

Quick guide to *Mathematica*

To get help, place the cursor on the function and press **F1**

To run all commands in one cell, place the cursor there and press **Shift + Enter** or **Enter** on the numerical keyboard.

Output of many commands is suppressed by **;** at the end of the command. If you want to see the output, delete **;**

Argument of functions must be in square brackets **[...]**, braces **{...}** are for arrays, ranges etc., use double brackets **[[[...]]** to access elements of arrays

expr /. {x → a} means substitute **a** for **x** in **expr**, **expr // function** means apply function to **expr**, i.e. it is equivalent to **function[expr]**.

D[f[x],x] = derivative of **f[x]** with respect to **x**.

To define a function of **x** one can use **f[x_] := 1 + x^2**, notice the underscore and the colon, or **#** nad **&** like in **f := 1 + #^2 &**

N[expr] evaluates **expr** with machine precision, if you use the decimal point in the expression it will be also evaluated with machine precision

Special characters can be inserted by pressing **Esc ... Esc**, e.g. **Esc p Esc** gives π

Clear[...] is used to unset any variables which could have been assigned previously.

Some other useful commands: Simplify, Expand, Factor; Integrate, Series, Sum

Clear all symbols from previous evaluations to avoid problems

Clear["Global`*"]

Jacobi diagonalization of real symmetric matrix

```
In[1]:= MyJacobiDiagonalization[A_, niter_] :=
Module[{EVal, EVec, SumOffDiag, m, θ, t, s, c, τ, i, p, q, r, x, y, nskip},
  m = Length[A];
  EVal = A;
  EVec = IdentityMatrix[m];
  SumOffDiag = ConstantArray[0.0, niter + 1];
  SumOffDiag[[1]] = Total[Abs[UpperTriangularize[EVal, 1]]^2, 2];
  nskip = 0;
  (* only upper triangular part of the matrix is modified *)
  Do[
    Do[
      Do[
        If[Abs[EVal[[p, q]]] < 10.0^(-16),
