

This is the command to get the latest release of QuESTlink:

```
Import["https://qtechtheory.org/questlink.m"];
```

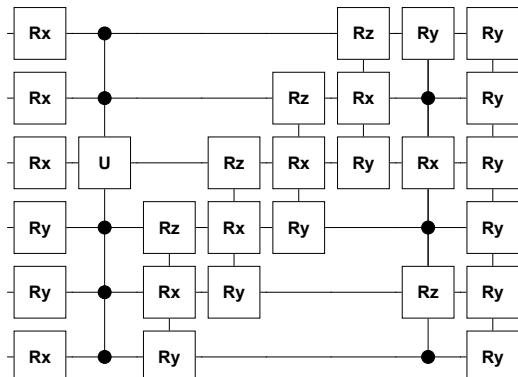
The next command downloads the pre-compiled single-thread simulator. For significantly faster multithreaded and GPU simulation, recompile QuESTlink (see guide here) and use **CreateLocalQuESTEnv[]**

```
CreateDownloadedQuESTEnv[];
```

Preparing an ansatz

Here's how we could hard-code a 6 qubit, 12 parameter ansatz. The parameters are labelled $\theta[i]$ for convenience, but they could be any symbols we want (θ_i , θ_i , etc)

```
Circuit[  
    Rx0[θ[1]] Ry1[θ[2]] Ry2[θ[3]] Rx3[θ[4]] Rx4[θ[5]] Rx5[θ[6]] ×  
    C0,1,2,4,5[U3[ $\begin{pmatrix} \text{Exp}[i\theta[7]] & 0 \\ 0 & \text{Exp}[-i\theta[7]] \end{pmatrix}]] ×  
    R[θ[7], Y0 X1 Z2] R[θ[8], Y1 X2 Z3] R[θ[9], Y2 X3 Z4] R[θ[10], Y3 X4 Z5] ×  
    C0,2,4[R[θ[11], Z1 X3 Y5]] ×  
    R[θ[12], Y0 Y1 Y2 Y3 Y4 Y5]];  
DrawCircuit[%]$ 
```



More often, we wish to generate an ansatz programmatically.

```

nQ = 6;
u = Block[{i = 1}, Flatten @ {
    Table[gq-1[θ[i++]], {q, nQ}, {g, {Rx, Ry, Rz}}],
    Table[CDeleteCases[Range[0, nQ - 1], q] [Uq[Exp[i θ[i++]] 0 1]], {q, 0, nQ - 1}],
    Table[R[θ[i++], Product[σq-1, {q, nQ}]], {σ, {Y, X, Z}}]];
nθ = Length[u];

DrawCircuit[u]

```

Since the parameters symbolic, we could study the ansatz operator analytically. Here's its first element, when every second parameter has value π .

```

u /. θ[_?EvenQ] → π;
CalcCircuitMatrix[%] // First // First
-  $\frac{1}{2} e^{\frac{1}{2} i \theta[3] + \frac{1}{2} i \theta[9] + \frac{1}{2} i \theta[15] - \frac{1}{2} i \theta[27]} \cos\left[\frac{\theta[1]}{2}\right] \cos\left[\frac{\theta[5]}{2}\right]$ 
 $\cos\left[\frac{\theta[7]}{2}\right] \cos\left[\frac{\theta[11]}{2}\right] \cos\left[\frac{\theta[13]}{2}\right] \cos\left[\frac{\theta[17]}{2}\right] \cos\left[\frac{\theta[25]}{2}\right] -$ 
 $i e^{-\frac{1}{2} i \theta[3] - \frac{1}{2} i \theta[9] - \frac{1}{2} i \theta[15] - \frac{1}{2} i \theta[27]} \sin\left[\frac{\theta[1]}{2}\right] \sin\left[\frac{\theta[5]}{2}\right] \sin\left[\frac{\theta[7]}{2}\right]$ 
 $\sin\left[\frac{\theta[11]}{2}\right] \sin\left[\frac{\theta[13]}{2}\right] \sin\left[\frac{\theta[17]}{2}\right] \sin\left[\frac{\theta[25]}{2}\right]$ 

```

Preparing a Hamiltonian

We could hard-code our own Hamiltonian...

```

h = .1 X0 Y1 Z2 + .3 Y0 X2 - .4 Z0 Z1 Z2;

