The Classical Complexity of Boson Sampling: A multi-core CPU implementation

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Level I 40cp project

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Computing the permanent of large matrices is one of the key parts of the Boson Sampling problem, as will be discussed in detail while explaining the problem in the subsequent sections of the paper.

3.2.1 Definition

Definition 3.1. /?/ The permanent of an $n \times n$ matrix $A = (a_{ij})$ is defined as

$$\operatorname{Per} A = \sum_{\sigma \in S_n} \prod_{i=1}^n a_{i\sigma(i)} \tag{1}$$

where the sum is over all elements of the symmetric group S_n i.e. over all permutations of the numbers in $[n] = \{1, 2, ..., n\}$

Example 3.2. For a 2×2 matrix, the permanent is calculated as follows

$$\operatorname{Per} \begin{bmatrix} a & b \\ c & d \end{bmatrix} = ad + bc \tag{2}$$

Example 3.3. For a 3×3 matrix, the permanent is calculated as follows

$$\operatorname{Per} \begin{bmatrix} a & b & c \\ d & e & f \\ g & h & i \end{bmatrix} = aei + bfg + cdh + ceg + bdi + afh \tag{3}$$

One can observe that the definition of the permanent is similar to the more commonly used determinant function, differing in the fact that the permanent definition lacks the alternating signs. An important property to note about the permanent function is that it is invariant to transposition i.e. Per $A = \text{Per } A^T$ [?].

3.2.2 Computing the permanent

Valiant showed that the problem of computing the permanent of a matrix is in the class #P-complete which implies that it is unlikely to have a polynomial time algorithm which implies that it is unlikely to have a polynomial-time algorithm [?]. The naive algorithm obtained by directly translating the formula into an algorithm would run in $\mathcal{O}(n!n)$ time.

A significant improvement on the naive approach, Rysers algorithm uses a variant of the inclusion-exclusion principle and can be evaluated in $\mathcal{O}(n^2 2^n)$ time [?]. Nijenhuis and Wilf sped this up to $\mathcal{O}(n2^n)$ time by iterating over the sum in Gray Code order [?].

Another formula that is as fast as Rysers was independently derived by Balasubramanian[?], Bax[?], Franklin and Bax[?], and Glynn[?], all using different methods. We shall henceforth refer to this as Glynn's formula and it is described as follows.

Let $M = (m_{ij})$ be an $n \times n$ matrix with $m_{ij} \in \mathbb{C}$, then

$$\operatorname{Per} M = \frac{1}{2^{n-1}} \sum_{\delta} \left(\prod_{k=1}^{n} \delta_k \right) \prod_{j=1}^{n} \sum_{i=1}^{n} \delta_i m_{ij}$$
 (4)

where $\delta \in \{-1,1\}^n$ with $\delta_1 = 1$. Hence there are 2^{n-1} such values for δ .

Implementing the formula as is would require $\mathcal{O}(2^n n^2)$ time. However, iterating over the δ arrays in Gray code order reduces it to $\mathcal{O}(n2^n)$ time.

In order to exploit this trick, let v_j be the symbol used to denote the innermost sum i.e. $v_j = \sum_{i=1}^n \delta_i m_{ij}$. If δ is iterated through in Gray code order, then one can notice that the terms $\{v_j, j \in [n]\}$ can actually be computed in $\mathcal{O}(n)$ time for a given value of δ rather than $\mathcal{O}(n^2)$ which is how long it would take if done naively. This is because successive elements of δ differ in only one position, so each successive v_j differs only by the addition or subtraction of some m_{ij} where i is the position at which δ was changed. Hence, the product of v_j terms can now be calculated in $\mathcal{O}(n)$ time.

4 Describing the paper and specifying the problem

Cliiford and Clifford [?] gave a study and analysis of the classical complexity of the exact Boson Sampling problem and proposed an algorithm that is significantly faster than previous algorithms for the problem. The algorithm is simple to implement, and is able to solve the Boson Sampling problem for system sizes much greater than quantum computing systems currently available, which reduces the likelihood of achieving quantum supremacy in the context of Boson Sampling in the near future rephrase last bit.

4.1 Explaining the problem

A summary of the problem in purely mathematical terms is as follows.