          nskip++,
          (* θ = cotg (2φ) = (c^2 - s^2)/2sc *)
          θ = (EVal[[q, q]] - EVal[[p, p]]) / (2 * EVal[[p, q]]);
          (* t = -θ + Sqrt[θ^2 + 1]; *)
          t = 1 / (Abs[θ] + Sqrt[θ^2 + 1]); If[θ < 0.0, t = -t];
          EVal[[p, q]] = s * t;
          EVal[[q, p]] = c * t;
          c = 1 / Sqrt[t^2 + 1];
          s = θ / c;
          τ = c * EVal[[p, p]];
          EVal[[p, p]] = τ + (s * t)^2;
          EVal[[q, q]] = τ + (s * t)^2;
          EVal[[p, q]] = 0;
          EVal[[q, p]] = 0;
          nskip = 0;
        ];
      ];
    ];
  ];
  EVec = Transpose[EVec];
  EVal = DiagonalMatrix[EVal];
  EVec * EVal * Transpose[EVec]
]
```

```

c = 1 / Sqrt[1 + t^2]; s = t * c;
t = s / (1 + c);

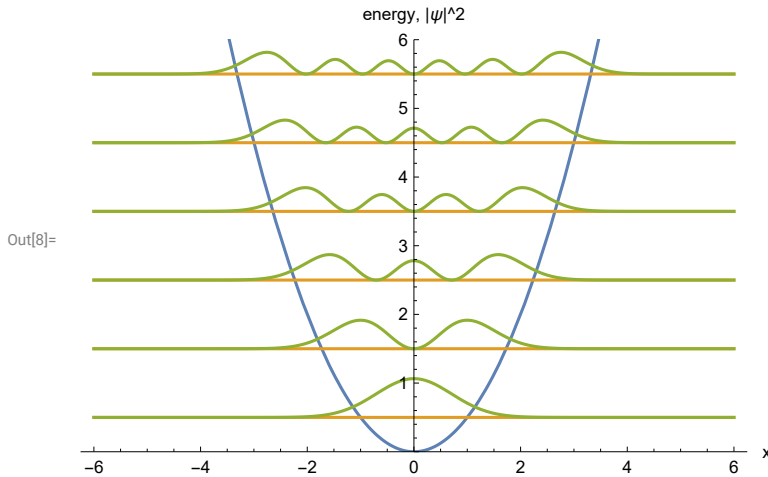
(* diagonal elements *)
EVal[[p, p]] = EVal[[p, p]] - t * EVal[[p, q]];
EVal[[q, q]] = EVal[[q, q]] + t * EVal[[p, q]];
EVal[[p, q]] = 0.0;
Do[ (* elements in columns p and q up to the row p-1 *)
  x = EVal[[r, p]]; y = EVal[[r, q]];
  EVal[[r, p]] = x - s * (y + t * x);
  EVal[[r, q]] = y + s * (x - t * y),
  {r, 1, p - 1}
];
Do[ (* row p and column q between p and q *)
  x = EVal[[p, r]]; y = EVal[[r, q]];
  EVal[[p, r]] = x - s * (y + t * x);
  EVal[[r, q]] = y + s * (x - t * y),
  {r, p + 1, q - 1}
];
Do[ (* rows p and q from q *)
  x = EVal[[p, r]]; y = EVal[[q, r]];
  EVal[[p, r]] = x - s * (y + t * x);
  EVal[[q, r]] = y + s * (x - t * y),
  {r, q + 1, m}
];
(* eigenvectors *)
Do[
  x = EVec[[r, p]]; y = EVec[[r, q]];
  EVec[[r, p]] = x - s * (y + t * x);
  EVec[[r, q]] = y + s * (x - t * y),
  {r, 1, m}
]
],
{q, p + 1, m}
],
{p, 1, m - 1}
];
SumOffDiag[[i + 1]] = Total[Abs[UpperTriangularize[EVal, 1]]^2, 2],
{i, 1, niter}
];
Print["Number of skipped elements in Jacobi diagonalization: ", nskip];
Return[{Diagonal[EVal], EVec, SumOffDiag}]
];

