

CPSC532W Homework 4

Justin Reiher
Student ID: 37291151
CWL: reiher

Link to public repository for homework 4:

https://github.com/justinreiher/probProg_Fall2021/tree/main/CS532-HW4

1 Program 0: BBVI Implementation Details

Implementing algorithm 11 and 12 from the book are outlined as follows: For algorithm 11, the sample and observe functions, where a new variable v is introduced which is the address to the node in question. If the node does not exist then it is added to the program environment variable σ . The implementation of the `sample` and `observe` case are as follows:

```
1 elif e == 'sample*':
2     Dobj = args[0]
3     if not(v in sig['Q']):
4         sig['Q'][v] = Dobj.make_copy_with_grads()
5     dg = sig['Q'][v].make_copy_with_grads()
6     ret = dg.sample()
7     sig['G'][v] = grad_log_prob(dg, ret)
8     logWv = Dobj.log_prob(ret) - sig['Q'][v].log_prob(ret)
9     sig['logW'] = sig['logW'] + logWv.detach().nan_to_num()
10 elif e == 'observe*':
11     Dobj = args[0]
12     ret = args[1]
13     sig['logW'] = sig['logW'] + Dobj.log_prob(ret).detach().nan_to_num()
```

Like in the textbook, the `sample` case computes the `gradLogProb` of the sample from Q :

```
1 def grad_log_prob(Dobj, x):
2     logProb = Dobj.log_prob(x)
3     logProb.backward()
4     params = Dobj.Parameters()
5     g = []
6     for i in params:
7         g.append(i.grad)
8
9     return g
```

Since we only want the gradient, a copy of the distribution object `Dobj` is made to then compute the `gradLogProb` and then subsequently the result is `detached`. If we do not do this, then the gradients are compounding and create an exponentially growing gradient compute tree. The BBVI algorithm is implemented as follows:

```

1  #set R_0, by sampling every node from the prior and set the
   initial map of G and Q to empty,
2  #they get populated in deterministic_eval
3  for v in vertices:
4      linkFuncsEval = bindingVars(R,linkFuncs[v])
5      R[v] = deterministic_eval(linkFuncsEval,sig,v)
6
7  for v in sig['Q']:
8      sig['opt'][v] = [torch.optim.Adam(sig['Q'][v].Parameters(),
   lr),sig['Q'][v]]
9      sig['Qp'][v] = [torch.stack(sig['Q'][v].Parameters()).
   detach()]
10
11  for t in range(T):
12      gradients.append([])
13      ret.append([])
14      for s in range(num_samples):
15          sig['G'] = {}
16          sig['logW'] = 0
17          for v in vertices:
18              # mb = getMarkovBlanket(v)
19              # for x in mb:
20                  linkFuncsEval = bindingVars(R,linkFuncs[v])
21                  R[v] = deterministic_eval(linkFuncsEval,sig,v)
22          gradients[t].append(sig['G'].copy())
23          logWeights[t,s] = sig['logW'].detach()
24
25          if type(E) == str:
26              retExp = R[E]
27          else:
28              retExp = bindingVars(R,E)
29          ret[t].append(deterministic_eval(retExp,sig,[]))
30          ghat = elbo_grad(gradients[t],logWeights[t,:])
31          print(t)
32          print(logWeights[t,:].mean())
33          optimize(sig['Q'],ghat,sig['opt'],sig['Qp'],t)
34          # input('before after')
35
36  return ret,logWeights,sig['Q'],sig['Qp']