Let m and n be positive integers. Consider all possible multisets¹ of size n with elements in [m], where $[m] = \{1, ..., m\}$. Let $z = [z_1, z_2, ..., z_n]$ be an array representation of such a multiset, with its elements in non-decreasing order. In other words, z is an array of n integers taken from [m] (with repetition) and arranged in non-decreasing order. Define $\Phi_{m,n}$ to be the set of all distinct values that z can take. Define $\mu(z) = \prod_{j=1}^m s_j!$ where s_j is the multiplicity of j in the array z i.e. the number of times it appears in z.

¹A multiset is a special kind of set in which elements can be repeated

 $A = (a_{ij})$ is a complex-valued $m \times n$ matrix constructed by taking the first n columns of a given $m \times m$ Haar random unitary matrix. For each z, build an $n \times n$ matrix A_z where the k^{th} row of A_z is the z_k^{th} row in A, for k = 1, ..., n. Finally, define a probability mass function over $\Phi_{m,n}$ as

$$q(z) = \frac{1}{\mu(z)} |\text{Per } A_z|^2 = \frac{1}{\mu(z)} \left| \sum_{\sigma} \prod_{k=1}^n a_{z_k \sigma_k} \right|^2, \quad z \in \Phi_{m,n}$$
 (5)

where Per A_z is the permanent of A_z and $\pi[n]$ is the set of all permutations of [n]. The computing task is to simulate random samples from the above pmf q(z).

4.1.1 Calculating the size of the sample space

The size of the sample space $\Phi_{m,n}$ can be calculated using the 'stars and bars' technique from combinatorics [?]. In our problem, we have n 'stars' representing the elements of the array z, and m 'buckets' representing all the values in [m]. Recall that z is a sorted array representation of a multiset, so multiple 'stars' can be in one 'bucket'. Since there are m 'buckets', we need m-1 'bars' to divide the 'stars' into 'buckets'. Therefore, from a total of m-1+n objects, we need to pick n of these to be the 'stars'. There are $\binom{m+n-1}{n}$ ways to do this. Hence, there are $\binom{m+n-1}{n}$ possible values of z.

5 The Boson Sampling Algorithm

5.1 The naive approach

Translating the formula above $(\ref{eq:condition})$ directly to an algorithm would require $\mathcal{O}(\binom{m+n-1}{n}n2^n)$ time to evaluate. The $\binom{m+n-1}{n}$ term comes from the size of the sample space of the pmf, and calculating the value of the permanent using the fastest known methods takes $\mathcal{O}(n2^n)$ time (Section $\ref{eq:condition}$). With $m = \mathcal{O}(n^2)$ as suggested in Section $\ref{eq:condition}$, the total running time is $\mathcal{O}(\binom{n^2+n-1}{n}n2^n) = \ref{eq:condition}$. Hence, even for relatively small values of n ((give some actual numbers), computing the Boson Sampling problem would be intractable for even very powerful supercomputers.

In the literature [?], two new algorithms are proposed for exact Boson Sampling. These are referred to as Algorithm A and Algorithm B, and both provide a significant speed up on the naive algorithm. The following subsections summarise the approach taken to obtain them.

5.2 Algorithm A

The approach to Algorithm A starts by expanding the sample space to a much larger size, which seems counterintuitive at first, but it allows us to express the pmf (Equation ??) in a form that is much easier to compute. The sample space is expanded to the space of all arrays $\mathbf{r} = (r_1, r_2, ..., r_n)$ where each element r_k is in [m], which implies that

we are considering a distribution on the product space $[m]^n$. It is stated and proved that sampling from $q(\mathbf{z})$ is equivalent to sampling from the pmf

$$p(\mathbf{r}) = \frac{1}{n!} \left| \operatorname{Per} A_{\mathbf{r}} \right|^2 = \frac{1}{n!} \left| \sum_{\sigma} \prod_{i=1}^{n} a_{r_i \sigma_i} \right|^2, \quad \mathbf{r} \in [m]^n$$
 (6)

where as before, σ is the set of all permutations of [n].