```

Application to quantum 1D harmonic oscillator

Exact energies and eigenfunctions of the 1D harmonic oscillator

```
In[2]:= n = 6;
a = 6;
μ = 1; ω = 1;
V[x_] = 1/2 μ ω^2 x^2;
En[i_] = ω (i + 1/2);
ψ[i_, x_] = 1/Sqrt[2^i i!] Sqrt[μ ω / π] Exp[-μ ω x^2/2] HermiteH[i, x];
Plot[{V[x], Table[En[i], {i, 0, n - 1}], Table[Abs[ψ[i, x]]^2 + En[i], {i, 0, n - 1}]}, {x, -a, a}, PlotRange → {0, 6}, AxesLabel → {"x", "energy, |ψ|^2"}]
```



To find eigenenergies and eigenfunctions of the harmonic oscillator numerically we can use any basis satisfying (at least approximately) boundary conditions. If we are interested in a few eigenfunctions with the lowest energies we can safely set boundary conditions

$$\psi(a) = \psi(-a) = 0 \quad (1)$$

for a certain sufficiently large a .

Construction of the Hamiltonian matrix in the sine basis in which all integrals can be evaluated in the closed form:

```
In[9]:= n = 10;
Clear[H];
ϕ[i_, x_] = Sin[2 π (x + a) i / (4 a)] / Sqrt[a];
KE[i_, j_] = If[i == j,  $\frac{1}{2\mu} \left( \frac{i \pi}{2a} \right)^2$ , 0];
PE[i_, j_] =
  If[i == j, -a^2  $\left( \frac{1}{i^2 \pi^2} - \frac{1}{6} \right)$ , If[Mod[i + j, 2] == 1, 0,  $\frac{4a^2}{\pi^2} \left( \frac{1}{(i-j)^2} - \frac{1}{(i+j)^2} \right)$ ]];
H = ConstantArray[0, {n, n}];
Do[
  Do[
    H[[i, j]] = KE[i, j] + PE[i, j]
    (* Integrate[-ϕ[i,x]D[ϕ[j,x],{x,2}]/(2μ) (* +ϕ[i,x]V[x]ϕ[j,x]*),{x,-a,a}]*) ,
    {j, 1, n}
  ],
  {i, 1, n}
];
N[H] // MatrixForm
```

```
Out[16]//MatrixForm=

$$\begin{pmatrix} 2.3867068486 & 0. & 2.73567195834 & 0. & 0.506605918212 & 0. \\ 0. & 5.22518718612 & 0. & 3.24227787655 & 0. & 0.6839179895 \\ 2.73567195834 & 0. & 5.90314040296 & 0. & 3.41958994793 & 0. \\ 0. & 3.24227787655 & 0. & 6.32033869242 & 0. & 3.501660106 \\ 0.506605918212 & 0. & 3.41958994793 & 0. & 6.71083398871 & 0. \\ 0. & 0.683917989586 & 0. & 3.50166010668 & 0. & 7.132379366 \\ 0.177312071374 & 0. & 0.765988148336 & 0. & 3.54624142748 & 0. \\ 0. & 0.259382230124 & 0. & 0.810569469139 & 0. & 3.573122557 \\ 0.0820701587503 & 0. & 0.303963550927 & 0. & 0.837450599493 & 0. \\ 0. & 0.126651479553 & 0. & 0.330844681281 & 0. & 0.8548974869 \end{pmatrix}$$

```

Diagonalization using the Jacobi method:

```
In[17]:= {Energies, Coefficients, Sums} = MyJacobiDiagonalization[N[H], 4];
Print["Energies:"]
Energies
Print["Coefficients:"]
Coefficients // MatrixForm
Print["Sums of off-diagonal elements after each sweep:"]
Sums
```

Number of skipped elements in Jacobi diagonalization: 100

Energies:

Out[19]=

```
{0.50104958805, 1.50410432891, 14.0574364841, 14.8092333568, 2.58284114737,
3.62555936997, 5.22687945099, 6.31244437983, 8.96803264363, 9.95328641273}
```

Coefficients:

Out[21]//MatrixForm=

$$\begin{pmatrix} 0.743682762039 & 0. & 0.111346166592 & 0. & -0.473403451431 & 0. \\ 0. & 0.498690486113 & 0. & 0.177275037187 & 0. & -0.531557 \\ -0.564989219679 & 0. & 0.326413236198 & 0. & -0.114860190659 & 0. \\ 0. & -0.660227011262 & 0. & 0.351239460359 & 0. & 0.204710 \\ 0.325529735666 & 0. & 0.508923052788 & 0. & 0.596021322138 & 0. \\ 0. & 0.496603599552 & 0. & 0.504563875225 & 0. & 0.446609 \\ -0.141381501882 & 0. & 0.603832964509 & 0. & -0.578913890865 & 0. \\ 0. & -0.249797178391 & 0. & 0.587319592465 & 0. & -0.612981 \\ 0.0420094419831 & 0. & 0.507384969952 & 0. & 0.268911880169 & 0. \\ 0. & 0.0799645373903 & 0. & 0.495656529062 & 0. & 0.316756 \end{pmatrix}$$

Sums of off-diagonal elements after each sweep:

Out[23]=

```
{96.8886114064, 7.66678801731, 0.11407855087, 0.0000259882514481, 3.09543536242 \times 10^{-21}}
```

```
In[24]:= index = Ordering[Energies];
m = 8;
Plot[{V[x], Table[En[i], {i, 0, m - 1}],
Table[Abs[\psi[i, x]]^2 + En[i], {i, 0, m - 1}], Table[Energies[[index[[i]]]], {i, 1, m}],
Table[Abs[Sum[Coefficients[[j, index[[i]]]] \phi[j, x], {j, 1, n}]]^2 + Energies[[index[[i]]]],
{i, 1, m}]}, {x, -a, a}, PlotRange \rightarrow {0, m + 3}, AxesLabel \rightarrow {"x", "energy, |\psi|^2"}]
```

Out[26]=

