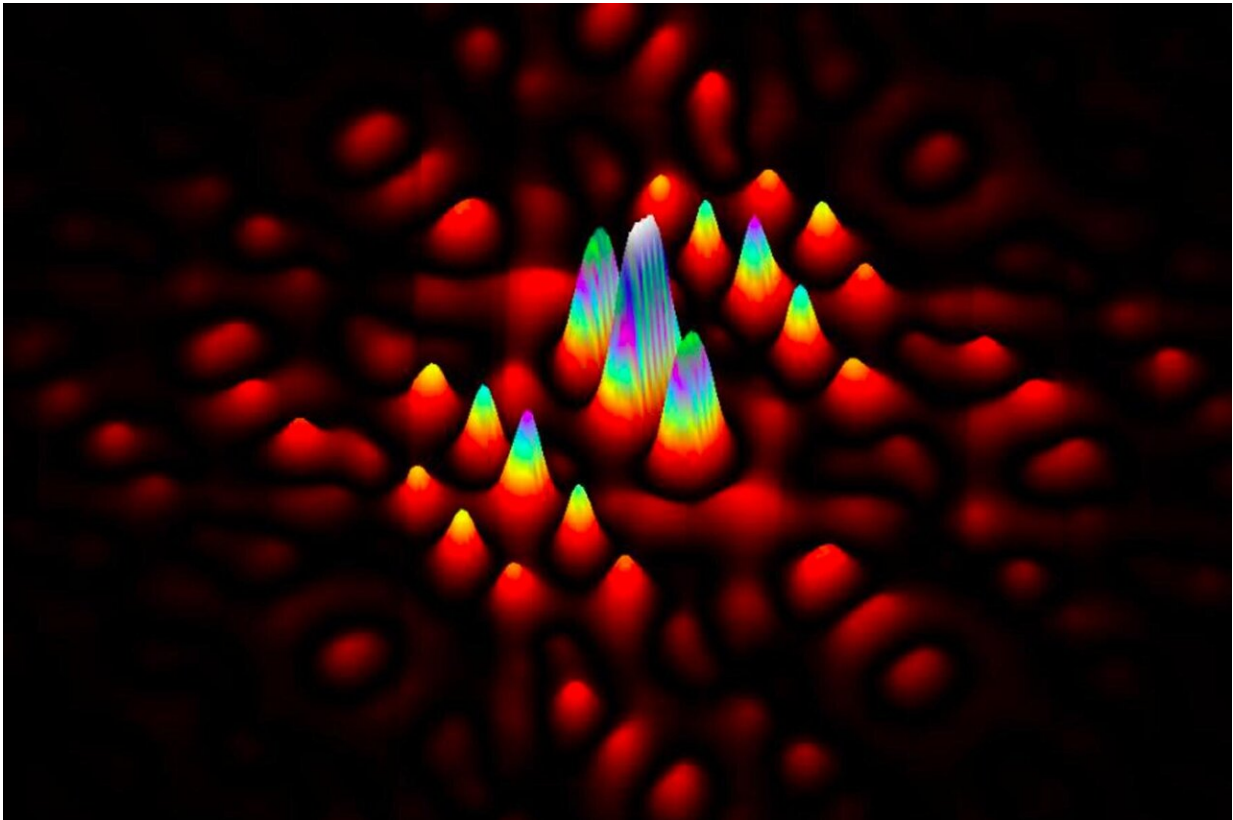


Machine learning to scale up the quantum computer

March 17 2020, by Dr Muhammad Usman and Professor Lloyd Hollenberg

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A map of electron wave function patterns, where the symmetry, brightness and size of features is directly related to the position of a phosphorus atom in silicon lattice. Credit: M.Usman/ University of Melbourne

Quantum computers are expected to offer tremendous computational

power for complex problems –currently intractable even on supercomputers—in the areas of drug design, data science, astronomy and materials chemistry among others.

The high technological and strategic stakes mean major technology companies as well as ambitious start-ups and government-funded research centers are all in the race to build the world's first universal quantum computer.

