

# Nested sampling with peers



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# Abstract

A nested sampling algorithm is a Bayesian approach to computing and comparing models and generating samples from posterior distributions. We introduce a general Monte Carlo method based on Nested Sampling, which name is Nested Sampling with peers, this method generates one particle above the threshold from the last iteration by querying the params from the server and updating it. We describe the new method over a test case and find that it has better accuracy than the original MCMC-based nested sampling with the same computational overhead. Put your abstract here. The abstract should contain a brief summary of the aim, methodologies, finding and conclusions of the dissertation. The abstract should normally be fewer than 350 words.



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# Chapter 1

## Introduction

### 1.1 Bayesian Analysis

The Bayesian Analysis problem is in fact a parameter estimation problem, The term *parameter* means *unknown quantity*, however in real world, we don't have enough information to decide a quantity, so we need Bayesian Analysis to help us. The parameter you interested in are denoted by  $\theta$ , to estimate them, first of all, you need to model a probability distribution on the *hypothesis space*, a *hypothesis space* is the collection of possibilities, this means that you are modeling the initial assumptions, and the probability distribution is called the *prior*, it is the distribution of  $\theta$ . The data set is called  $D$ , after that we can use Bayes' Theorem to determine the *posterior distribution*:

$$p(\theta|D) = \frac{p(\theta)p(D|\theta)}{p(D)} \quad (1.1)$$

in the above equation,

1. The posterior distribution is  $p(\theta|D)$ , given the data set  $D$  with the conditional distribution of  $\theta$ .
2. The prior distribution is  $p(\theta)$ .
3. The likelihood is  $p(D|\theta)$ , it means the conditional probability of observing the data.
4. The denominator  $p(D)$  is called marginal likelihood or evidence, it doesn't depend on the parameter  $\theta$ .

The posterior distribution is usually more constricted than the prior distribution, indicating that we have gained some insights from the data and that our level of uncertainty about the parameter values has been lowered. See the Figure 1.1 for how we often see the updating from the prior distribution to a posterior distribution.

### 1.1.1 Marginal Likelihood

The marginal likelihood shows how does the model be constructed. From probability theory we can learn that If you integrate over the posterior distribution, you must get 1 at the last. So we can write the marginal likelihood in the following form:

$$p(D|I) = \int p(\theta|I)p(D|\theta, I)d\theta \quad (1.2)$$

where  $I$  represents the underlying assumptions and information, and the integral is N-dimensional, go through all the continuous parameter space, the  $I$  is usually ignored and the marginal likelihood can be denoted as  $Z$ :

$$Z = \int p(\theta)p(D|\theta)d\theta \quad (1.3)$$

when sum the discrete parameter  $\theta$ , it also has the following form:

$$Z = \sum p(\theta)p(D|\theta) \quad (1.4)$$

The ratio of evidence values is always called Bayes factors, calculating the value of  $Z$  is important for comparing different model assumptions, so the Z-value allows the results to be predicted into the future because the predicted model can be compared with the current model without the need for recalculation, in short, this parameter measures how well a model fits the data. However, the standard algorithm such as Markov chain Monte Carlo (MCMC) are considered for posterior distribution, it cannot provide the evidence like Bayesian Inference, for it only gives a collection of normalised posterior, it will be a challenge to get the evidence in Bayesian analysis.

## 1.2 Nested Sampling

### 1.2.1 Model Selection with Nested Sampling

Consider two mutually exclusive model  $M_1$  and  $M_2$ ,  $M_1$  has the parameter  $\theta_1$  and  $M_2$  has the parameter  $\theta_2$ , from previous method we discussed, we can calculate the posterior distribution of  $\theta_1$  :

$$p(\theta_1|D, M_1) = \frac{p(\theta_1|M_1)p(D|\theta_1, M_1)}{p(D|M_1)} \quad (1.5)$$

Same, calculate the posterior distribution of  $\theta_2$

$$p(\theta_2|D, M_2) = \frac{p(\theta_2|M_2)p(D|\theta_2, M_2)}{p(D|M_2)} \quad (1.6)$$



For simplicity, we often calculate the ratio of posterior distributions to compare models:

$$\frac{p(M_1|D)}{p(M_2|D)} = \frac{p(M_1)}{p(M_2)} \times \frac{p(D|M_1)}{p(D|M_2)} \quad (1.7)$$

The result of this formula is called *posterior odds*, the posterior probability of M2 over M1 depends on the prior probability: is M2 more credible than M1 before considering the data? Another ratio is the ratio of likelihoods, sometimes called the Bayesian coefficient; how likely is the data for hypothetical M2 versus hypothetical M1? These possibilities are not the likelihood of a particular value of the parameter, but the likelihood of the entire model. We have to use the marginal likelihood to make a better progress.

### 1.2.2 Sorting

In multiple dimensions, the direct integration calculation for the edge likelihood becomes unrealistic, we define the  $X$  as the model's prior mass, it can be integrated in this form:

$$X(\lambda) = \int_{L(\theta) > \lambda} \pi(\theta) d\theta \quad (1.8)$$

we can write the inverse function  $L(X)$ , the marginal likelihood *i.e.* evidence will be in this form:

$$Z = \int_0^1 L(X) dX \quad (1.9)$$

To get the inverse function, we can divide the prior into little particles and then sort all the particles by likelihood. In geometrical terms, the function  $L(X)$  is monotonically diminishing, and the likelihood decreases as the value of  $X$  increases from 0 to 1, as shown in Figure 1.1. From the figure 2.1 we can find that when approaching the maximum likelihood,  $X$  will keep decreasing.

