

COM SCI 239

QUANTUM PROGRAMMING:  
ALGORITHMS IN QISKIT

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# 1 Design

## 1.1 Mathematical explanation<sup>1</sup>

All four algorithms take as input a function  $f$  in the form:

$$f : \{0, 1\}^n \rightarrow \{0, 1\}^m$$

In the case of Deutsch-Josza, Bernstein-Vazirani, and Grover, we have  $m = 1$ . Simon, on the other hand, has  $m = n$ .

For Deutsch-Josza, Bernstein-Vazirani, and Simon, the quantum oracle of  $f$ , denoted  $U_f$ , is defined as:

$$U_f |x\rangle |b\rangle = |x\rangle |b \oplus f(x)\rangle$$

Here, we have  $b \in \{0, 1\}^m$ . The  $\oplus$  operator represents bitwise XOR, or equivalently bitwise addition mod 2.

Grover's algorithm makes use of a gate denoted by  $Z_f$ , which is defined as:

$$Z_f |x\rangle = (-1)^{f(x)} |x\rangle$$

Notice that this is precisely the application of the phase kickback trick for gates of a form equivalent to  $U_f$  where  $|b\rangle = |-\rangle$ :

$$Z_f |x\rangle |-\rangle = (-1)^{f(x)} |x\rangle |-\rangle$$

Clearly, we can also define  $Z_f$  as a quantum oracle for Grover's algorithm in the same way  $U_f$  was defined for the three other algorithms. Given that all four algorithms' quantum oracles have the same general form, we created a single function which could generate a quantum oracle and reused it for each program.

**Lemma 1.**  $\sum_{q \in \{0,1\}^k} |q\rangle \langle q| = I_{2^k}$  where  $I_{2^k}$  is the identity matrix in  $\mathcal{H}_{2^k}$  (corresponding to  $k$  qubits).

*Proof.* Let  $q \in \{0, 1\}^k$ . We have  $|q\rangle = [q_1 \ q_2 \ \dots \ q_{2^k}]^T$  where  $q_i = 1$  for some  $1 \leq i \leq 2^k$  and  $q_j = 0$  for all  $j \neq i$ . Then,  $|q\rangle \langle q|$  is a  $2^k \times 2^k$  matrix of the form  $[q_{ab}]$  where:

$$q_{ab} = \begin{cases} 1 & \text{if } a = b = i \\ 0 & \text{otherwise} \end{cases}$$

For  $\alpha, \beta \in \{0, 1\}^k$  such that  $\alpha \neq \beta$ , we have  $|\alpha\rangle \neq |\beta\rangle \implies |\alpha\rangle \langle \alpha| \neq |\beta\rangle \langle \beta|$ . Thus:

$$\sum_{q \in \{0,1\}^k} |q\rangle \langle q| = \begin{bmatrix} 1 & 0 & \dots & 0 \\ 0 & 0 & \dots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \dots & 0 \end{bmatrix} + \begin{bmatrix} 0 & 0 & \dots & 0 \\ 0 & 1 & \dots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \dots & 0 \end{bmatrix} + \dots + \begin{bmatrix} 0 & 0 & \dots & 0 \\ 0 & 0 & \dots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \dots & 1 \end{bmatrix} = I_{2^k}$$

□

---

<sup>1</sup>**NB:** This section is identical to Section 1.1 in the *Algorithms in PyQuil* report. Regardless, it is repeated here for the sake of completeness.