Building a quantum computer

In contrast to today's classical computers, where information is encoded in bits (0 or 1), quantum computers process information stored in quantum bits (qubits). These are hosted by quantum mechanical objects like electrons, the negatively charged particles of an atom.

Quantum states can also be binary and can be put in one of two possibilities, or effectively both at the same time—known as quantum superposition—offering an exponentially larger computational space with an increasing number of qubits.

This unique data crunching power is further boosted by entanglement, another magical property of quantum mechanics where the state of one qubit is able to dictate the state of another qubit without any physical connection, making them all 1's for example. Einstein called it a 'spooky action at distance'.

Different research groups in the world are pursuing different kinds of qubits, each having its own benefits and limitations. Some qubits offer potential for scalability, while others come with very long coherence times, that is the time for which quantum information can be robustly stored.



Over the next decade, quantum computers will move beyond physics research laboratories. Credit: Connie Zhou/IBM

Qubits in silicon are highly promising as they offer both. Therefore, these qubits are one of the front-runner candidates for the design and implementation of a large-scale quantum computer architecture.

One way to implement large-scale quantum computer architecture in silicon is by placing individual phosphorus [atoms](#) on a two-dimensional grid.

The single and two qubit logical operations are controlled by a grid of nanoelectronic wires, bearing some resemblance to classical logic gates for conventional microelectronic circuits. However, key to this scheme

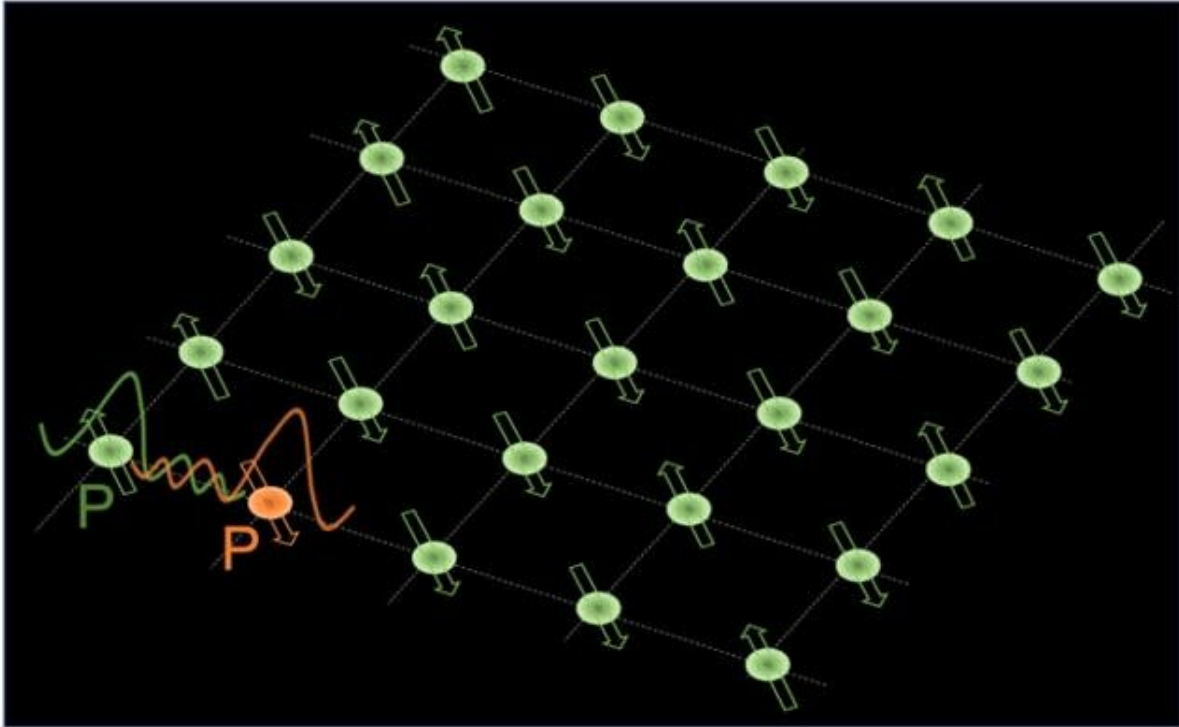
is ultra-precise placement of phosphorus atoms on the silicon grid.