This method requires $p(\mathbf{r}) = p(r_1, ..., r_n)$ to be rewritten as a product of conditional probabilities using the chain rule i.e.

$$p(\mathbf{r}) = p(r_1)p(r_2|r_1)p(r_3|r_1, r_2)...p(r_n|r_1, r_2, ..., r_{n-1})$$
(7)

We first sample r_1 from $p(r_1), r_1 \in [m]$. Then for k = 2, ...n, we sample r_k from the conditional pmf $p(r_k|r_1, r_2, ..., r_{k-1})$ with $r_1, ..., r_{k-1}$ fixed. After sampling all values of r_k , sort $(r_1, r_2, ..., r_n)$ in non-decreasing order, and that results in the array representation of a multiset sampled from the Boson Sampling distribution $q(\mathbf{z})$. In order to obtain the conditional probabilities, a formula for the joint pmf of the leading subsequences of $(r_1, ..., r_k)$ is given in Lemma 1 of the paper, and proved using arithmetic techniques and facts about probability measures. The formula in Lemma 1 is as follows:

$$p(r_1, ..., r_k) = \frac{(n-k)!}{n!} \sum_{c \in \mathcal{C}_k} \left| \operatorname{Per} A_{r_1, ..., r_k}^c \right|^2, \quad k = 1, ..., n$$
 (8)

where C_k is the set of k-combinations taken without replacement from [n] and A_{r_1,\ldots,r_k}^c is the matrix formed by taking only columns $c \in \mathcal{C}_k$ of the rows $(r_1, ..., r_k)$ of A.

Notice in equation ??, we need to sample r_k from the conditional probability distribution $p(r_k|r_1,...,r_{k-1})$. This can be rewritten as $p(r_1,...,r_k)/p(r_1...r_{k-1})$. Since r_k does not appear in the denominator, in order to sample r_k from the conditional pmf, we can equivalently sample from the pmf proportional to the numerator, as $(r_1,...r_{k-1})$ are fixed, known values at this stage in the algorithm. Therefore, the formula is Lemma 1 is used to calculate the conditional pmfs at each stage, giving way to the following algorithm.

Algorithm 1 Boson Sampler: Single sample **z** from $q(\mathbf{z})$ in $\mathcal{O}(mn3^n)$ time

Require: $m, n \in \mathbb{Z}_+$; A formed by first n columns of $m \times m$ Haar random unitary matrix

- 1: $\mathbf{r} \leftarrow \emptyset$
- 2: for $k \leftarrow 1 \rightarrow n$ do
- $w_i \leftarrow \sum_{c \in \mathcal{C}_k} \left| \operatorname{Per} A^c_{(\mathbf{r},i)} \right|^2, i \in [m]$ $x \leftarrow \operatorname{Sample}(w)$
- $\mathbf{r} \leftarrow (\mathbf{r}, x)$
- 6: $\mathbf{z} \leftarrow \operatorname{IncSort}(\mathbf{r})$
- 7: return z

The correctness of the algorithm is clear as it is simply an evaluation of the required pmf using the chain rule of probability. The literature gives a mathematical proof to derive the runtime, however here we shall give an informal explanation to obtain the runtime using the algorithm above. The runtime is dominated by the **for** loop in lines 2 to 6, as the sorting step in line 7 takes only $\mathcal{O}(n \log n)$ using the fastest methods for sorting (insert reference). In each iteration of the **for** loop, we first construct the conditional pmf on the sample space [m]. This is represented by a weighted array **w**. This involves calculating the permanents of $|\mathcal{C}_k| \ k \times k$ matrices for each $i \in [m]$, where k is the loop index variable. Sampling from this distribution (line 4) takes $\mathcal{O}(m)$ time [?], but this too is dominated by the calculations in line 3, so we can ignore it. Therefore, the total time taken in the kth iteration is $\mathcal{O}(m\binom{n}{k}k2^k)$, as $|\mathcal{C}_k| = \binom{n}{k}$ and calculating the permanent of a $k \times k$ matrix takes $\mathcal{O}(k2^k)$ using the fastest methods. So for n iterations,

$$\sum_{k=1}^{n} mk2^{k} \binom{n}{k} = m\frac{2}{3}n3^{n} = \mathcal{O}(mn3^{n})$$
 (9)

Therefore the total time taken is $\mathcal{O}(mn3^n)$. In terms of space complexity, we only need to store one permanent calculation at a time, and the weight array for each pmf is of size m. These can also be stored only one at a time, so $\mathcal{O}(m)$ additional space is required.