```

A topological graph sort is performed on the nodes before hand and stored. The nodes are then initially populated and for every random variable with an associated $Q(v)$ gets an optimization object initialized with the parameters of the prior associated with node v . The optimizer used in this case is `Adam`. This distribution object is used to make an update step, copy over the update to $Q(v)$ and continue. The optimization requires computing the weighted gradients. The gradient keys are merged together because under differing control flow it's possible that some gradients don't exist in some traces. In these instances the gradients are computed to zero:

```

1 def elbo_grad(G, logW):
2     #merging list of dictionaries:
3     # https://stackoverflow.com/questions/3494906/how-do-i-merge-a-
4     # list-of-dicts-into-a-single-dict
5     # this line is horribly unreadable, but:
6     # for every key 'k' in each dictionary 'd' of list 'G'
7     # insert its value 'v'.
8     # note that the values in this merged gradient object are
9     # irrelevant
10    # we just want to know all the keys that exists in all L
11    # gradient samples
12    # because different traces may have invoked different gradient
13    # addresses (or nodes)
14    Gmerg = {k: v for d in G for k, v in d.items()}
15    # return Gmerg
16    #initialize my F dictionary
17    #with the correct number of keys and empty lists
18    F = {k: [] for k in Gmerg}
19    Gs = {k: [] for k in Gmerg}
20    L = len(logW)
21    ghat = {}
22    for v in Gmerg:
23        sizeG = torch.stack(Gmerg[v]).shape
24        for s in range(L):
25            if v in G[s].keys():
26                F[v].append(torch.stack(G[s][v])*logW[s])
27            else:
28                F[v].append(torch.zeros(sizeG))
29                G[s][v].append(torch.zeros(sizeG))
30        #this picks out all gradients from 1:L of G in a list for
31        #variable v
32        GL = torch.stack([torch.stack(g[v]) for g in G])
33        Fv = torch.stack(F[v])
34        bhat = computeBaseline(Fv, GL)
35        ghat[v] = torch.sum((Fv - bhat*GL)/L, 0)
36    return ghat

```

and subtracting the *baseline*. The *baseline* is computed by computing the element-wise covariance of each mini-batch trace and the weighted gradients and then summed. Any results resulting in *nan* are turned to zeros using `torch.nan_to_num`. Using `numpy.cov` routine returns a `dtype=float64` result, this is truncated back to `torch.float32` (This baseline computation was discussed at length with many of my classmates)

```

1 def computeBaseline(Fv, GL):
2     C = []
3     #Check to see if the elements of Fv are multi-dimensional
4     if len(Fv[0].shape) > 1:
5         n, d = Fv[0].shape
6         for i in range(n):
7             C_i = []
8             #collect the cov(Fi, Gi) terms
9             for j in range(d):
10                C_ij = np.cov(Fv.detach().numpy()[i, :], GL.numpy())

```

```

11    [:,i,j], rowvar = True)
12         C_i.append(C_ij[0,1])
13     C.append(C_i)
14     Cov = torch.tensor(C)
15     #need to get ride of NaN, sum all the cov(Fi,Gi) terms
16     bhat = torch.nan_to_num(Cov.sum()/GL.var(0).sum())
17 else:
18     for i in range(Fv[0].shape[0]):
19         C_i = np.cov(Fv.detach().numpy()[ :,i],GL.numpy()[ :,i],
20 rowvar = True)
21         C.append(C_i[0,1])
22         Cov = torch.tensor(C)
23         #again need to get ride of NaN
24         bhat = torch.nan_to_num(Cov.sum()/GL.var(0))
25
26 # for some reason numpy turns the above into float64,
27 everything else (and default torch floating point numbers) is
28 in torch.float32
29 # so truncate back to torch.float32
30 return torch.tensor(bhat, dtype = torch.float32)

```

The optimization step retrieves the optimization object associated with the $\mathcal{Q}(v)$ that is being optimized, a step is taken the gradients are then set to zero (same reason as to why the log-weights are detached after being computed) and the distribution object in $\mathcal{Q}(v)$ is updated with the new parameters:

```

1 def optimize(Q,ghat,optimizers,Qp,t):
2     for v in ghat:
3         [opt,Dobj] = optimizers[v]
4         grad = ghat[v]
5         i = 0
6         for p in Dobj.Parameters():
7             p.grad = -torch.nan_to_num(grad[i].detach())
8             i+=1
9         opt.step()
10        opt.zero_grad()
11
12        i = 0
13        for p in Q[v].Parameters():
14            p.data = Dobj.Parameters()[i] #+ 1e-8/(t+1)*grad[i]
15            i += 1
16        Qp[v].append(torch.stack(Q[v].Parameters()).detach())

```

Finally, the results of the samples, the log-weights are stored and then returned.