The challenges

However, even with state-of-the-art fabrication technologies, placing phosphorus atoms at precise locations in silicon lattice is a very challenging task. Small variations, of the order of one atomic lattice site, in their positions are often observed and may have a huge impact on the efficiency of two qubit operations.

The problem arises from the ultra-sensitive dependence of the exchange interaction between the electron qubits on phosphorus atoms in silicon. Exchange interaction is a fundamental quantum mechanical property where two sub-atomic particles such as electrons can interact in real space when their wave functions overlap and make interference patterns, much like the two traveling waves interfering on water surface.

Exchange interaction between electrons on phosphorus atom qubits can be exploited to implement fast two-qubit gates, but any unknown variation can be detrimental to accuracy of quantum gate. Like logic gates in a conventional computer, the quantum gates are the building blocks of a quantum circuit.



An artist's impression of a large-scale quantum computer architecture based on phosphorus (P) atom qubits in silicon. The wave functions of electrons bound to P atom exhibit oscillations and constructive/destructive spatial overlaps of these wave functions give rise to large variations in interaction, introducing errors in quantum gates. Determination of Exact P atom positions could eliminate errors, paving the way for the ultimate goal of fault-tolerant universal quantum computing. Credit: M.Usman/ University of Melbourne

For phosphorus qubits in silicon, even an uncertainty in the location of qubit atom of the order of one atomic lattice site can alter the corresponding exchange interaction by orders of magnitude, leading to errors in two-qubit gate operations.

Such errors, accumulated over the large-scale architecture, may severely impede the efficiency of quantum computer, diminishing any quantum

advantage expected due to the quantum mechanical properties of qubits.