The explanation of 1.5 is that the  $L(X)$  is the probability corresponding to  $\lambda$  that makes the probability,  $L(\theta)$ , greater than  $\lambda$  for  $X$ .  $L(0.4) = 0.01$  may be interpreted as a 40% probability that  $\pi(\theta)$  will be greater than 0.01 for  $\theta$  chosen from the prior. To evaluate the mapped integral in 1.5, Skilling proposed a simple trapezoidal rule by calculating weighted sums:

$$Z \approx \sum_{i=1}^j w_i L_i(w_i = \frac{1}{2}(X_{i-1} - X_{i+1})) \quad (1.10)$$

In every iteration  $i$ , the likelihood is  $L(X_i)$ , and  $j$  is total number of iterations.

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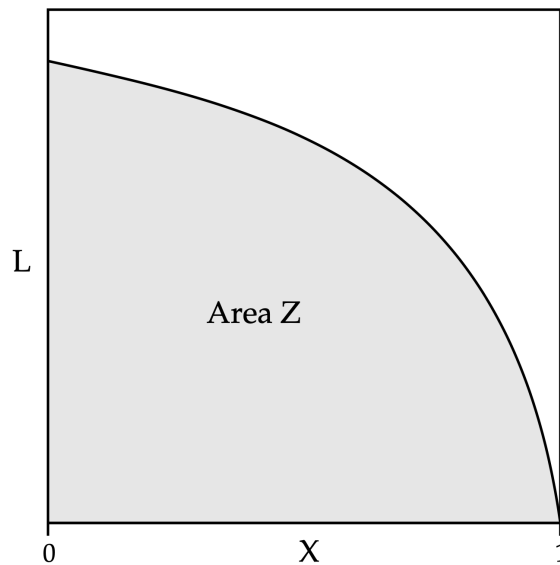


Figure 1.1: Likelihood function with area  $Z$ .

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### 1.2.3 A easy example

Imagine we have a model whose parameter is a single  $X$ , and if the parameter  $X$  follows the uniform prior distribution  $\pi(\theta)$  from 0 to 1, it also has a likelihood function  $L(X)$  which is a decreasing function of  $X$ , the Figure 1.2. illustrate the graph of the example, It's easy to find that in this situation, the evidence will be  $Z = \int_0^1 L(X)DX$

### 1.2.4 Nested Sampling Algorithms

This section will discuss the main idea of Nested Sampling, it is a Monte Carlo algorithm and not a typical MCMC algorithm in technical. John Skilling proposed the algorithm specifically for approximating these marginal integrals, *i.e.*, evidence, and it also generates samples from

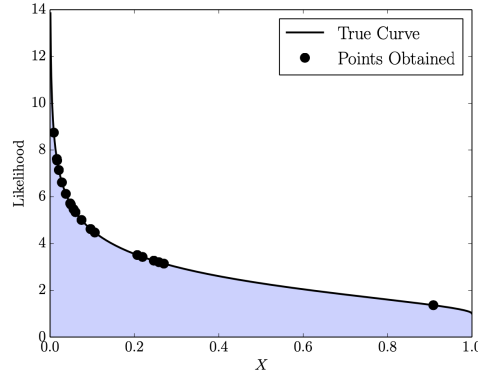


Figure 1.2: A simple model estimate problem with only one parameter

the posterior distribution. In the prior distribution  $\pi(\theta)$ , it needs to generate  $N$  *particles or live points*  $\theta$ , just like an vector to initialize the starting state. To approximating the evidence, there's some important steps, if we find the points with the worst likelihood, always denoted by  $L_{worst}$  and this is often associated with the highest  $X$  value, If different particle  $N$  are taken, the highest  $X$  value will always be close to  $X = 1$  and not close to  $X = 0$ . Skilling (2006) gives a simple treatment by estimating  $X_{worst} = e^{(\frac{-1}{N})}$ , and we also know the probability of this point.

By generating a new point  $X_1$  from the prior distribution  $\pi(\theta)$  to replace the worst  $X_{worst}$ , more points can be obtained,

and the restriction is that the likelihood of the point must be higher than  $L_{worst}$ . Also, the prior distribution  $\pi(\theta)$  is sampled from the conditional prior,  $\pi(\theta|L(\theta) > \lambda_{min,1})$ , we can denote the new set of particles as  $\omega_1 = \{\theta|L(\theta) > \lambda_{min,1}\}$ , it's a subset of the whole parameter space  $\omega$ . And in the second iteration, the estimation of the  $X$  value who has a worst particle is  $e^{(\frac{-1}{N})} \times e^{(\frac{-1}{N})} = e^{(\frac{-2}{N})}$ , because we already know the likelihood, so we can continue the iteration. So the second step to generate the new point  $X_2$  and the  $X_{worst}$ ,  $X$  will be repeated, we use  $\omega_2$  to denote them, and we should know that  $\omega_1 \subset \omega_2$ , so the  $X_{worst}$  will be replaced, too, from the prior distribution  $\pi(\theta)$  from the conditional prior,  $\pi(\theta|L(\theta) > \lambda_{min,2})$ , so we can write when the iteration run  $i$  times and it will generate a sequence of  $\lambda_{min,i}$  will be like following:

$$\lambda_{min,1} < \lambda_{min,2} < \dots < \lambda_{min,i-1} < \lambda_{min,i}$$

,because every  $\omega_i$  is subset of the whole space  $\omega$ , so it will decrease continuously, so it can be viewed as a reduction or compressing of the parameter space  $\omega$  and the subspaces are nested within each other, so we have the term *nested sampling*. The points  $i$  discarded within each loop can be sampled using the MCMC algorithm by finding a random copy of a particle

as the initial value, proposed by Skilling (2006). Nowadays, there are also studies to investigate whether this approximation can be satisfied efficiency requirements of MCMC based on the shape of the likelihood function and the MCMC chosen to explore the restricted prior distribution. (Syarafana Abdul Rahman)