Consider the application of  $U_f$  to the outer product of  $|x\rangle |b\rangle$  with itself:

$$\begin{aligned} U_f |x\rangle |b\rangle \langle x| \langle b| &= |x\rangle |b \oplus f(x)\rangle \langle x| \langle b| \\ &= |x\rangle \langle x| |b \oplus f(x)\rangle \langle b| \end{aligned}$$

Let  $k = n + m$ . By taking the sum of this over all bit strings  $xb \in \{0, 1\}^k$ , we get:

$$\begin{aligned} \sum_{xb \in \{0, 1\}^k} U_f (|x\rangle |b\rangle \langle x| \langle b|) &= U_f \left( \sum_{xb \in \{0, 1\}^k} |x\rangle |b\rangle \langle x| \langle b| \right) \\ &= U_f \circ I_{2^k} \\ &= U_f \end{aligned}$$

And so:

$$U_f = \sum_{xb \in \{0, 1\}^k} |x\rangle \langle x| |b \oplus f(x)\rangle \langle b| \quad (1)$$

## 1.2 Python implementation

In our PyQuil assignment, we noted that we had to use the `numpy` library in order to implement [Equation 1](#) in Python, as PyQuil does not have native methods of creating custom gates from scratch. Qiskit, on the other hand, possesses its own way to define and manipulate custom gates on the matrix level: the `Operator` class [[Qiskit, 2020a](#)], as seen in [Code Block 1](#).

We chose to make this change for a few reasons. For one, using the `Operator` class would allow for easier interoperability with actual Qiskit QuantumCircuits [[Qiskit, 2020c](#)], and while we could just as easily use `numpy` to calculate  $U_f$  and then convert it to an `Operator` just before appending it to a circuit, we felt that using the `Operator` class for said calculation would be more idiomatic Qiskit. Also, we found that interestingly, the `Operator` class was capable of performing the necessary tensor products rather faster than `numpy`. Specifically, the time taken to generate  $U_f$  using `Operator` methods was about 0.847% faster than doing so with `numpy` (the full set of readings is omitted for brevity).

Apart from this minor change, our implementation of generating  $U_f$  is the same it was with PyQuil. That is, as seen in [Code Block 1](#), we still accept a function definition `f` in the form of a Python dictionary, not a physical function, for the same reasons as discussed in our PyQuil report (it allows more flexibility of input, and it makes it far easier to parametrize our solutions in `n`).

Also, the `oracle.gen_matrix` method, along with any other methods which require the dimension of the domain and/or range of  $f$ , still accepts `n` and `m` (as defined previously) as arguments. While `m` is algorithm-dependent and so assigned inside the code, the user can pass in a value for  $n$  using the `--num` option<sup>2</sup>.

---

<sup>2</sup>Discussed in further detail in the README.

```

134     for xb in range(0, int(2**(n+m))):
135         # Convert index to binary and split it down the middle
136         x = f'{xb:0{n+m}b}'[:int(n)]
137         b = f'{xb:0{n+m}b}'[int(n):]
138         # Apply f to x
139         fx = f[x]
140         # Calculate b + f(x)
141         bfx = f'{int(b, 2) ^ int(fx, 2):0{n}b}'
142
143         # Vector representations of x, b, and b+f(x)
144         xv = Operator(np.zeros((2**n,1)))
145         xv.data[int(x, 2)] = 1.
146         bv = Operator(np.zeros((2**m,1)))
147         bv.data[int(b, 2)] = 1.
148         bfxv = Operator(np.zeros((2**m,1)))
149         bfxv.data[int(bfx, 2)] = 1.
150
151         # Accumulate (|x><x| (*) |b + f(x)><b|) into the sum
152         # (*) is the tensor product
153         U_f += (xv * xv.transpose()).tensor(bfxv * bv.transpose())
154
155     return U_f

```

Code Block 1: Excerpt from the `gen_matrix` function in `oracle.py` showing the implementation of Equation 1 using Qiskit’s `Operator` class.

### 1.2.1 Readability

In terms of the readability of our oracle functions such as the one seen in Code Block 1, there is not much we can say here that is different from what we described in our PyQuil report. Our reasons for implementing `gen_matrix` as seen above are the same—we felt it is more concise to iterate through the decimal equivalents of `x` and `b` and extract relevant bits on the fly—and we still comment our code with PyDocs, loosely conforming to PEP/8 guidelines [van Rossum et al., 2001].