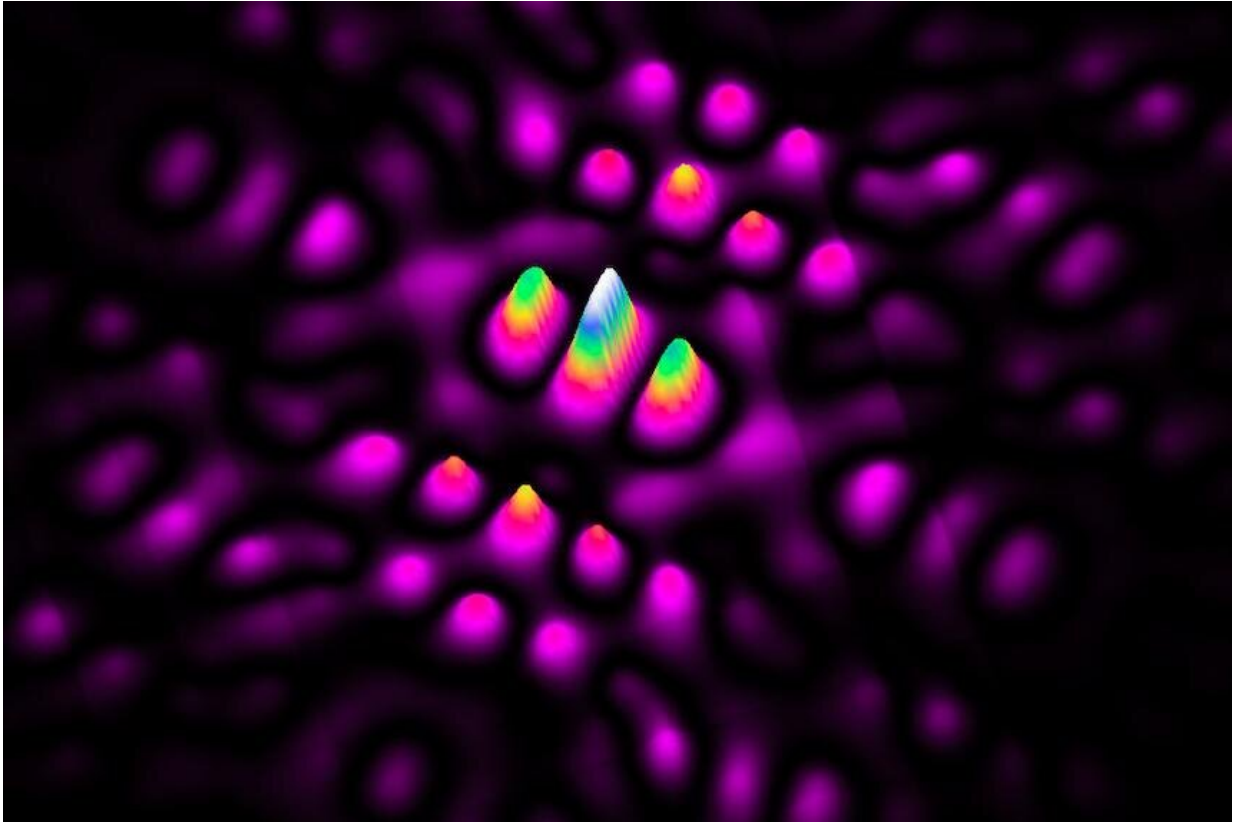
Finding exact coordinates of qubit atom

So in 2016, we worked with the Center for Quantum Computation & Communication Technology researchers at the University of New South Wales, to develop a technique that could pinpoint exact locations of phosphorus atoms in silicon.

The technique, reported in *Nature Nanotechnology*, was the first to use computed scanning tunneling microscope (STM) images of phosphorus atom wave functions to pinpoint their spatial locations in silicon.

The images were calculated using a computational framework which allowed electronic calculations to be performed on millions of atoms utilizing Australia's national supercomputer facilities at the Pawsey supercomputing center.

These calculations produced maps of electron wave function patterns, where the symmetry, brightness and size of features was directly related to the position of a phosphorus atom in silicon lattice, around which the electron was bound.



A map of electron wave function patterns, where the symmetry, brightness and size of features is directly related to the position of a phosphorus atom in silicon lattice. Credit: M.Usman/ University of Melbourne

The fact that each donor atom positions led to a distinct map, pinpointing of qubit atom locations, known as spatial metrology, with single lattice site precision was achieved.

The technique worked very well at the individual qubit level. However, the next big challenge was to build a framework that could perform this exact atom spatial pinpointing with high speed and minimal human interaction coping with the requirements of a universal fault tolerant quantum computer.