```

```

CalcPauliSumMatrix[h] // Chop // MatrixForm

```

$$\begin{pmatrix} -0.4 & 0 & 0 & 0. - 0.1i & 0 & 0. - 0.3i & 0 & 0 \\ 0 & 0.4 & 0. - 0.1i & 0 & 0. + 0.3i & 0 & 0 & 0 \\ 0 & 0. + 0.1i & 0.4 & 0 & 0 & 0 & 0 & 0. - 0.3i \\ 0. + 0.1i & 0 & 0 & -0.4 & 0 & 0 & 0. + 0.3i & 0 \\ 0 & 0. - 0.3i & 0 & 0 & 0.4 & 0 & 0 & 0. + 0.1i \\ 0. + 0.3i & 0 & 0 & 0 & 0 & -0.4 & 0. + 0.1i & 0 \\ 0 & 0 & 0 & 0. - 0.3i & 0 & 0. - 0.1i & -0.4 & 0 \\ 0 & 0 & 0. + 0.3i & 0 & 0. - 0.1i & 0 & 0 & 0.4 \end{pmatrix}$$

but more likely we'll want to produce one from a file of coefficients and Pauli strings. Here we download a 369-term 6-qubit Lithium Hydride Hamiltonian

```

h = GetPauliSumFromCoeffs[
  "https://questlink.qtechtheory.org/demo_hamiltonian.txt"];

Length[h]
h[[;; 30]]

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- 6.52209 - 0.00168947 X0 + 0.000335609 X1 + 0.00233908 X0 X1 -
  0.00518865 X2 - 2.32678 × 10-6 X0 X2 - 0.00238276 X1 X2 - 0.000333484 X0 X1 X2 +
  0.0561302 X3 + 0.0000211588 X0 X3 + 0.0000198838 X0 X1 X3 - 0.000133652 X2 X3 -
  0.0000311241 X1 X2 X3 + 0.000547046 X4 + 0.00165752 X3 X4 - 0.00600013 X0 X3 X4 -
  0.00812442 X2 X3 X4 + 0.000447423 X5 - 0.0000517994 X0 X5 + 0.00019084 X0 X1 X5 -
  0.0000777401 X2 X5 - 0.0001908 X1 X2 X5 - 0.00630859 X3 X5 + 0.0661688 X4 X5 +
  0.0000169999 X0 X4 X5 + 0.00634842 X0 X1 X4 X5 - 0.000177071 X2 X4 X5 -
  0.00636179 X1 X2 X4 X5 + 0.00562722 X3 X4 X5 - 0.000346402 Y0 Y1
```

We can determine its ground-state exactly, which we'll later compare to that which our variational algorithm estimates

```

gs = Min @ Eigenvalues @ CalcPauliSumMatrix[h]
-7.88074
```

Preparing quantum states

To perform numerical simulation of our ansatz , we need to create quantum registers, which are stored in the backend QuEST process.

```

ψ = CreateQureg[nQ];
InitPlusState[ψ];

GetQuregMatrix[ψ] // Chop
{0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125,
 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125,
 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125,
 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125,
 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125, 0.125}
```