In terms of the readability of our actual implementations of the different algorithms, we found that Qiskit’s way of building quantum circuits with custom gates was in some ways more intuitive and cleaner than that of PyQuil. Compare the following lines of code:

```

# p is a PyQuil Program, u_f is a NumPy matrix, n is the number of qubits
U_f_def = DefGate('U_f', u_f)
U_f = U_f_def.get_constructor()
p += U_f_def
p += U_f(*(tuple(range(n))))

```

```
# circuit is a Qiskit QuantumCircuit, u_f is a NumPy matrix, n is the
number of qubits
U_f = Operator(u_f)
circuit.append(U_f, list(range(n))[::-1])
```

The equivalent Qiskit implementation is much less verbose, without sacrificing clarity. By accepting a list of qubit indices instead of requiring them to be passed as arguments, it is far easier to parametrize a gate in this manner.

However, note the addition of `[::-1]` to the Qiskit code. One perhaps unintuitive aspect of Qiskit’s multi-qubit gates is that qubits are ordered in reverse, where qubit 0 is the least significant qubit, and qubit  $n$  is the most significant qubit. As such, the way we generate matrices for each  $U_f$  require that we pass in a reversed list of qubits; similarly, for multi-qubit outputs, we must again reverse the output.

## 2 Evaluation

### 2.1 Shared code

Just as with PyQuil, as opposed to copying identical sections of code between each of our four programs, we opted to have a central module, `oracle.py`, which contains common code relating to the generation of functions  $f$  (bit mappings, to be precise) and quantum oracles, which was discussed in the previous section.

Of course, there is still some duplication of code between programs. Each program has a `getUf` function (named `getZf` for Grover) which checks if a  $U_f$  matrix for the desired value of  $n$  has already been generated and stored<sup>3</sup>, in which case it loads it from its file; otherwise, it simply generates a new  $U_f$ . All the implementations of this function are identical.

Also, our programs use the `argparse` library [Van Rossum and Drake, 2009] to read and process user-defined options<sup>4</sup>. The code for doing so is largely the same across programs, save for some minor algorithm-specific differences (like the names). Furthermore, we included code that measures each program’s execution time as well.

Overall, we estimate that each program file has an average of approximately 33 lines of code that is shared (within a reasonable threshold of a few characters’ difference) with the other files. Excluding `oracle.py`, the average length of a single program’s Python file is 167 lines (not including `.`). As such, the percentage of shared code between programs is approximately 20%.

Notice that this number is lower than the 31% code reuse percentage quoted for PyQuil. We attribute this drop to the fact that, overall, Qiskit is less verbose than PyQuil, as well as the

<sup>3</sup>This generation and storing of matrices ahead of actually running the algorithms is discussed in the README.

<sup>4</sup>Also discussed in the README.

fact that we added some more wrapping code in order to measure execution time provide more verbose output if desired.

It is possible for us to minimize code reuse. A lot of the aforementioned code to measure execution time is largely identical between programs, save for small differences. With more time, perhaps this could be removed and placed into a separate utility Python module.

## 2.2 Visualization

Qiskit comes packaged with a number of ways to visually express a built quantum circuit. This was extremely useful for us in running and testing our programs to make sure we had understood Qiskit's `QuantumCircuits` properly, especially in regards to qubit ordering (discussed below in Section 3).

For example, consider the generated diagram for a 2-bit input Deutsch-Josza circuit in Figure 1 below:

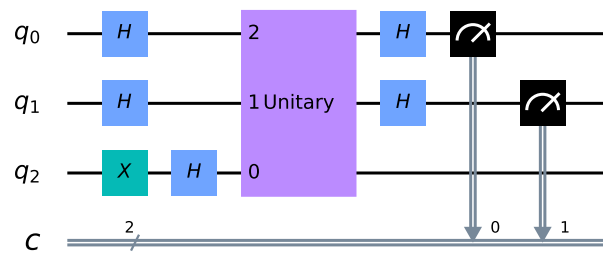


Figure 1: Circuit diagram for 2-bit input Deutsch-Josza.