5.3 Algorithm B

The main theorem of the paper (Theorem 1) states that the time complexity of the Boson Sampling problem is $\mathcal{O}(n2^n + \text{poly}(m, n))$ where $\text{poly}(m, n) = \mathcal{O}(mn^2)$, with $\mathcal{O}(m)$ additional space required. This is achieved using the second algorithm proposed in the paper i.e. Algorithm B.

Algorithm B makes use of the Laplace Expansion [?] of permanents to obtain a significant speed up. The Laplace expansion allows us to calculate the permanent of a matrix using its permanent minors (the permanent of a submatrix with a column and a row removed). For any $k \times k$ matrix $B = (b_{i,j})$,

$$\operatorname{Per} B = \sum_{l=1}^{k} b_{k,l} \operatorname{Per} B_{k,l}^{\diamond}, \tag{10}$$

where $B_{k,l}^{\diamond}$ is the submatrix of B with row k and column l removed. Hence the permanent of B can be calculated in $\mathcal{O}(k)$ steps provided the permanent minors i.e. the values $\{\operatorname{Per} B_{k,l}^{\diamond}\}$ are known.

Computing $\{\text{Per }B_{k,l}^{\diamond}\}$ by calculating each of the permanents independently would collectively take $\mathcal{O}(k^22^k)$ time as there are k permanents to compute and to compute the value of each one takes $\mathcal{O}(k2^k)$ time. Lemma 2 of the paper states that the collection $\{\text{Per }B_{k,l}^{\diamond}\}$ can be collectively evaluated in $\mathcal{O}(k2^k)$ time and $\mathcal{O}(k)$ extra space. using Glynn's formula to evaluate a permanent minor for a given value of l, we have

$$\operatorname{Per} B_{k,l}^{\diamond} = \frac{1}{2^{k-2}} \sum_{\delta} \left(\prod_{i=1}^{k-1} \delta_i \right) \prod_{j \in [k] \setminus l} v_j(\delta), \quad l \in [k],$$
(11)

where $\delta \in \{-1,1\}^{k-1}$ with $\delta_1 = 1$ and $v_j(\delta) = \sum_{i=1}^{k-1} \delta_i b_{ij}$. The usual trick of iterating over the values of δ in Gray code order is required to compute $\operatorname{Per} B_{k,l}^{\diamond}$ in $\mathcal{O}(k2^k)$ time for each value of l, but this would still have to be repeated k times to cover all $l \in [k]$. The second and more novel trick used to achieve an additional speed up is used while calculating the products $\prod_{j \in [k] \setminus l} v_j(\delta)$. Observe that for each l, $\prod_{j \in [k] \setminus l} v_j(\delta) = \prod_{j \in [k]} v_j(\delta)/v_l(\delta)$. However, we cannot compute this method to compute the partial product as it would not work for cases when $v_l(\delta) = 0$. Instead, we first precompute two arrays \mathbf{f} and \mathbf{b} containing the forwards and backwards cumulative products of $v_j(\delta)$ respectively as follows,

$$f_i = \prod_{j=1}^i v_j(\delta) \quad b_i = \prod_{j=i}^k v_j(\delta), \quad i \in [k].$$

$$(12)$$

Then each partial product $\prod_{j\in[k]\setminus l} v_j(\delta)$ can be expressed as a product of two terms from \mathbf{f} and \mathbf{b} , $f_{l-1}b_{l+1}$ for any l. Since it takes $\mathcal{O}(k)$ time to compute the forward and backward cumulative product arrays and $\mathcal{O}(1)$ time to obtain each partial product $\prod_{j\in[k]\setminus l} v_j(\delta)$, for each value of δ , the total time taken to simultaneously calculate the values of $\{\operatorname{Per} B_{k,l}^{\diamond}\}$ is $\mathcal{O}(k2^k)$. Additional space required to store the partial products is $\mathcal{O}(k)$.

