(requests) ▷ wait for MPI requests for this
    pair of machines to complete
23:
            permuted_set!(dist_array, chain, j) > set value
    of dist_array[chain] to to_machine
24:
         end if
25:
         c_t \leftarrow \mathtt{chain}
26: end for
27: return \{(x_t, c_t)\}_{t=1}^T
```

number, and machine number j, permuted_set!() updates the PermutedDistributedArray variables with the updated permutation.

4. Strong parallelism invariance

In this section we describe potential violations of strong parallelism invariance (SPI) that can occur in a distributed setting. We also explain how we avoid these issues by using special distributed reduction schemes and splittable random number generators. Insights provided in this section can be applied to general distributed software beyond Julia.

We have identified two factors that can cause violations of our previously-defined SPI that standard Julia libraries do not automatically take care of:

- (1) Non-associativity of floating point operations: When several workers perform distributed reduction of floating point values, the output of this reduction will be slightly different depending on the order taken during reduction. When these reductions are then fed into further random operations, this implies that two randomized algorithms with the same seed but using a different number of workers will eventually arbitrarily diverge.
- (2) Global, thread-local, and task-local random number generators: These are the dominant approaches to parallel random number generators in current languages, and an appropriate understanding of these RNGs is necessary. In particular, in Julia

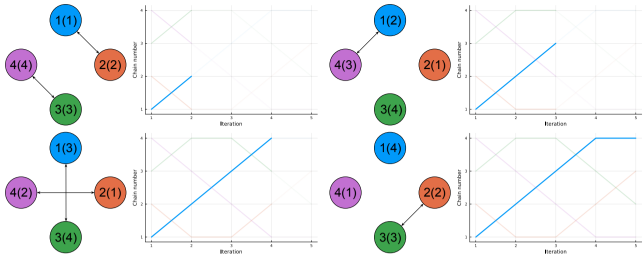


Fig. 6: A summary of communication between four machines with N=4 PT chains. Circles represent machines with the machine number in the center of the circle. Different colours are also used for different machines. Numbers in parantheses represent the chain that is currently being explored by that machine. Arrows indicate communication occuring between machines to exchange chain indices. To keep track of which chains are stored on which machine, we introduce the dist_array of type PermutedDistributedArray, described in Section 3.3.1, which is of length N. The j-th element of dist_array at a given time step indicates which machine number is storing chain index j. The bolded curve in the figures to the right indicate the trajectory of the first Replica over the course of each of the communication steps. Top left: the PT replicas are initialized and the first communication step is proposed. At this time step, dist_array = [1, 2, 3, 4]. Top right: after the first successful communication step, we now have dist_array = [2, 1, 4, 3]. Bottom left: The second communication step is completed and dist_array = [4, 2, 3, 1].

it is important to understand the behaviour of the @threads macro.

Our focus in the remainder of this section is to describe how our implementation solves the two above issues.

4.1 Distributed reduction and floating point non-associativity

After each round of PT, the machines need to exchange information such as the average swap acceptance probabilities, statistics to adapt a variational reference and adjust annealing parameters, and so on. For instance, suppose our state is univariate and real-valued and that each Replica keeps track of the number of times the target chain N is visited as well as the mean of the univariate states from the target chain. To obtain the final estimate of the mean of the target distribution, we would like to pool the mean estimates from each of the Replicas, weighted by the number of times that each Replica visited the target chain. This process involves summing floating-point values that are located on each of the Replicas/machines and is an example of a reduction scheme.