Here, we can see precisely what is going on with each of our input qubits. Furthermore, by observing the input indices on the purple *Unitary* gate, we can ensure that the order of qubits being supplied is as desired.

By supplying the `--verbose` or `-v` option when running our programs, one can view such a circuit diagram within their terminal as well.

## 2.3 Testing

As mentioned earlier, our implementation of generating and storing quantum oracle gate matrices and usage of `argparse` to handle user input greatly facilitated testing. As such, there was no compilation time to speak of, and execution was incredibly fast. Instead, we simply used Qiskit's Aer simulator as the backend for running our programs.

For our implementation of Grover, we configured the program such that it would automatically calculate the minimum number of trials required to minimize error based on the size of the

input using the equation  $k = \lfloor \pi\sqrt{N}/4 \rfloor$  with  $N = 2^n$ . For our implementation of Simon, given that a full "iteration" was  $n - 1$  applications of the quantum oracle (to generate  $n - 1$  values for  $y$ ), we multiplied the user's input number of trials by 4 to minimize error using the equation  $\mathcal{P}(\text{not linearly independent}) = \exp(-t/4)$  with  $t$  being the number of trials.

We tested our code with a variety of values of  $n$  (the length of the input bit string). For Simon and Grover, we also tested with differing values of  $t$  (the number of trials); this was unnecessary for Deutsch-Josza and Bernstein-Vazirani, which are deterministic algorithms. We found that our results very closely matched the expected outcomes of applying our algorithms.

For each value of  $n$  up to about 5-9 depending on the algorithm, we generated randomized functions (following the appropriate constraints for its corresponding algorithm) and subsequently  $U_f$  matrices, storing them locally for later use. We limited  $n$  to 5-9 mostly because any values greater than those resulted in massive  $U_f$  matrices (the file size of a 10-qubit  $U_f$  matrix stored as an `.npy` array exceeded 100 MB), or otherwise took too much time to generate such a  $U_f$ .

For smaller  $n$ , however, we were able to make some observations:

- Deutsch-Josza was the fastest algorithm of the four, able to reach 9 qubits before crashing.
- Whether the input function to Deutsch-Josza was balanced or constant did not seem to significantly affect execution time.
- Despite being probabilistic, Simon's and Grover's algorithms are in fact quite accurate even for a smaller number of trials. Of course, this is in an ideal setting on a simulator without noise.
- For PyQuil, we found that compiling our code was incredibly CPU-intensive, taking many minutes to simply compile our quantum circuit into the QUIL assembly language. On the other hand, Qiskit needed to do no such thing, perhaps due to the fact that Qiskit's fundamental gate set is far larger than PyQuil—PyQuil at its core uses the gate operators  $R_Z(\theta)$ ,  $R_X(\frac{k\pi}{2})$ ,  $CZ$  [Rigetti, 2020], while Qiskit uses the universal gate set  $U_1$ ,  $U_2$ ,  $U_3$ ,  $CX$  and  $I$  [Qiskit, 2020b], allowing for much more flexibility in implementing other gates [Zhang et al., 2018].
- We were only able to test Simon's algorithm with up to 5 input qubits; beyond that, the time taken to generate  $U_f$  was unreasonably long. This is expected behavior; while the other three algorithms have one additional helper qubit, Simon's algorithm must use  $n$  additional qubits. Thus, increasing the input bitstring by 1 causes the resulting quantum oracle matrix to balloon to  $2^4 = 16$  times its original size.

## 2.4 Scalability

As with PyQuil, we tested our code on a machine with an **8-core Intel(R) Core(TM) i7-8550U @ 1.80GHz** and a **RAM of 16GiB System Memory (x2 8GiB SODIMM DDR4 Synchronous Unbuffered 2400 MHz)**.