1.1 Program 1

Number of samples per optimization is: 200, number of optimization steps is: 350, Adam learning rate: 0.5

Collect samples denoted by program 1:

Elapsed time for program 1 .daphne is: 0:01:28.018433 seconds

Mean of set of samples: tensor(7.2636)

and variance of samples: tensor(0.0539)

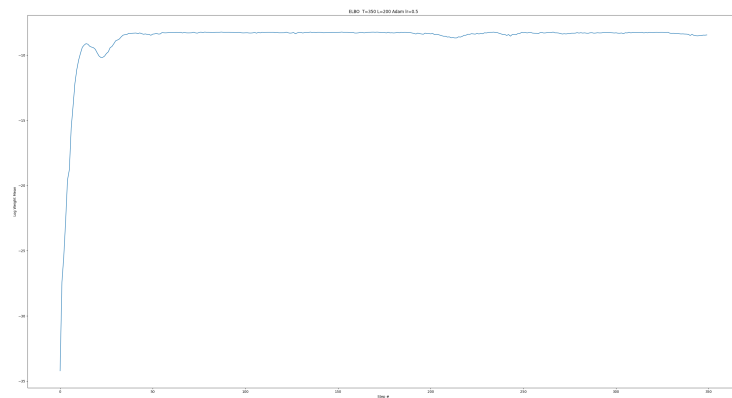


Figure 1: ELBO plot

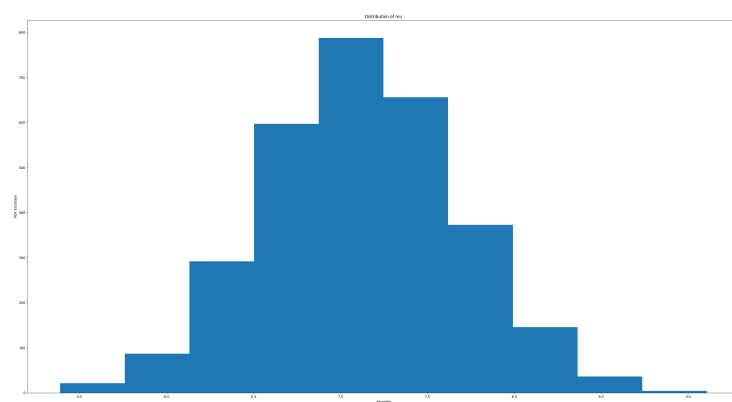


Figure 2: Estimate of Distribution μ from \mathcal{Q}

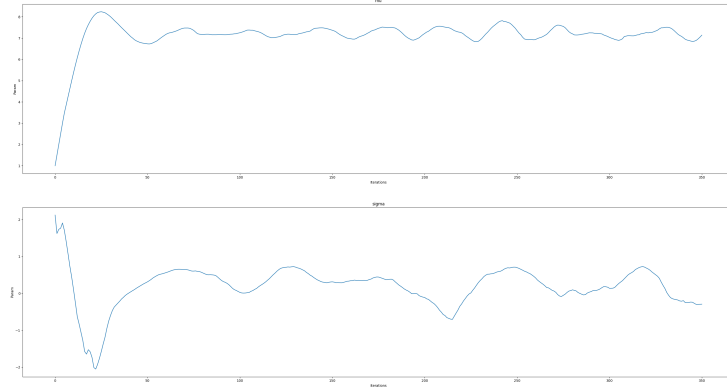


Figure 3: Parameters for Q for μ and Σ over iterations

1.2 Program 2

Number of samples per optimization is: 200, number of optimization steps is: 350, Adam learning rate: 0.25

Collect samples denoted by program 2:

Elapsed time for program 2 .daphne is: 0:03:02.441880 seconds

Mean of set of samples: `tensor([1.8603, 0.6375])`

and variance of samples: `tensor([0.0843, 1.3375])`

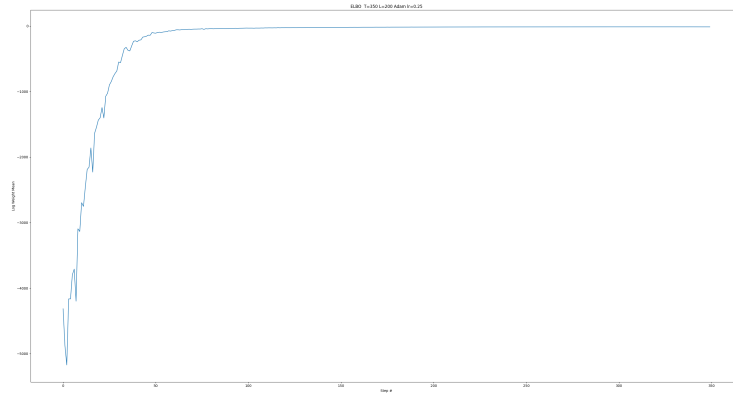


Figure 4: ELBO plot

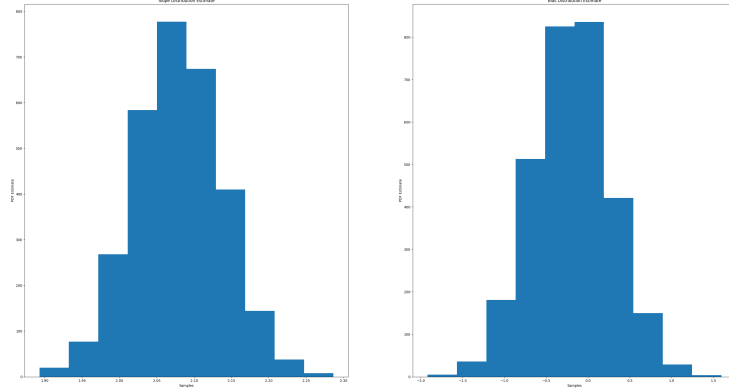


Figure 5: Estimate of Distributions for Slope and Bias from Q respectively

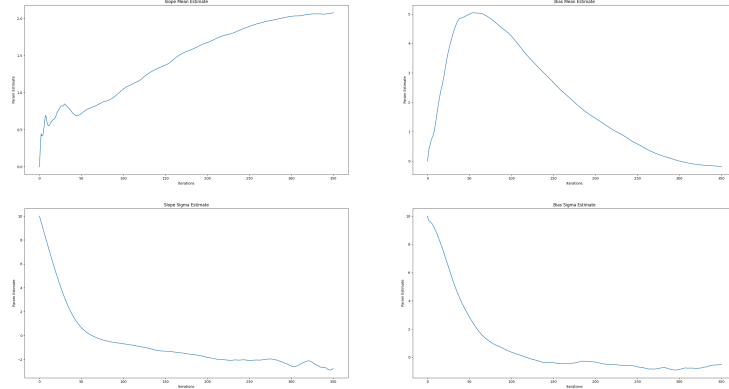


Figure 6: Parameters for Q for Slope and Bias over iterations

1.3 Program 3

Number of samples per optimization is: 25, number of optimization steps is: 350, Adam learning rate: 0.05

Collect samples denoted by program 3:

Elapsed time for program 3 .daphne is: 0:01:49.841174 seconds

Mean of samples: tensor(0.5729)

and variance of samples: tensor(0.2132)