To apply our ansatz circuit to a state, we must replace the parameters with numerical values

```

ApplyCircuit[u /. θ[_] :> RandomReal[], ψ];

GetQuregMatrix[ψ]

{0.0224731 + 0.111599 I, 0.00523535 - 0.0521917 I,
 -0.0111565 - 0.0767799 I, -0.00378012 + 0.00136288 I, 0.00123122 - 0.0731973 I,
 -0.00457615 + 0.00651081 I, 0.00978255 - 0.00067529 I,
 -0.0267522 - 0.0498707 I, -0.0235968 - 0.0586924 I, -0.0191137 - 0.00769477 I,
 -0.00438554 - 0.0141594 I, -0.00728107 - 0.0635589 I, -0.00530986 - 0.0091749 I,
 -0.0142068 - 0.0662965 I, -0.0103575 - 0.0461789 I, -0.00615036 - 0.134759 I,
 0.0153867 - 0.0281887 I, -0.0302696 + 0.0147383 I, -0.0151482 + 0.00568313 I,
 -0.0364413 - 0.0803933 I, -0.0157579 + 0.0110711 I, -0.0448814 - 0.0796178 I,
 -0.0309596 - 0.0620335 I, -0.0692182 - 0.151883 I, -0.0319339 - 0.00383458 I,
 -0.0233075 - 0.111581 I, -0.0101306 - 0.0822589 I, -0.0323024 - 0.21812 I,
 -0.0189583 - 0.0844727 I, -0.0556178 - 0.218061 I, -0.0195894 - 0.167156 I,
 0.373976 + 0.176453 I, 0.0481885 - 0.116956 I, 0.0218496 - 0.0050421 I,
 0.0379506 - 0.0147816 I, -0.0269679 - 0.025903 I, 0.0375501 - 0.00818858 I,
 -0.0296966 - 0.0290688 I, -0.0344551 - 0.0168195 I, -0.0071038 - 0.0670466 I,
 0.0193431 - 0.021617 I, -0.0122202 - 0.0328829 I, -0.0153121 - 0.016552 I,
 0.0102312 - 0.0824418 I, -0.01866 - 0.020663 I, 0.00323491 - 0.0849524 I,
 0.00804622 - 0.0662416 I, 0.0994447 + 0.109923 I, 0.0100959 - 0.000568641 I,
 -0.0268904 - 0.0495882 I, -0.0245207 - 0.0355702 I, -0.0215095 - 0.099105 I,
 -0.0280162 - 0.0379397 I, -0.0312304 - 0.0979598 I, -0.0138876 - 0.0789094 I,
 0.17717 - 0.0295302 I, -0.0106074 - 0.0458956 I, -0.00641741 - 0.133966 I,
 0.00722081 - 0.101594 I, 0.237598 + 0.0403602 I, -0.00222075 - 0.103981 I,
 0.2454 - 0.000476521 I, 0.171626 + 0.0649399 I, 0.462416 - 0.101295 I}

```

We must create a “working Qureg” $\mathbf{h}\psi$ to compute expected values. We see our starting state isn’t especially close to the ground-state.

```

hψ = CreateQureg[nQ];
CalcExpecPauliSum[ψ, h, hψ]
-6.55243

```

We’ll also need additional Quregs to keep track of our fixed input state $\mathbf{in}\psi = |\+\rangle$, and $\mathbf{n}\theta$ derivative states $\mathbf{d}\psi[i] = \left| \frac{\partial\psi}{\partial\theta[i]} \right\rangle$ (where $\psi = u(\theta) \mathbf{in}\psi$)

```

inψ = CreateQureg[nQ];
InitPlusState[inψ];

dψ = CreateQuregs[nQ, nθ];

```

Computing variational observables

Let’s re-randomise the ansatz parameters.

```
vθ = Table[θ[i] → RandomReal[], {i, nθ}]
{θ[1] → 0.302214, θ[2] → 0.701127, θ[3] → 0.250832,
θ[4] → 0.366587, θ[5] → 0.184505, θ[6] → 0.134426, θ[7] → 0.2041,
θ[8] → 0.869532, θ[9] → 0.209075, θ[10] → 0.221281, θ[11] → 0.415592,
θ[12] → 0.598776, θ[13] → 0.91903, θ[14] → 0.0748652, θ[15] → 0.824325,
θ[16] → 0.0585614, θ[17] → 0.4362, θ[18] → 0.434674, θ[19] → 0.250115,
θ[20] → 0.0151225, θ[21] → 0.71593, θ[22] → 0.408774, θ[23] → 0.793994,
θ[24] → 0.943991, θ[25] → 0.711386, θ[26] → 0.455652, θ[27] → 0.0252188}
```

We set each $\mathbf{d}\psi[i] = \left| \frac{\partial \mathbf{u}}{\partial \theta[i]} \right| \mathbf{in}\psi \rangle$ for the given assignment of θ

```
CalcQuregDerivs[u, inψ, vθ, dψ];
```

The imaginary time ‘tensor’ is simply $\text{Re}[\langle \frac{\partial \psi}{\partial \theta[i]} || \frac{\partial \psi}{\partial \theta[j]} \rangle]$

```
m = Re @ CalcInnerProducts[dψ];
```

```
m // Chop // MatrixForm
```