As mentioned earlier, there was no compilation time to speak of as there was with PyQuil, and we were therefore able to measure execution time simply by using Python's built-in `time`

module, measuring the time taken to call our functions (from the creation of a circuit to the obtaining of a `counts` object).

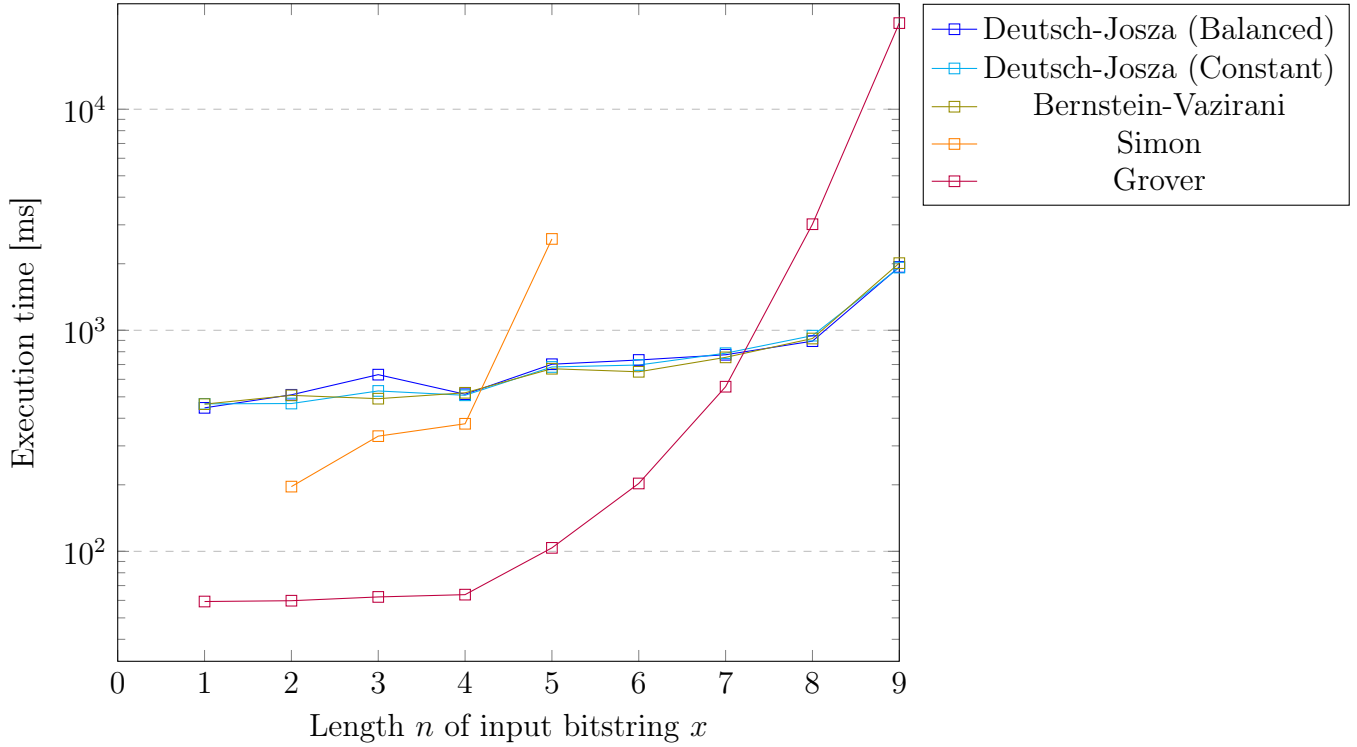


Figure 2: Graph of execution time (ms) vs. the length of the input bitstring. Note that the  $y$ -axis is logarithmically scaled.

Figure 2 portrays execution time against  $n$ . The execution time growths of Deutsch-Josza and Bernstein-Vazirani are both very similar, with a soft exponential growth (shown on a logarithmic graph as apparently linear). Simon’s algorithm appears to grow exponentially, at a much, much faster rate than DJ and BV.