To estimate the evidence  $Z$  we can use the equation give in 1.10, and changing the  $w_i = \frac{1}{2}(X_{i-1} - X_{i+1})$  into  $\lambda_{min,i}$ , which means that the  $Z$  can be estimate like this:

$$\hat{Z} \approx \sum_{i=1}^j w_i L_i(w_i = \frac{1}{2}(X_{i-1} - X_{i+1})) = \sum_{i=1}^j w_i \lambda_{min,i} \quad (1.11)$$

The value of  $Z$  will be updated each time based on the results of this approximation until the algorithm terminates according to some termination criterion.

### 1.2.5 Prior Mass, X

Skilling(2006) proposed a method that can decide the prior mass  $X$  statistically: start from  $X_0 = 1$ , and at each step  $i$ , a new point  $X_i$  is drawn randomly from the prior, and the  $X_i$  has the restriction that  $X_i < X_{i-1}$ , this will generate  $N$  live points that follows the *Uniform*(0,  $X_{i-1}$ ). The  $X$  can be generated like this:

$$X_0 = 1, X_i = t_i X_{i-1} \quad (1.12)$$

The  $t_i$  is from the *Beta* distribution where  $Beta(N, 1)$ , it's the Beta shrinkage, which lowers estimation errors, and  $t_i = \frac{X_i}{X_{i-1}}$ , where  $t_i \in [0, 1]$ . The recurrence is:

$$P(t_i) = N t_i^{N-1} \quad (1.13)$$

, and it has the expectation that  $E(\log(t_i)) = -\frac{1}{N}$ , and also has the variance that  $Var(\log(t_i)) = \frac{1}{N^2}$ , after  $i$  iterations, and the  $\log^t$  are independent, a rough approximation for the prior mass is  $\log(x_i) = \sum_{j=1}^i \log(t_j)$ , (Skilling, 2006), and there's another estimation is  $\log(x_i) \in [\frac{-(i-\sqrt{i})}{N}, \frac{-(i+\sqrt{i})}{N}]$ , proposed by (Sivia and Skilling, 2006), this means that The reduction of the prior mass will be linear along the logarithmic X scale.

### 1.2.6 Measuring

There's another output of Nested Sampling Algorithm is information, this is to find how much were we learnt from the data can be measured by the negative relative entropy(Russel, 2017), also Burnham and Anderson had proposed the Kullbeck Liebler divergence (Burnham and Anderson, 2001), denoted by information  $H$ , this method measures the discrepancy between two distributions, prior distribution and posterior distribution:

$$H = \int p(\theta|D) \log\left(\frac{p(\theta|D)}{\pi(\theta)}\right) d\theta \quad (1.14)$$

here  $p(\theta|D)$  is the posterior distribution. The  $\pi(\theta)$  is the prior distribution, and it can greatly influence the evidence  $Z$ , if the size of the region of prior mass was concentrated by posterior distribution. Skilling (2006) proposed that the region is  $e^{-H}$ , naively, a prior distribution that overlaps with the likelihood function so that the parameter values are consistent will yield less information. This can estimate the uncertainty of  $\log(Z) = \sqrt{H/N}$  (Skilling, 2006). On the other hand, if the prior distribution and the likelihood function are concentrated in different regions, more information will be obtained. After looking at the data, if the prior greatly changed, more information is obtained from the data (Russel, 2017). If we use  $L(X_i)$  to represents the likelihood  $X$  at every iteration  $i$ , the value  $H$  will be like this:

$$H = \sum_i w_i L(X_i)/Z \log(L(X_i)/Z) \quad (1.15)$$

### 1.2.7 Nested sampling procedure

In the begging, the Nested Sampling Algorithm will generating  $N$  particles  $\{\theta_1, \theta_2, \dots, \theta_n\}$  from the prior, also has every likelihood  $\{L(\theta_1), L(\theta_2), \dots, L(\theta_n)\}$ , and the lowest value is denoted by  $L_{min,i}$ , in every iteration  $i$ , there are  $j$  steps.

The Nested Sampling Algorithm has the following steps:

---

#### Algorithm 1 A

---

Begin with N particles  $\{\theta_1, \theta_2, \dots, \theta_n\}$  from the prior.

**initialise**  $Z = 0, X_0 = 1, i = 0$

**repeat**

for  $i = \{1, 2, \dots, j\}$

**Find** the particle with the current lowest likelihood values as  $L_{min,i}$ .

**Crude** estimate the value as  $X_i = \exp(\frac{-i}{N})$

**Set**  $w_i = X_{i-1} - X_i$

**Save** the propertied  $(X_i, L_{min,i})$

**Increment**  $Z$  by  $L_{min,i} w_i$

**Generate** a new particle from the prior to replace the point above found with  $L(\theta) > L_{min,i}$

**Increment** the  $Z$  by  $N^{-1}(L(\theta_1) + \dots + L(\theta_N))X_j$

**until** Enough iterations have been performed

---

### 1.2.8 Nested Sampling Termination Criterion

To terminate by default, the algorithm has to set some termination criterion. The termination criterion will depends on the increment of the evidence, which means that sum of the evidence

has been found enough. The current found likelihood will make the evidence more significant and  $Z_j$  will times a small fraction  $f$ :

$$\max(L(\theta_1), L(\theta_2), L(\theta_3), \dots, L(\theta_N))X_j < fZ_j \quad (1.16)$$