Coming back to formulating the second Boson Sampling algorithm, the sample space is now expanded even further with an auxiliary array $\alpha = (\alpha_1, \alpha_2, ..., \alpha_n)$ where $\alpha \in \pi(n)$, which is the set of permutation of [n]. The approach is similar to that of Algorithm A in that the goal is to create a succession of pmfs for leading subsequences of \mathbf{r} and then progressively sampling r_i from conditional pmfs on $(r_1, ..., r_{i-1})$. The conditioning variable α helps us reformulate the pmf in such a way that we can use the permanent minors that we are able to calculate efficiently, while still ensuring that it is equivalent to sampling from the pmf $p(\mathbf{r})$ in equation ??. Define

$$\phi(r_1, ... r_k | \alpha) = \frac{1}{k!} \left| \text{Per } A_{r_1, ... r_k}^{[n] \setminus \{\alpha_{k+1}, ..., \alpha_n\}} \right|^2, \quad k = 1, ..., n - 1.$$
 (13)

In Lemma 3 of the paper, it is shown that $p(\mathbf{r}) = \mathbb{E}_{\alpha} \{ \phi(\mathbf{r}|\alpha) \}$, which is the expectation taken over α , uniformly distributed over $\pi(n)$ for fixed \mathbf{r} . The chain rule of expectation is used to prove the equality, and this full proof is provided in the literature. As in Algorithm A, a chain of conditional pmfs as follows is used,

$$\phi(\mathbf{r}|\alpha) = \phi(r_1|\alpha)\phi(r_2|r_1,\alpha)\phi(r_3|r_1,r_2,\alpha)...\phi(r_n|r_1,...r_{n-1},\alpha),\tag{14}$$

where $\phi(r_k|r_1,...r_{k-1},\alpha) = \phi(r_1,r_2,...r_k|\alpha)/\sum_{r_k}\phi(r_1,...r_k|\alpha)$, using conditional probabilities and the law of total probability. The algorithm starts off by sampling r_1 from the pmf $\phi(r_1|\alpha)$. Then for stages k=2,...,n, r_k is sampled from $\phi(r_k|r_1,...r_{k-1},\alpha)$ with $(r_1,...,r_{k-1})$ fixed. Since the terms in the denominator are fixed, known values, sampling from $\phi(r_k|r_1,...r_{k-1})$ is equivalent to sampling from the numerator, which can be evaluated by taking advantage of the Laplace Expansion of permanents. At stage n, the array $(r_1,...,r_n)$ will have been sampled from $p(\mathbf{r})$. Sorting $(r_1,...,r_n)$ in non-decreasing

Algorithm 2 Boson Sampler: Single sample \mathbf{z} from $q(\mathbf{z})$ in $\mathcal{O}(n2^n + \text{poly}(m, n))$ time Require: $m, n \in \mathbb{Z}_+$; A formed by first n columns of $m \times m$ Haar random unitary matrix

```
1: \mathbf{r} \leftarrow \emptyset
                                                                                                                                   ▶ Empty array
 2: A \leftarrow \text{Permute}(A)
                                                                                             \triangleright Randomly permute columns of A
 3: w_i \leftarrow |a_{i,1}|^2, i \in [m]
                                                                                                \triangleright Make indexed weighted array w
 4: x \leftarrow \text{Sample}(w)
                                                                                                               \triangleright Sample index x from w
 5: \mathbf{r} \leftarrow (\mathbf{r}, x)
                                                                                                                               \triangleright Append x to \mathbf{r}
 6: for k \leftarrow 2 to n do
            B_{k}^{\diamond} \leftarrow A_{\mathbf{r}}^{[k]}
Compute {Per B_{k,l}^{\diamond}, l \in [k]}
                                                                                                                              ⊳ From Lemma 2
            w_i \leftarrow \left| \sum_{l=1}^k \operatorname{Per} B_{k,l}^{\diamond} \right|^2, i \in [m]
                                                                                                            10:
            \mathbf{r} \leftarrow (\mathbf{r}, x)
12: \mathbf{z} \leftarrow \operatorname{IncSort}(\mathbf{r})
                                                                                                   \triangleright Sort r in non-decreasing order
13: return z
```

order gives \mathbf{z} , the array representation of the multiset sampled from the Boson Sampling distribution $q(\mathbf{z})$.