Note the execution time growth for Grover’s algorithm, on the other hand, which has the shape of an exponential curve despite being plotted on a logarithmic scale. Although enough data could not be collected for Simon’s algorithm, it is expected that a similar growth would be observed for it as well. This is expected behavior, and appears for a specific reason: the number of times Simon’s and Grover’s algorithm are to be run is directly dependent on the number of input qubits. In the case of Simon’s algorithm, to minimize error, the entire circuit must be executed  $(n - 1) \times 4m$  times, where  $m$  is a user-defined parameter (chosen to be equal to 5 for our tests). For Grover’s algorithm, the entire circuit is to be called once, but the Grover rotation gate is appended to the circuit  $\lfloor \frac{\pi}{4} \sqrt{2^n} \rfloor$  times. As such, it is natural that the execution time for each of these algorithms increases at a double-exponential rate. On a log-log graph, the plots for these two algorithms’ growth rates would indeed appear in a linear form.



## 3 Reflection on Qiskit

### 3.1 Learning to use Qiskit

Some aspects of Qiskit that were rather easy to learn are:

- For basic circuits, Qiskit is incredibly intuitive. When we look at how to utilize multiple basic gates such as Hadamard or CNOT, we can easily instantiate them into our circuit by applying a range, with `circuit.H(range(n))`. This proved to be extremely useful and much easier to read when debugging in the future. It also gave a lot of flexibility in terms of easily maneuvering our gates across qubits.
- The fact that we didn't have to start a server to compile our Qiskit circuit was actually amazing. The `qasm-simulator` proved to be so easy to use and incredibly fast. As opposed to PyQuil where we ran into many problems with the `qvm` and `qcuil` servers, we didn't face any problems with compilation whatsoever.
- The ability to customize and set your measure registers in Qiskit was quite easy to pick up. It provided an easy and intuitive way to set a range, list or integer to determine which qubits will be measured and to which classical bits.

```
circuit.measure(n,n)
circuit.measure(range(1, n+1), range(n))
```

This made circuit customization super easy and by having it all in one line, made our life super easy in terms of debugging and understanding how it was used.

Some aspects of Qiskit that were hard to learn are:

- The fact that Qiskit has its bits flipped was a tough concept to grasp. Mainly due to the fact that our matrices were generated in a way that fits a more traditional qubit model, so when it came to debugging, it was quite common to second guess oneself as to whether or not we need to invert the range: As it turns out, 100% of the time the answer to that

```
circuit.append(U_f, range(n)[::-1])
```

question was yes, and we're sure after more exposure to programming in Qiskit that this notation will become second nature.

- Qiskit operators have amazing functionality and usage, however, using them correctly proved to be a learning curve. One, it's quite hard to find amazing documentation on how operators work and how to utilize it completely on the Qiskit documentation website. Two, if you generate your Operator from a numpy array, you can't name/label it because apparently labeling happens at the Gate level.
- The different Qiskit elements proved to be challenging to wrap our heads around in terms of looking through the documentation. When looking for API references, it was definitely hard at first to determine where to look. Most of our API's we found in Terra, but understanding where to look at first definitely was a challenge

Qiskit is great when it comes to usability and testing. Their embedded print functionality made debugging simple and reduced a lot of the overhead complexity of the PyQuil implementation. We also found that the fact that there is no quilc or qvm server allowed for greater abstraction and usability as developers. So long are the days when I had to worry about running a server to compile my quantum circuit. Finally, the speed at which Qiskit performs our algorithms far surpasses that of PyQuil. The compilation time of Qiskit with the Aer backend makes development quick and responsive. It was extremely fast to tell whether or not our circuits are working and to simply bootstrap code.

Qiskit doesn't allow you to label your custom operators/instructions when appending it to a circuit. This can help for debugging however it's not too necessary as the developer is responsible for understanding what their circuit generally looks like. We mention this later in our suggestion, but a way to reverse the qubit ordering would be fantastic. You can do it already when it comes to drawing the circuit through:

```
circuit.draw('mpl', reverse_bits=True)
```

However, letting it work on an implementation level would be fantastic.