0.25	0	-0.16127	0.25	0	-0.0458649
0	0.25	0	0	0	0
-0.16127	0	0.25	-0.16127	0	0.0295865
0.25	0	-0.16127	0.25	0	-0.0458649
0	0	0	0	0.25	0
-0.0458649	0	0.0295865	-0.0458649	0	0.25
0.25	0	-0.16127	0.25	0	-0.0458649
0	0	0	0	0	0
-0.191007	0	0.123215	-0.191007	0	0.035042
0.25	0	-0.16127	0.25	0	-0.0458649
0	0	0	0	0	0
-0.100933	0	0.0651097	-0.100933	0	0.0185171
0.25	0	-0.16127	0.25	0	-0.0458649
0	0	0	0	0	0
-0.0186988	0	0.0120622	-0.0186988	0	0.00343048
0.25	0	-0.16127	0.25	0	-0.0458649
0	0	0	0	0	0
-0.105625	0	0.0681363	-0.105625	0	0.0193779
-0.0512571	0.0213583	-0.0512571	-0.0512571	-0.00824045	0.0512571
-0.167895	-0.000900317	0.167895	-0.167895	0.00233352	-0.167895
-0.0385019	-0.0124444	0.0385019	-0.0385019	0.00419934	0.00860778
-0.114527	-0.0181217	0.114527	-0.114527	0.00807583	0.0226566
-0.14819	-0.0466023	0.14819	-0.14819	0.0293635	0.0392888
-0.0793218	-0.0358798	0.0793218	-0.0793218	0.015727	0.0224444
-0.00813621	0.00265984	0	-0.00813621	0.00320796	0
0.039606	-0.0140054	0	0.039606	-0.00428338	0
0.0002884	0	-0.000447077	0.0002884	0	-0.00157201

We set $|\psi\rangle = u(\theta) |\mathbf{in}\psi\rangle$, and $|\mathbf{h}\psi\rangle = \mathbf{h} |\psi\rangle$

```
ApplyCircuit[u /. vθ, CloneQureg[ψ, inψ]];
ApplyPauliSum[ψ, h, hψ];
```

The energy gradient is then simply $\text{Re}[\langle \psi | \mathbf{h} | \frac{\partial \psi}{\partial \theta[i]} \rangle]$

```
v = Re @ CalcInnerProducts[hψ, dψ];
v // MatrixForm

$$\begin{pmatrix} -0.013409 \\ -0.00539944 \\ 0.0379003 \\ -0.013409 \\ -0.0105197 \\ 0.0604042 \\ -0.013409 \\ 0.00468087 \\ 0.0250343 \\ -0.013409 \\ -0.0564997 \\ 0.00145499 \\ -0.013409 \\ -0.0671572 \\ -0.0213131 \\ -0.013409 \\ -0.0855882 \\ 0.0411782 \\ -0.0196398 \\ -0.0389648 \\ -0.010845 \\ 0.022991 \\ 0.0449788 \\ 0.000658498 \\ 0.00793236 \\ -0.0329018 \\ 0.018517 \end{pmatrix}$$

```

A single iteration of imaginary time evolution would use these observables to update the parameters via $\mathbf{m}\Delta\theta = -\mathbf{v}\Delta t$

```

Δθ = Δt LinearSolve[m, -v];
Δθ // MatrixForm

```

$$\begin{pmatrix} -0.112232 \Delta t \\ -0.247915 \Delta t \\ 1.37134 \Delta t \\ -0.112232 \Delta t \\ 0.191551 \Delta t \\ -1.42699 \Delta t \\ -0.112232 \Delta t \\ -0.0810106 \Delta t \\ 0.0389451 \Delta t \\ -0.112232 \Delta t \\ 0.408794 \Delta t \\ -0.26508 \Delta t \\ -0.112232 \Delta t \\ 0.501992 \Delta t \\ -0.0693181 \Delta t \\ -0.112232 \Delta t \\ 0.741091 \Delta t \\ -0.583384 \Delta t \\ 1.50634 \Delta t \\ -1.15593 \Delta t \\ 0.414318 \Delta t \\ -0.365748 \Delta t \\ -0.275681 \Delta t \\ -0.706214 \Delta t \\ -0.0677737 \Delta t \\ 0.379378 \Delta t \\ -0.0555886 \Delta t \end{pmatrix}$$