there's some small fraction  $f$ . In the previous section we discuss about information, this can be used as a criterion of termination. The  $L_i w_i$  is related to the likelihood  $L_i$  and width  $w_i$ , it will increase faster than the widths  $w_i$  decreases, and this leads to more regions. (Skilling, 2006). So In the prior mass, the regions we are interested in is  $X \approx e^{-H}$ , and  $X_i \approx e^{-i/H}$ , this means the region from the prior found most of the posterior mass. So Skilling proposed a plausible termination criterion that when the iteration  $i$  has exceeded  $NH$ , which means that the information  $H$  times the count of live points  $N$ , and practically the number should be set to  $2NH$  (Russel, 2007). There's also a scaling constant to ensure that the nested is appropriate, we have:

$$ri > N_r H = \sqrt{r} H N \quad (1.17)$$

So in this inequality, we can see that we need at least  $\sqrt{r} H N$  samples to substitute (Henderson and Goggans, 2013). However, in most circumstances, the user decide when to stop the iteration, because user knows how much information do they need in the current region.

## 1.3 Distributed System

A distributed system is a computing environment in which various components are decentralized among multiple computers (or other computing devices) on a network. These devices work together and synchronize their efforts to get the job done more efficiently than if a single device were responsible for the task. And a distributed system has two main characteristics:

- Autonomous computing elements are collected together, and they are independent. In a distributed system, when a computer or node has some crash will not affect other computers in the same system, it will not made known to the other components with which it communicates immediately.
- Running program in the same time are a very normal thing in the computer world, all of us doing things without disturbing others, and when it is needed, we can share our resources together. The synchronization of concurrently executing programs for shared resources is also an important and recurring topic.

### 1.3.1 Example of a distributed system

Google is the best example of distributed system, it is a pioneer and leader in the field of web search technology, and it has a very big and complicated sophisticated distributed system, such

as searching service and other applications such as youtube and Google Earth. To design and build this sophisticated distributed system, google has to put efforts from the following points:

1. Underlying infrastructure such as big network rooms and databases,as well as various matching settings such as radiators Google has numerous data centers scattered around the world. At least 12 significant Google data center installations are located in the United States. The largest known centers are located in The Dalles, Oregon; Atlanta, Georgia; Reston, Virginia; Lenoir, North Carolina; and Moncks Corner, South Carolina (Wikipedia).
2. To support hundreds of millions files fast and stable reading, writing and searching google has to optimize the distributed file system in many ways such as writing better programs.
3. To synchronize the files' reading and writing, it also needs a lock system.
4. A programming model to supports managing very large parallel and distributed computations on the underlying physical infrastructure.

### 1.3.2 Distributed system models

There're three type of models that can provide an abstract of distributed system:

1. *Physical models* show an aspect of distributed system from hardware, explains their underlying connected network and data systems.
2. *Architectural models* show an aspect from computational elements, computational elements are a single computer or a collection of computers supported by an appropriate network interconnection.
3. *Fundamental models* To examine individual aspects of distributed systems, we take an abstract perspective, it includes three kinds of models: interaction models, which consider the structure and sequencing of the communication between the elements of the system; failure models, which consider the ways in which a system may fail to operate correctly and security models, which consider how the system is protected against attempts to interfere with its correct operation or to steal its data (George, 2006).

In the individual process environment, most of computers use client-server architecture, it's a very simple but very important architecture in the history and also use in today's world, the process in this architecture is a role of client or server, in particular, client processes communicate with individual server processes in potentially independent hosts to ask for the resources like images, videos they manage. Also, a server can also be clients and asks for resources from other servers, for example a search is not only a server, but also a client, for the reason

it have to respond queries from users, which means clients, then it needs to run crawlers that act as clients of other web servers. Responding to user queries can be called server task, and requesting other web servers is the client task, these two tasks are independent with each other, so they can run concurrently.

The figure 1.3 shows the architecture of client-server modes.

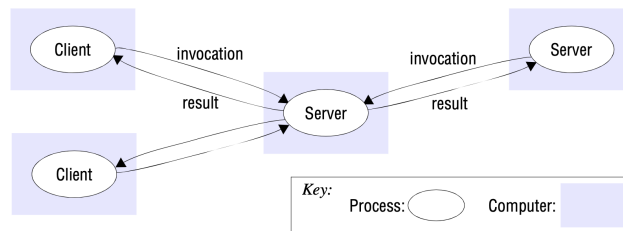


Figure 1.3: Client-server architecture

Although client-server architecture offers a straightforward and relatively simple way to share data and manage other resources, it is poorly scalable. The server address is relatively single, so it is difficult to expand the computer capacity and network bandwidth. In the peer-to-peer structure, all processes are of the same status and no distinction is made between client and server processes or the computers on which they run. In fact, all participating processes run the same programs and provide the same set of APIs to each other. API stands for application programming interface, it's a connection between computers or between computer processes. Peer-to-peer architecture can lead to a very useful result that the resources will grow with the number of users. This is mainly due to the fact that peer-to-peer was designed with the idea that the network and computing resources owned by the users of a service could also be used to support that service. The figure 1.4 shows the architecture of client-server modes.

Peer-to-peer architectures can utilize the resources (hardware resources, software resources, data) in almost any computer in the system to accomplish a specific task. Here's the advantages of peer-to-peer architecture:

1. Better parallel processing capabilities compared to Client-server architecture, which gets slower and slower as more and more clients are added, but p2p networks do not change very much, and all clients can provide resources, including bandwidth, storage space, and computing power.