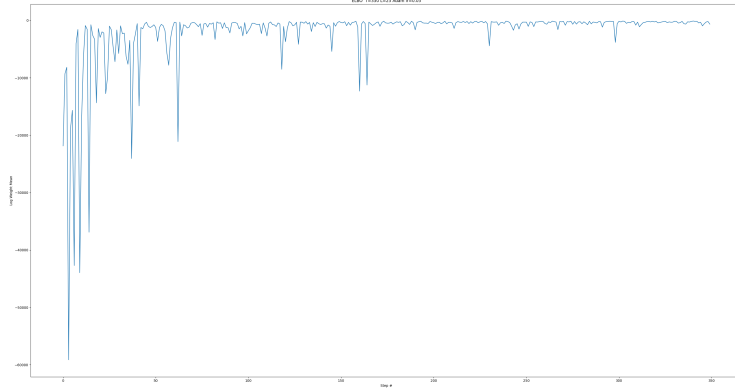


Figure 7: ELBO plot

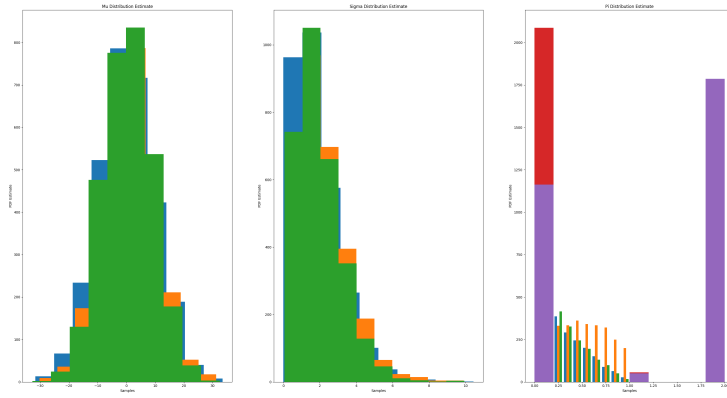


Figure 8: Estimates of Distributions for the 3 Modes (μ and Σ) and Π

During optimization there are sudden jumps where the ELBO drops significantly likely arising from the fact that the modes switch, i.e. sampling from one mode for awhile and then sampling from another. In these instances a sudden drop in the importance weights will arise from estimating the distribution from the "wrong" place. The convergence back to the previous ELBO is quick, however this bouncing around makes optimization hard.

1.4 Program 4

Number of samples per optimization is: 150, number of optimization steps is: 200, Adam learning rate: 0.1


```

Collect samples denoted by program 4:
Elapsed time for program 4 .daphne is: 0:32:30.898768 seconds
Mean of samples W0 : tensor([[ -0.0538],
[ 0.0759],
[ -0.0523],
[ -0.4615],
[ -0.0378],
[ -0.0117],
[ -0.0077],
[ -0.0430],
[ 0.2541],
[ -0.2181]])
and variance of samples W0 : tensor([[0.5128],
[0.5521],
[0.2950],
[0.4300],
[0.4077],
[0.2621],
[0.6241],
[0.5681],
[0.4060],
[0.3981]])
Mean of samples b0 : tensor([[ 0.0283],
[ -0.1436],
[ 0.0596],
[ -0.1415],
[ -0.0643],
[ -0.1060],
[ 0.0333],
[ -0.3151],
[ 0.2745],
[ 0.2415]])
and variance of samples b0 : tensor([[0.3771],
[0.3458],
[0.4271],
[0.3895],
[0.4588],
[0.3073],
[0.4415],
[0.2625],
[0.4780],
[0.2059]])
Mean of samples W1 : tensor([[ -0.3377, 0.1622, 0.2904, 0.3412, -0.0154, -0.2529, 0.1700,
-0.1905, 0.1341],
[ 0.6109, -0.0588, 0.1595, -0.2266, -0.2677, -0.1859, 0.2364, 0.0217,
-0.0459, 0.0779],