Correctness holds as Algorithm B samples from successive conditional pmfs, which is equivalent to sampling from $\phi(r|\alpha)$ due to the chain rule of probability (Equation ??), for a given α . Since α is uniformly distributed over the sample space $\pi(n)$, the algorithm equivalently samples from $\mathbb{E}_{\alpha}\{\phi(\mathbf{r}|\alpha)\}$ which is shown to be equal to p(r) in Lemma 3.

Once again, we will analyse the time complexity of the algorithm by going through the actual steps of the algorithm rather than the more mathematical explanation given in [?]. The time complexity of the algorithm is dominated by the **for** loop as expected. Permuting the columns of A (Line 2) can be carried out in $\mathcal{O}(n)$ time using the Knuth shuffling algorithm [?]. Lines 3 and 4 take $\mathcal{O}(m)$ time, in order to generate the weight array representing the pmf and sample from it. In the **for** loop from lines 6 to 11, computing the permanent minors (Line 8) takes $\mathcal{O}(k2^k)$ time in the k^{th} iteration, as discussed previously. Subsequently, the weight array in Line 9 is computed in $\mathcal{O}(mk)$ time as it is an array of size m, with each element taking $\mathcal{O}(k)$ time to compute as it is a sum of k known terms. Sampling from this pmf takes $\mathcal{O}(m)$ time. The final sorting step outside the loop, in Line 12 takes $\mathcal{O}(n\log n)$ time, so it is dominated by the loop. Each iteration of the loop takes $\mathcal{O}(k2^k) + \mathcal{O}(mk)$ time in total. So for all iterations, this takes

$$\sum_{k=2}^{n} \mathcal{O}(k2^k) + \mathcal{O}(mk) = \mathcal{O}(n2^n) + \mathcal{O}(mn^2)$$
(15)

time.

The space is dominated by the weight array in Line 9, which is of size $\mathcal{O}(m)$.

Hence, the Boson Sampling Problem can be run in $\mathcal{O}(n2^n + \text{poly}(m, n))$ time using Algorithm B.

6 Implementation

In this section, we will discuss the actual implementation of Algorithm B described above. The paper by Clifford and Clifford which demonstrated the two new algorithms is still relatively new, which explains why there have not been any open-source large scale benchmark tests for actual implementations of the algorithm.

6.1 Original R Implementation

At the time of writing this paper, the only open-source implementation is at [?], by the authors of the paper. The implementation is in written as a package written in R and is freely available to download and use from CRAN. This package, named 'Boson Sampling' contains an implementation of Algorithm B, along with the required functions that it depends on. The relevant functions are described below.

bosonSampler

The package provides a function 'bosonSampler(A, sampleSize, perm=FALSE)', which implements the Boson Sampling Algorithm B from [?]. As the function definition suggests, it takes three arguments

- A: The first n columns of an $m \times m$ random unitary matrix.
- sampleSize: The number of independent sample values to be taken from the distribution, where each value is a vector of size n.
- perm: Takes the value TRUE if the permanents and pmfs associated to each sample are required to be returned. By default, perm=FALSE unless specified otherwise.

Note that we do not need to explicitly provide the values m and n as the algorithm suggests since those values are obtained by simply checking the dimensions of the input $m \times n$ matrix using the $\operatorname{ncol}(A)$ and $\operatorname{nrow}(A)$ functions from R. This implementation allows us to obtain multiple samples from the Boson Sampling distribution using the same input matrix A by repeatedly looping over the algorithm the desired number of times. One notable difference is that the input matrix A is transposed right at the start, with the reason being that R stores matrices in column order so transposing the matrix makes operations easier. When accessing elements of A, the indices have been adjusted accordingly. Semantically, the function implements the algorithm exactly as it is, which ensures its' correctness (provided the inputs received and additional functions used are also correct).

cxPermMinors

The package provides a set of functions for computing permanents of matrices: cxPerm(A), rePerm(B), cxPermMinors(C). Algorithm B requires the values

 $\{\operatorname{Per} B_{k,l}^{\diamond}, l \in [k]\}$ to be computed in each iteration of the loop, and in order to do this, the bosonSampler function makes a call to the cxPermMinors function.

7 Results

8 Conclusion

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