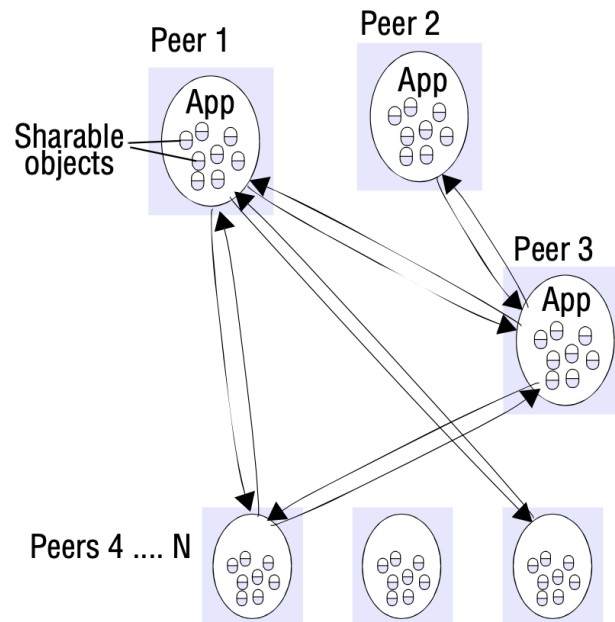


Figure 1.4: Peer-to-peer architecture

2. No central server, relying on user groups (peers) to exchange information, reducing network transmission nodes
3. No need to invest a lot of money in server software and hardware.

Also, there's some disadvantages of peer-to-peer:

1. More complex architecture, NAT isolation usually exists between nodes and direct communication is not possible, resulting in the need for additional pathway construction process and coordinated management of resources for each node
2. P2P nodes are spread throughout the Internet, bringing monitoring problems to developers, organizations, and governments
3. Used in large-scale networks, resource sharing is disorderly and difficult to manage
4. Lower security

### 1.3.3 Interprocess communication: sockets

There are various methods of inter-process communication under unix/linux system, such as

- Anonymous pipeline (pipe): communication between parent and child processes
- Named pipes (fifo): communication with unrelated processes
- Shared memory (shm): can communicate with unrelated processes, need to handle their own process synchronization, such as with semaphore.
- Unix domain sockets (unix socket): can communicate with unassociated processes, using the socket interface.
- Network socket: can communicate across hosts, using the socket interface.

Sockets is one of the most important communication style in distributed system, A socket is an abstraction of an endpoint for bi-directional communication between application processes on different hosts in a network. A socket is one end of the communication between processes on the network and provides a mechanism for application layer processes to exchange data using network protocols. In terms of its position, a socket is connected to the application process and to the network protocol stack, and is the interface through which the application communicates with the network protocol and interacts with the network protocol stack. The sockets can be defined as:

$$\text{sockets} = \text{protocol} / \text{ip} : \text{ports} \quad (1.18)$$

The C++ offers many APIs for socket programming:

1. **socket** Use the function to create a socket:

---

```
int sockfd = socket(domain, type, protocol)
```

---

2. **bind** function after create the socket, the **bind** function binds the socket to the address and port number all together.

---

```
int bind(int sockfd, const struct sockaddr *addr,
socklen_t addrlen);
```

---

3. **listen** the server will listen to the ports and waiting for client's connect request.

---

```
int listen(int sockfd, int backlog);
```

---

4. **accept** The `textbfaccept` function is called by the TCP server to return the next completed connection from the completed connections queue. If the completed connections queue is empty, then the process is put to sleep (assuming the default blocking mode of socket silence)

---

```
int listen(int sockfd, int backlog);
int new_socket= accept(int sockfd, struct sockaddr *addr,
socklen_t *addrlen);
```

---

5. **connect** The **connect** function is usually used to establish a tcp connection on the client side.

---

```
int connect (int sockfd, struct sockaddr * serv_addr, int
addrlen);
```

---

Here is a basic of network programming procedure using sockets APIs in C++:

---

```
// server.cpp
// socket
int listenfd = socket(AF_INET, SOCK_STREAM, 0);
// bind
struct sockaddr_in addr;
addr.sin_family = AF_INET;
addr.sin_port = htons(8000);
addr.sin_addr.s_addr = INADDR_ANY;
if (bind(listenfd, (struct sockaddr*)&addr, sizeof(addr)) == -1)
{
    std::cout << "Error: bind" << std::endl;
    return 0;
}
// listen
if(listen(listenfd, 5) == -1) {
    std::cout << "Error: listen" << std::endl;
    return 0;
}
// accept
int conn;
char clientIP[INET_ADDRSTRLEN] = "";
struct sockaddr_in clientAddr;
socklen_t clientAddrLen = sizeof(clientAddr);
while (true) {
    conn = accept(listenfd, (struct sockaddr*)&clientAddr,
        &clientAddrLen);
    inet_ntop(AF_INET, &clientAddr.sin_addr, clientIP,
        INET_ADDRSTRLEN);

    char buf[255];
    while (true) {
```

```

        memset(buf, 0, sizeof(buf));
        int len = recv(conn, buf, sizeof(buf), 0);
        buf[len] = '\0';

        std::cout << buf << std::endl;
        send(conn, buf, len, 0);
    }

    close(conn);
}
close(listenfd);
return 0;

```

---

```

// client.cpp
// socket
int client = socket(AF_INET, SOCK_STREAM, 0);

// connect
struct sockaddr_in serverAddr;
serverAddr.sin_family = AF_INET;
serverAddr.sin_port = htons(8000);
serverAddr.sin_addr.s_addr = inet_addr("127.0.0.1");
if (connect(client, (struct sockaddr*)&serverAddr,
    sizeof(serverAddr)) < 0) {
    std::cout << "Error: connect" << std::endl;
    return 0;
}
std::cout << "...connect" << std::endl;
char data[255];
char buf[255];
while (true) {
    send(client, data, strlen(data), 0);
    int len = recv(client, buf, sizeof(buf), 0);
}
close(client);
return 0;

```

---

The figure 1.5 shows diagram of client-server modes in socket programming, which needs many procedure to communicate with other computers.