```

```

[-0.0839, -0.2009, -0.3588, -0.1991,  0.1542, -0.0972,  0.2010, -0.1792,
-0.1038, -0.0384],
[ 0.0038,  0.0847, -0.0733, -0.2610,  0.0972,  0.1078, -0.0553,  0.2599,
-0.1014, -0.2961],
[ 0.0242,  0.0829, -0.0083,  0.2522,  0.1779,  0.1928, -0.2569, -0.4040,
-0.1642,  0.0741],
[ 0.1149, -0.2028, -0.1752, -0.1942, -0.3036,  0.5337,  0.0388,  0.1312,
-0.0115, -0.0357],
[ 0.4245,  0.3058, -0.2490,  0.0654,  0.1084, -0.0598,  0.3495,  0.0311,
-0.2384,  0.5183],
[-0.1928, -0.1891, -0.2955, -0.3391,  0.2615, -0.1450, -0.2682, -0.1339,
-0.2855,  0.1191],
[ 0.2049, -0.0804, -0.1176, -0.0216, -0.0917, -0.0542,  0.1835,  0.0915,
-0.1915, -0.0375],
[-0.0047,  0.1943, -0.1376, -0.2776,  0.0288,  0.1535,  0.0938, -0.0937,
-0.1021, -0.0475]])
and variance of samples W1 : tensor([[0.3923, 0.4234, 0.4946, 0.2481, 0.4493, 0.2450, 0.27
0.3278],
[0.4420, 0.3859, 0.4415, 0.4125, 0.3175, 0.3814, 0.3217, 0.4444, 0.3659,
0.3045],
[0.3325, 0.4457, 0.4203, 0.6208, 0.4533, 0.2631, 0.4323, 0.5770, 0.3698,
0.5566],
[0.3912, 0.3266, 0.3773, 0.3905, 0.5574, 0.6455, 0.5060, 0.3169, 0.4468,
0.4718],
[0.4554, 0.6540, 0.2060, 0.3671, 0.4822, 0.3046, 0.2851, 0.4852, 0.4505,
0.6185],
[0.5007, 0.4577, 0.4148, 0.3821, 0.4859, 0.3172, 0.4985, 0.4503, 0.2416,
0.5182],
[0.4381, 0.5384, 0.4589, 0.3488, 0.5986, 0.4541, 0.4134, 0.3126, 0.2671,
0.3327],
[0.5090, 0.4212, 0.4048, 0.5165, 0.2403, 0.1082, 0.6374, 0.3966, 0.2565,
0.5461],
[0.3849, 0.4273, 0.3088, 0.5978, 0.3867, 0.5247, 0.4416, 0.3489, 0.5252,
0.4045],
[0.3821, 0.3247, 0.4055, 0.5226, 0.4852, 0.3786, 0.3103, 0.4447, 0.6368,
0.3510]])
Mean of samples b1 : tensor([[ 0.0669],
[ 0.2564],
[-0.0540],
[-0.3482],
[ 0.1357],
[-0.1493],
[ 0.3596],
[-0.0459],
[-0.5084],
[-0.0625]])

```

```

and variance of samples  b1 :  tensor([[0.3432],
[0.3910],
[0.5833],
[0.2678],
[0.3929],
[0.6498],
[0.4430],
[0.4306],
[0.3789],
[0.3163]])