### 3.2 A suggestion

Using other quantum programming languages and through academia, we have been continuously taught by making the top qubit in a graphical diagram to be the most significant qubit and thus our custom Operators will reflect that understanding. Furthermore, in most conventions we've seen, the tensor product  $|a\rangle \otimes |b\rangle$  is expressed as  $|ab\rangle$ ; with Qiskit, it is expressed as  $|ba\rangle$ .

It would be extremely helpful if Qiskit made their qubit ordering such that the most significant qubit is qubit 0. Mainly because our brains are wired to look out the circuit display top to bottom and having to continually invert our Operators and our outputs are quite cumbersome and unintuitive. A homogenization on the circuit level with other quantum programming languages and academia would greater support the usage and bootstrapping of Qiskit.

### 3.3 Reading the documentation

Three things we found great about Qiskit documentation are:

- The ease of starting with Qiskit through “Installing Qiskit” and “Getting Started with Qiskit” proved to be really helpful in terms of bootstrapping Qiskit for the first time. Qiskit has an amazing “Tutorials” section that was documentation amazingly.
- The “Advanced Circuits” subsection we found to be extremely helpful and made the process of programming the algorithms extremely smooth.
- The builtin print functionality which allowed us to simply run `print(circuit)` in order to visualize our circuit, as mentioned earlier. No documentation really needed, but they

also included a “Visualizing a Quantum Circuit” subsection that covers great material in terms of drawing to `matplotlib` and `LATEX`.

Three things we found challenging about Qiskit documentation are:

- The separation of API references through the Elements was quite confusing at first. Sometimes, as a developer, you just need to see all options and be able to scan through the API with just the objects themselves. Working with Qiskit Elements it was challenging at first as to where to look for specific API references and to check whether or not certain function calls were possible.
- If they gave more scripture describing the specific objects and their functionality. For example, if you look at the Terra API Reference in [Figure 3](#).



Figure 3: Terra API reference.

There is literally no description of what each object is used for and how it can be utilized with respect to a quantum circuit, transpiling, or visualization. A little description of each of them would make it more beginner-friendly and easier to reference.

- Following the discussion of the API reference, it would be great if while giving an example of how functions worked, if the API reference could also link you to the return object in question and give an example output. For example, consider [Figure 4](#).

It describes the API reference for `execute`. While it does have a link to the return object, it doesn’t tell us how we can utilize the job instance of Base Job. Some functions don’t even have example usage, so there is quite a good amount of flipping back and forth to understand the API.

Figure 4: `execute` API reference.

### 3.4 Translating key concepts

As is our understanding, below are some key concepts in quantum programming as they are referred to with Qiskit:

- **Backend:** Overarching class that envelops different providers (Aer or IBMQ) and their simulators
- **Shots:** Number of Trials
- **Operator:** Objects with a matrix that can be applied onto a Qiskit circuit
- **Transpiler:** Circuit  $\rightarrow$  Circuit. Primarily used to optimize of gates for a quantum program. It is used to optimize and produce better circuits for more robust implementation
- **Terra:** Foundation of Qiskit and has the underlying modules to build circuits, transpile, providers, visualization etc.
- **Aer:** High Performant simulators that allow the development of quantum programs to happen faster
- **Ignis:** Combatting noise through mitigation and filters and control over errors and gates
- **Aqua:** For real-world application and algorithms

### 3.5 Type checking in Qiskit

Qiskit automatically checks if custom matrices are unitary. Extremely important in the case of ensuring nothing exceedingly strange happens to our output. Qiskit also checks whether or not the dimensions of the custom gate you created matches the range you specified when you append it to the circuit.

If the length of the range you supply doesn't match the dimensions of the Operator, Qiskit will throw an error to prevent undefined behavior.

```
circuit.append(Uf, range(n))
```

## References

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