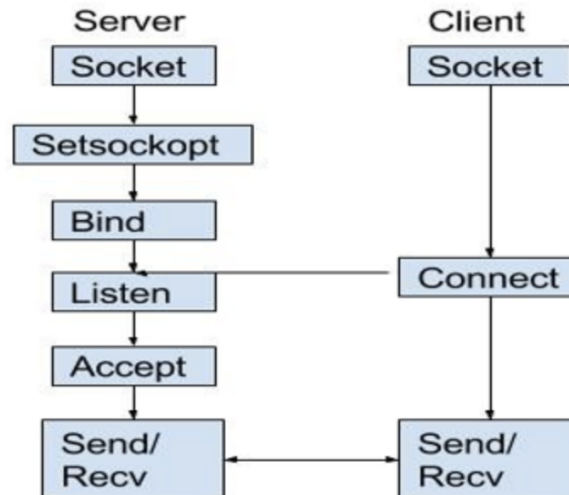


Figure 1.5: Socket diagram

### 1.3.4 JavaScript Object Notation: (JSON)

*JSON* is a text format for data exchange, the current standard is ECMA-404, Although JSON is derived from the JavaScript language, it is a data format that can be used in any programming language. There are formats with similar functionality, such as XML and YAML, of which JSON has the simplest syntax.

For example, when a dynamic web page wants to get data from the server, the server looks up the data from the database and converts it into JSON text format:

---

```

{
  "title": "Distributed System",
  "subtitle": "How to design better distributed system",
  "author": [
    "Feishuang Wang",
    "Json Jack",
  ],
  "year": 2010,
  "weight": 1.6,
  "hardcover": false,
  "publisher": {
    "Company": "Education",
    "Country": "China"
  }
}

```

```
    },  
    "blog": null  
}
```

---

The scripting code of the web page can then parse this JSON text into internal data structures for use.

As you can see from this example, JSON is a tree structure, while JSON contains only 6 data types.

1. null: it represents by **null**.
2. boolean: it represents by **true** or **false**.
3. number: it usually represents by float numbers.
4. string: it represents by ‘...’
5. array: it represents by [...]
6. object: it represents by {...}

JSON parsing fulfills 3 main requirements.

- Parse JSON text into a tree data structure (parse).
- Provide an interface to access the data structure (access).
- Convert the data structure into JSON text (stringify).

The figure 1.6 the process of parsing a json object:



Figure 1.6: Json parsing

There're many advantages to use json as a format to exchange data:

- Compared with XML, JSON is more concise, we can see the content at a glance, easy to check and troubleshoot.
- JSON is lighter, more efficient in writing, transferring, and parsing.

- JSON uses compression technology in the transmission process, which saves broadband.
- Numerous language support, such as javascript, python, C, C++ and other mainstream languages are supported.

### 1.3.5 Berkeley Open Infrastructure for Network Computing: BOINC

*BOINC* is one of the leading distributed computing platforms, a decentralized computing system developed by the Department of Computer Science at the University of California, Berkeley. Originally designed for the *SETI@home* project, this project is SETI stands for Search for Extraterrestrial Intelligence. What this project does is to analyze the data observed by radio telescopes around the world to find three equally spaced convex waves. It is not easy to observe such convex waves because the cosmic waves are very weak and are strongly influenced by the electromagnetic properties of the universe itself.

This kind of task is very suitable for distributed computing, and SETI joined BOINC in the sixth year after the project was established. The nodes that contribute computational power around the world are allowed to join together to find it. It mainly for research institutions to provide a distributed computing platform. BOINC was developed because many research institutions lack funds, but many research projects require very large computation. So we want to use the distributed computing method to handle large amount of data. The *BOINC* has many excellent ideas for developers to learn:

- Provide dedicated support for the development of advanced algorithms.
- Applying artificial intelligence techniques to analyze and process gravitational wave big data, developing more efficient and intelligent algorithms for gravitational wave data analysis, and increasing the possibility of identifying theoretically unknown gravitational waves.
- Open up gravitational wave astronomical data and hold challenges to attract companies and citizen scientists to participate and use group intelligence to overcome the problem.

## 1.4 Spakslab Problem

Spike and slab is a more graphic description of a prior. For a random variable  $X$ , a spike and slab prior will either gets some value  $v$ , which can be called *spike*, or gets some other prior  $p_{slab}(x)$ , which will be called *slab*. The figure 1.7 shows the spike and slab prior:

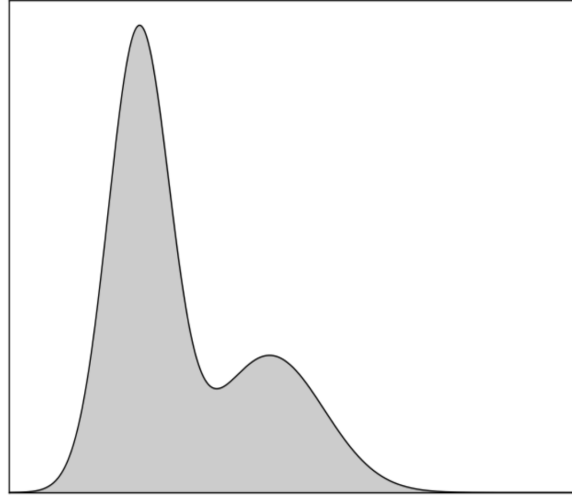


Figure 1.7: A view of the spike and slab prior in 2 dimensions. The higher peak is the "spike" and the lower peak is the "slab"