```

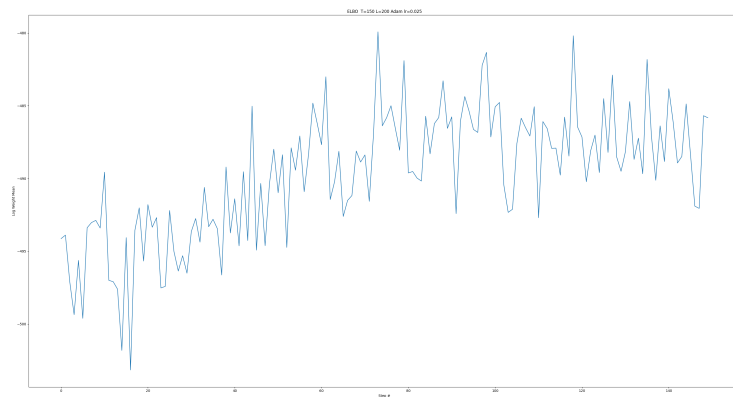


Figure 9: ELBO plot



Figure 10: Heatmap of expectation of W_0



Figure 11: Heatmap of expectation of b_0

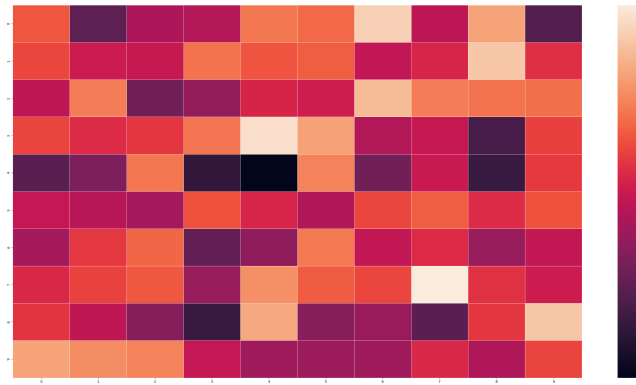


Figure 12: Heatmap of expectation of W_1



Figure 13: Heatmap of expectation of b_1

1.5 Program 5

For this program I am going to get creative about approximating the *Uniform* distribution using tanh functions. I have used Wolfram Alpha to check some concrete examples, but a derivation to my approach:

$$\begin{aligned} f_+(x) &= 0.5 \tanh(x) + 0.5 \in [0, 1] \\ \lim_{x \rightarrow \infty} f_+(x) &= 1 \\ \lim_{x \rightarrow -\infty} f_+(x) &= 0 \end{aligned} \quad (1)$$

If we define $f_-(x) = -0.5 \tanh(x) + 0.5$, then $x \rightarrow \infty f_-(x) = 0$ and $x \rightarrow -\infty f_-(x) = 1$. We can scale how fast the transition occurs with a parameter $s > 0$ and translate where the transition occurs with a :

$$f(x) = 0.5 \tanh(s(x - a)) + 0.5 \quad (2)$$

To approximate `Uniform(0,1)` we compose f_+ and f_- as follows:

$$\begin{aligned} \text{Uniform}(x)[0, 1] &\approx 0.5 (\tanh(s(x)) - \tanh(s(x - 1))) \\ \hat{\mathcal{U}}(x)[a, b] &= \frac{1}{b-a} (0.5 (\tanh(s(x + a)) - \tanh(s(x - b)))) \end{aligned} \quad (3)$$

The question is the above a real probability density function? Using Wolfram Alpha and as an example $s = 10000$, $b = 1$ and $a = -1$ then $\int_{-\infty}^{\infty} \hat{\mathcal{U}}(x)[a, b] dx = 1$. Why use tanh functions? They are differentiable everywhere, as $s \rightarrow \infty$ the approximation gets closer and closer to the real distribution.

Collect samples denoted by program 5:

```
Elapsed time for program 5 .daphne is: 0:00:11.613676 seconds
tensor([2.9748, 4.0338])
```

```

tensor([[5.3001, 1.2459],
[5.3001, 1.2459],
[5.3001, 1.2459],
...,
[2.9748, 4.0338],
[2.9748, 4.0338],
[2.9748, 4.0338]])
Mean of trace: tensor([2.9516, 4.0578])
and variance of trace: tensor([0.7594, 0.7662])

```

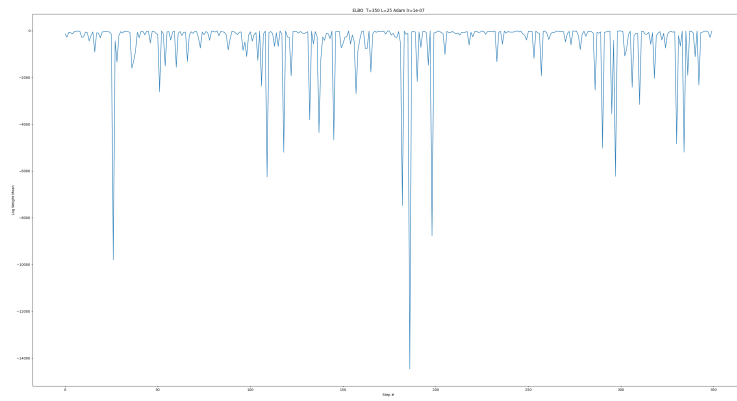


Figure 14: ELBO plot

Figure 15: Distribution Estimate for s from \mathcal{Q}