hence the updated parameters, dependent on Δt , would be

```

vθ[[All, 2]] += Δθ
{0.302214 - 0.112232 Δt, 0.701127 - 0.247915 Δt, 0.250832 + 1.37134 Δt,
 0.366587 - 0.112232 Δt, 0.184505 + 0.191551 Δt, 0.134426 - 1.42699 Δt,
 0.2041 - 0.112232 Δt, 0.869532 - 0.0810106 Δt, 0.209075 + 0.0389451 Δt,
 0.221281 - 0.112232 Δt, 0.415592 + 0.408794 Δt, 0.598776 - 0.26508 Δt,
 0.91903 - 0.112232 Δt, 0.0748652 + 0.501992 Δt, 0.824325 - 0.0693181 Δt,
 0.0585614 - 0.112232 Δt, 0.4362 + 0.741091 Δt, 0.434674 - 0.583384 Δt,
 0.250115 + 1.50634 Δt, 0.0151225 - 1.15593 Δt, 0.71593 + 0.414318 Δt,
 0.408774 - 0.365748 Δt, 0.793994 - 0.275681 Δt, 0.943991 - 0.706214 Δt,
 0.711386 - 0.0677737 Δt, 0.455652 + 0.379378 Δt, 0.0252188 - 0.0555886 Δt}

```

Performing variational imaginary time

We simply repeat these steps to perform variational imaginary time minimization. We'll measure and record the energy at each iteration.

```

vθ = Table[θ[i] → RandomReal[], {i, nθ}];

Δt = .01;
nt = 100;

en = Table[
    (* compute imaginary time tensor *)
    CalcQuregDerivs[u, inψ, vθ, dψ];
    m = Re @ CalcInnerProducts[dψ];

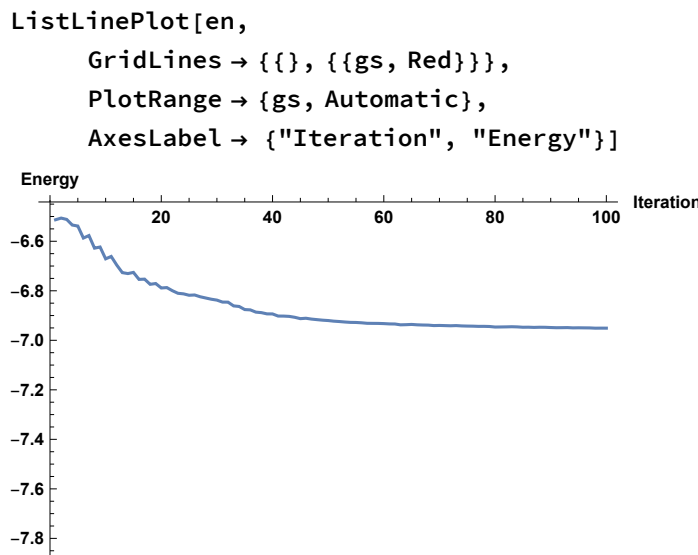
    (* compute energy gradient *)
    ApplyCircuit[u /. vθ, CloneQureg[ψ, inψ]];
    ApplyPauliSum[ψ, h, hψ];
    v = Re @ CalcInnerProducts[hψ, dψ];

    (* update parameters under imaginary time *)
    Δθ = Δt LinearSolve[m, -v];
    vθ[[All, 2]] += Δθ;

    (* record the current energy *)
    CalcExpecPauliSum[ψ, h, hψ],
    (* perform nt iterations *)
    nt
];

```

Here's how the energy evolved



and the final energy, and its corresponding parameters

```
Row[{Last[en], " / ", gs}]
vθ
- 7.27344 / - 7.88074

{θ[1] → 0.310389, θ[2] → 0.0898798, θ[3] → 1.48073, θ[4] → 0.0527977,
θ[5] → 0.801902, θ[6] → 0.262244, θ[7] → 0.722303, θ[8] → 1.53805,
θ[9] → 0.913898, θ[10] → 0.26469, θ[11] → 1.53865, θ[12] → -0.483215,
θ[13] → 0.0282503, θ[14] → 1.54369, θ[15] → 0.687921, θ[16] → 0.584671,
θ[17] → 1.52099, θ[18] → 0.533196, θ[19] → 0.826099, θ[20] → -0.714602,
θ[21] → -0.401971, θ[22] → 0.00317968, θ[23] → -0.189258,
θ[24] → -0.157878, θ[25] → 1.5827, θ[26] → 1.0375, θ[27] → 0.231741}
```