The spike and slab prior is sparsity inducing in the case that  $v = 0$  and  $X = 0$ , it will give a principled alternative to sparsity-inducing regularisers. It is used to do Bayesian sparse learning, to obtain a sparse model from which further tasks such as feature selection and model compression can be done. From Bayesian linear regression to Bayesian neural networks, any probabilistic model in a Bayesian framework can be used in spike and slab prior. The spike and slab correspond to the sampling process that

1. For each parameter  $w_i$ , we first assign a binary indicator  $s_i$ , which is sampled from a Bernoulli distribution  $Bern(p_0)$  and has only two possible values, 0 or 1.
2. If  $s_i = 1$ , then we continue to draw from a relatively normal and flat Gaussian distribution, such as  $N(0, 1)$ , at which point the value of  $w_i$  has a high probability of not being 0. Conversely, if  $s_i = 0$ , then we continue to draw from an abnormal and steep Gaussian distribution, such as  $N(0, 0.000000...1)$ , at which point the value of  $w_i$  is very close to 0. The probability of the value of  $w_i$  is very close to 0, and the effect of sparsity is achieved.

The variance of the spike and slab distribution is  $0.000...001$ , but its theoretical value is actually the limit of convergence to 0, so it is actually a Dirac delta function  $\delta(w_i)$  ( $y = 1$  except at  $x = 0, y = 0$  at other points). As shown in the figure 1.8.



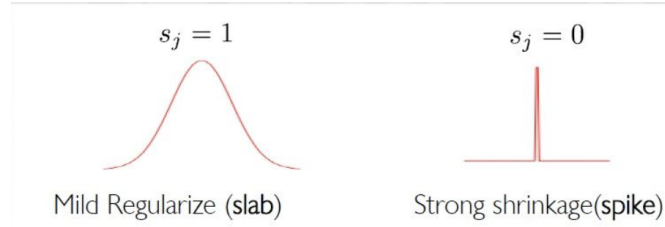


Figure 1.8: A sampling proecess of spike and slab.

The above steps can be abstracted into the formula:

$$P(w_j) = P(s_j)P(w_j|s_j) = \text{Bern}(s_j|p_0)(s_j N(w_j|0, \sigma_0^2) + (1 - s_j)\delta(w_j)) \quad (1.19)$$

where  $p_0, \sigma_0$  is all hyperparameters. Multiply this in front of your data-likelihood  $P(D|W)$  to get a joint distribution of probability models  $P(S)P(W|S)P(D|W)$ .

Compared to sparse methods such as L1 loss, spike and slab gives a more precise and fine-grained sparse modeling that we can place a unique "sparse prior  $p_0$ " for each parameter in the model, and also obtain a "sparse probability  $c(p_j)$ " after extrapolation that describes the probability of each parameter being selected. However, some of the most commonly used inference frameworks, such as variational inference and Expectation propogation, cannot be used directly under this kind of prior due to the form of the spike and slab prior (with discontinuous delta functions and mixed Gaussian forms).



## Chapter 2

# Implementation

### 2.1 The classic nested sampling implementation

In this section, i implement the classic nested sampling algorithm,

#### 2.1.1 Implement the sampler

#### 2.1.2 Sqlite database

### 2.2 Flask webserver

### 2.3 Query from the server

### 2.4 Calculating the MSE

1. The basic functions:  $\cos(x), \sin(x), \ln(x)$ , ( $\backslash\cos(x), \backslash\sin(x), \backslash\ln(x)$ ).
2. Greek letters:  $\alpha\beta\gamma\delta\epsilon\dots$  ( $\backslash\alpha\backslash\beta\backslash\gamma\backslash\delta\backslash\epsilon\dots$ ).
3. Mathematical symbols:  $\int \oint \sum \lim \bigcup \bigcap$  ( $\backslash\int\backslash\oint\backslash\sum\backslash\lim\backslash\bigcup\backslash\bigcap$ ).
4. Fractions:  $\frac{1}{2}, \frac{1}{2-x}$  ( $\backslash\frac{1}{2}, \backslash\frac{1}{2-x}$ ).

The following matrix

$$\begin{bmatrix} U_r & r & W_r \\ 0 & 1 & V_x \\ 0 & 0 & W_x \end{bmatrix}, \quad (2.1)$$

is generated using the `equarray` environment:

```

\begin{eqnarray}\label{eqn:matrix}
\left[
\begin{array}{ccc}
U_{\{r\}} & r & W_{\{r\}} \\
0 & 1 & V_{\{x\}} \\
0 & 0 & W_{\{x\}}
\end{array}
\right],
\end{eqnarray}

```

The `\label{eqn:matrix}` command labels the equation with `{eqn:matrix}` which can be referred to somewhere else in the text by using `\ref{eqn:matrix}` or `\eqref{eqn:matrix}`.