To illustrate why distributed reduction with floating point values can violate strong parallelism invariance if not properly implemented, we consider the following toy example. Suppose we have 8 machines storing the floating point numbers $\{1x, 2x, \ldots, 8x\}$, as illustrated in Fig. 7, where we use $x = 10e^1 \approx 27.1828$ in the following examples. In this case, if our reduction procedure is to sum the floating point numbers, we know that our final answer should be approximately 36x. However, depending on the exact order in which floating point addition is carried out, the answers might not all be the same and exactly equal to 36x. For instance, in Fig. 7 we see that the order of operations for eight machines would be given

by
$$((1x+2x)+(3x+4x))+((5x+6x)+(7x+8x)) \\ \approx 978.5814582452562.$$

In contrast, with two machines, one possible order of operations might be

$$(((1x+2x)+3x)+4x)+(((5x+6x)+7x)+8x)$$

 $\approx 978.5814582452563.$

A possible reduction tree for two machines is illustrated in Fig. 8.

To avoid the issue of non-associativity of floating point arithmetic, we ensure that the order in which operations are performed is exactly the same, irrespective of the number of processes/machines and threads. This is achieved by making sure that every value to be added—if addition is our reduction operation—is a leaf node in the reduction tree, irrespective of the number of machines available to perform the reduction. For instance, if N values are to be reduced, then the reduction tree would have N leaf nodes. If M machines are available, these machines are then assigned in such a way that the order of operations is as if there were N machines available. To do so, it may be necessary for a machine to "communicate with itself", imitating the behaviour that would be present if there were N machines available. Fig. 9 and Fig. 10 illustrate the reduction procedure for eight and two machines, respectively.

4.2 Splittable random streams

Another building block towards achieving SPI is a *splittable ran-dom stream* [17, 6]. Julia uses *task-local* random number generators, a notion that is related to (but does not necessarily imply) strong parallelism invariance. A *task* is a unit of work on a machine. A task-local RNG would then mean that a separate RNG is

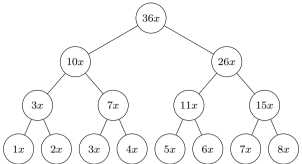


Fig. 7: Adding eight floating point numbers $\{1x,2x,\ldots,8x\}$ across eight machines. Additions in each row of the tree can be performed in parallel. The final result is stored in the root node of the tree and can be represented by the expression ((1x+2x)+(3x+4x))+((5x+6x)+(7x+8x)), indicating the order of operations.

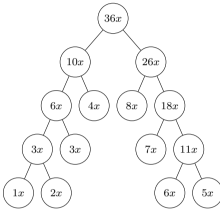


Fig. 8: One possible way of adding eight floating point numbers $\{1x, 2x, \ldots, 8x\}$ across two machines. The final result is stored in the root node of the tree and can be represented by the expression (((1x+2x)+3x)+4x)+(((5x+6x)+7x)+8x)). Note that the order of operations in this expression is different from the one presented in Fig. 7.

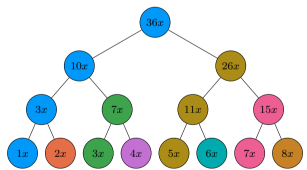


Fig. 9: Addition of eight floating point numbers across eight machines with a guarantee on SPI. Each machine is represented by a different colour. The final result can be represented by the expression ((1x+2x)+(3x+4x))+((5x+6x)+(7x+8x)).

used for each unit of work, hopefully implying strong parallelism invariance if the number of tasks is assumed to be constant. Unfortunately, this is not the case when a separate task is created for each thread of execution in Julia. We note that the @threads macro in Julia creates nthreads() tasks and thus nthreads() pseudoran-

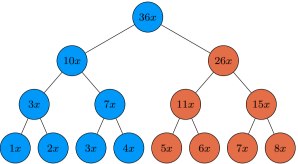


Fig. 10: Addition of eight floating point numbers across two machines with a guarantee on SPI. Each machine is represented by a different colour. The final result can be represented by the expression ((1x+2x)+(3x+4x))+((5x+6x)+(7x+8x)), which is the same as that given by eight machines in Fig. 9.

dom number generators. This can break strong parallelism invariance as the output may depend on the number of threads.

To motivate splittable random streams, consider the following toy example that violates our notion of SPI:

```
using Random
import Base.Threads.@threads

println("Num. of threads: $(Threads.nthreads())")

const n_iters = 10000
result = zeros(n_iters)
Random.seed!(1)
@threads for i in 1:n_iters
    result[i] = rand()
end
println("Result: $(last(result))")
```

With eight threads, this outputs:

```
Num. of threads: 8
Result: 0.25679999169092793
```

Julia guarantees that if we rerun this code, as long as we are using eight threads, we will always get the same result, irrespective of the multi-threading scheduling decisions implied by the @threads-loop. Internally, Julia works with task-local RNGs and the @threads macro spawns nthreads() number of task-local RNGs. For this reason, with a different number of threads, the result is different:

```
Num. of threads: 1
Result: 0.8785201210435906
```

In this simple example above, the difference in output is perhaps not too concerning, but for our parallel tempering use case, the distributed version of the algorithm is significantly more complex and difficult to debug compared to the single-threaded one. We therefore take task-local random number generation one step further and achieve SPI, which guarantees that the output is not only reproducible with respect to repetitions for a fixed number of threads, but also for different numbers of threads or processes.

A first step to achieve this is to associate one random number generator to each Replica. To do so, we use our SplittableRandoms.jl package, which is a Julia implementation of Java SplittableRandoms. Our package offers an implementation of SplitMix64 [25], which allows us to turn one seed into an arbitrary collection of pseudo-independent RNGs. A quick example of how to use the SplittableRandoms.jl library is given below. By splitting a master RNG using the split() function, we can achieve SPI even with the use of the @threads macro.

```
using Random
using SplittableRandoms: SplittableRandom, split
import Base.Threads.@threads

println("Num. of threads: $(Threads.nthreads())")

const n_iters = 10000
const master_rng = SplittableRandom(1)
result = zeros(n_iters)
rngs = [split(master_rng) for _ in 1:n_iters]
Random.seed!(1)
@threads for i in 1:n_iters
    result[i] = rand(rngs[i])
end
println("Result: $(last(result))")
```

With one and eight threads, the code above outputs

```
Num. of threads: 1
Result: 0.4394633333251359
```

```
Num. of threads: 8
Result: 0.4394633333251359
```

5. Related work

Automated software packages for Bayesian inference have revolutionized Bayesian data analysis in the past two decades, and are now a core part of a typical applied statistics workflow. For example, packages such as BUGS [19, 20, 21], JAGS [14], Stan [9], PyMC3 [23], and Turing.jl [11] have been widely used in many scientific applications.

These software packages often provide two key user-facing components: (1) a probabilistic programming language (PPL), which allows users to specify Bayesian statistical models in code with a familiar, mathematics-like syntax; and (2) an inference engine, which is responsible for performing computational Bayesian inference once the model and data have been specified. Pigeons.jl focuses on the development of a new inference engine that employs distributed, high-performance computing.

Inference engines available in existing software packages are varied in their capabilities and limitations. For instance, the widely-used Stan inference engine is only capable of handling real-valued (i.e., continuous) parameters. Other tools, such as JAGS, are capable of handling discrete-valued parameters but are limited in their capability to handle custom data-types (e.g. phylogenetic trees). Of those that have the capability to model arbitrary data types, none have an automatically distributed implementation to our knowledge. Turing, jl [11] offers another popular inference engine and PPL, how-

ever Pigeons.jl allows one to interface with several different PPLs as well as to easily perform distributed computation.

There is also a vast literature on distributed and parallel Bayesian inference algorithms [3, 5, 8, 16, 15, 18, 24, 28, 29, 32]. These methods unfortunately do not lead to widely usable software packages because they either introduce unknown amounts of approximation error, involve significant communication cost, or reduce the generality of Bayesian inference.

6. Conclusion

Pigeons.jl is a Julia package that enables users with no experience in distributed computing to efficiently approximate posterior distributions and solve challenging Lebesgue integration problems over a distributed computing environment. The core algorithm behind Pigeons.jl is distributed, non-reversible parallel tempering [27, 26]. Pigeons.jl can be used in a multi-threaded context, as well as distributed over up to thousands of MPI-communicating machines. Further, Pigeons.jl is designed so that for a given seed, the output is *identical* regardless of the number of threads or processes used.

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