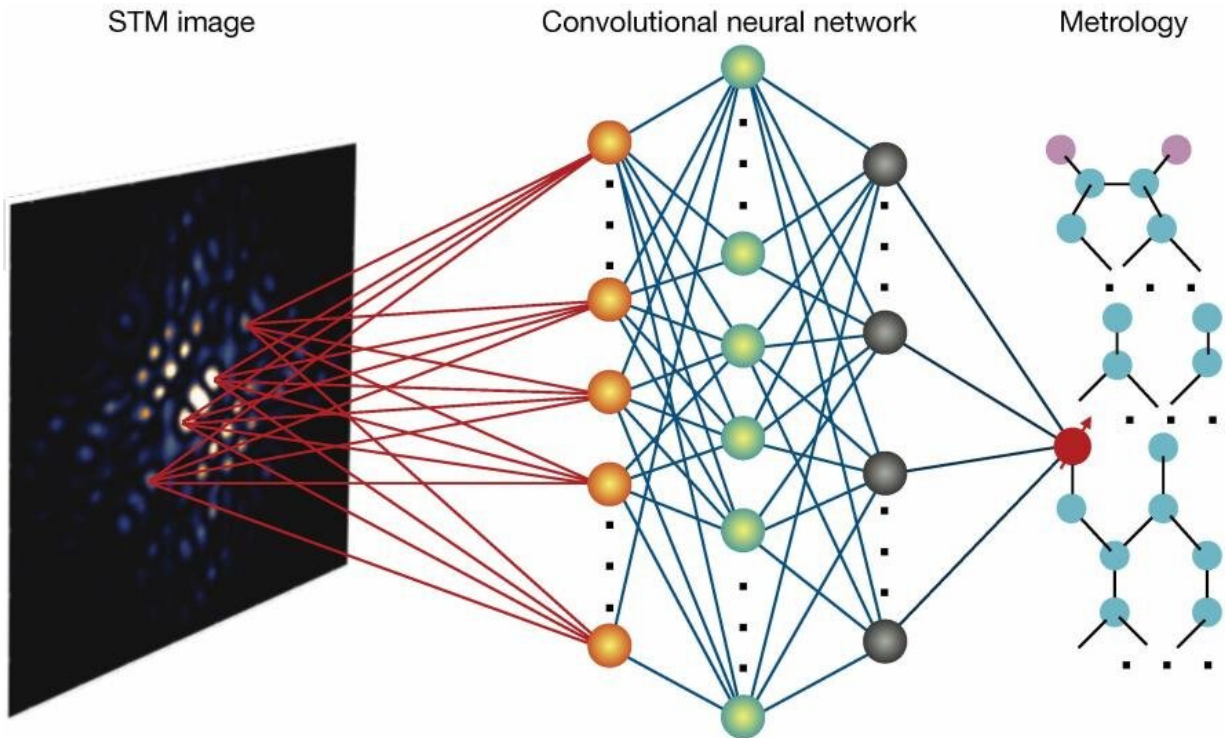
Machine learning

Machine learning is an emerging area of research which is revolutionizing almost every field of research, from medical science to image processing, robotics, and material design.

A carefully trained machine learning algorithm can process very large data sets with enormous efficiency.

One branch of machine learning is known as convolutional neural network (CNN) – an extremely powerful tool for image recognition and classification problems. When a CNN is trained on thousands of sample images, it can precisely recognize unknown images (including noise) and perform classifications.

Recognising that the principle underpinning the established spatial metrology of qubit atoms is basically recognizing and classifying feature maps of STM images, we decided to train a CNN on the computed STM images. The work is published in the NPJ Computational Materials journal.



Computed scanning tunneling microscope (STM) images of phosphorus atoms qubits in silicon used to train a convolutional neural network (CNN), capable of autonomous and high-throughput qubit characterization with an exact atom precision in both, their spatial locations and atom count. Credit: M.Usman/ University of Melbourne

The training involved 100,000 STM images and achieved a remarkable learning of above 99 percent for the CNN. We then tested the trained CNN for 17600 test images including blurring and asymmetry noise typically present in the realistic environments.

The CNN classified the test images with an accuracy of above 98 percent, confirming that this machine learning-based technique could process qubit measurement data with [high-throughput](#), high precision, and minimal human interaction.

This technique also has the potential to scale up for qubits consisting of more than one phosphorus atoms, where the number of possible image configurations would exponentially increase. However, machine learning-based framework could readily include any number of possible configurations.

In the coming years, as the number of qubits increase and size of quantum devices grow, [qubit](#) characterization via manual measurements is likely to be highly challenging and onerous.

This work shows how [machine learning](#) techniques such as developed in this work could play a crucial role in this aspect of the realization of a full-scale fault-tolerant universal quantum computer—the ultimate goal of the global research effort.

More information: M. Usman et al. Spatial metrology of dopants in silicon with exact lattice site precision, *Nature Nanotechnology* (2016). [DOI: 10.1038/nnano.2016.83](https://doi.org/10.1038/nnano.2016.83)

Muhammad Usman et al. Framework for atomic-level characterisation of quantum computer arrays by machine learning, *npj Computational Materials* (2020). [DOI: 10.1038/s41524-020-0282-0](https://doi.org/10.1038/s41524-020-0282-0)

Provided by University of Melbourne

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