The command `\notag` eliminates the numbering of the first equation,

$$\begin{aligned}
 \lambda^{(1)} &= \text{tr}[T^{(1)}P], \\
 \lambda^{(2)} &= \text{tr}[T^{(2)}P - T^{(1)}ST^{(1)}P].
 \end{aligned}
 \tag{2.2}$$

```

\begin{eqnarray} \label{eqn:lambda_trace}
\lambda^{(1)} &=& \text{tr}[T^{(1)}P], \notag \\
\lambda^{(2)} &=& \text{tr}[T^{(2)}P - T^{(1)}ST^{(1)}P].
\end{eqnarray}

```

### 2.4.1 Itemized lists

Example of an itemized list:

- muscle and fat cells remove glucose from the blood,
- cells use glucose for protein synthesis.

```

\begin{itemize}
\item muscle and fat cells remove glucose from the blood,
\item cells use glucose for protein synthesis.
\end{itemize}

```

This can be done by an enumerated list:

1. muscle and fat cells remove glucose from the blood,
2. cells use glucose for protein synthesis.

```

\begin{enumerate}
\item muscle and fat cells remove glucose from the blood,
\item cells use glucose for protein synthesis.
\end{enumerate}

```

### 2.4.2 Inserting figures

You may save your Matlab figures as jpg files. Figures should be stored in the same folder as the latex files. For the graphicx package to work you usually need to ask latex to create a pdf file (e.g., command `pdflatex` or `latexpdf`).

An example of an inserted image is given in Figure 2.1.

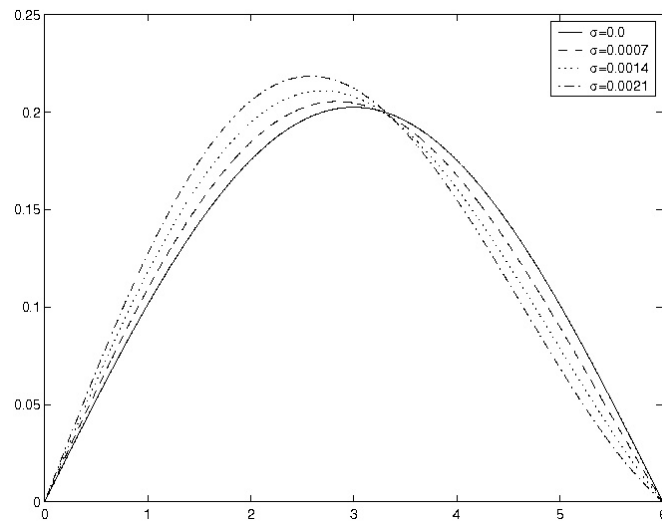


Figure 2.1: Mode shapes

### 2.4.3 Tables

Example of a table,

Gene	GeneID	Length
human latexin	1234	14.9 kbps
mouse latexin	2345	10.1 kbps
rat latexin	3456	9.6 kbps

Table 2.1: **title of table** - Overview of latexin genes.

```
\begin{table}[htp]
\centering
\begin{tabular}{ccc}
```

```
% ccc means 3 columns, all centered; alternatives are l, r
{\bf Gene} & {\bf GeneID} & {\bf Length} \\
\hline % draws a line under the column headers
human latexin & 1234 & 14.9 kbps \\
mouse latexin & 2345 & 10.1 kbps \\
rat latexin   & 3456 & 9.6 kbps \\
\end{tabular}
\caption[title of table]{\textbf{title of table} - Overview of latexin genes.}
\label{latexin_genes} % label for cross-links with \ref{latexin_genes}
\end{table}
```

See how to add two vertical lines in the table (Simply change {ccc} to {c|c|c})

Gene	GeneID	Length
human latexin	1234	14.9 kbps
mouse latexin	2345	10.1 kbps
rat latexin	3456	9.6 kbps

Table 2.2: **title of table** - Overview of latexin genes.

#### 2.4.4 How to Refer to Equations, Sections, etc

- References can be linked to equations, figures, tables or sections using the command `\ref`: Equation (2.2), Figure 2.1, Table 2.2 and Section 2.4.3.  
`Equation~(\ref{eqn:lambda_trace}), Figure~\ref{modes},`  
`Table~\ref{latexin_genes2} and Section~\ref{table}.`
- Equations can be conveniently referred to using `\eqref`. See, for example, Equation (2.2).  
`Equation \eqref{eqn:lambda_trace}`  
 Note that `\eqref` includes the round brackets by itself.
- Citations are in a similar way but using the command `\cite`:  
`[2], [3], and [4], or [2, 3, 4].`

```
\cite{Salmond}, \cite{Stull}, and \cite{TandC},
or \cite{Salmond,Stull,TandC} .
```

There are many different styles for writing citations – you should follow the norms for your subject area.

A more advanced way to do citations is to use `bibtex`. This is a powerful tool and we encourage you to try it. There is plenty of information about it on the web.





## **Chapter 3**

# **Methodologies and analysis**

### **3.1 Methodologies**

### **3.2 Analysis**



## **Chapter 4**

# **Discussion**

### **4.1 Main results**

### **4.2 Discussion**



## **Chapter 5**

## **Conclusions**

You may add more chapters as needed in the file.



# References

- [1] Farge Marie, *Wavelet Transforms and Their Applications to Turbulence*, Ann. Rev. Fluid Mech. volume 24, pages 395-457, 1992.
- [2] Salmond Jennifer, *Vertical Mixing of Ozone in the Very Stable Nocturnal Boundary Layer*, PhD Thesis, University of British Columbia, 2001.
- [3] Stull B. Ronald, *Introduction to Boundary Layer Meteorology*, Dordrecht; Boston: Kluwer Academic Publishers, 1988.
- [4] Torrence Christopher, Compo Gilbert P., *A Practical Guide to Wavelet Analysis*, Bulletin of the American Meteorological Society volume 79, pages 61-78, 1998.





## **Appendix A**

### **Some extra things**

This is an optional chapter for any additional material that does not fit conveniently into the body of the text (e.g., data, copies of computer programmes). Note that appendices won't necessarily be marked.