

1D-FEM in Fortran: Steady State Heat Conduction

Kengo Nakajima
Information Technology Center

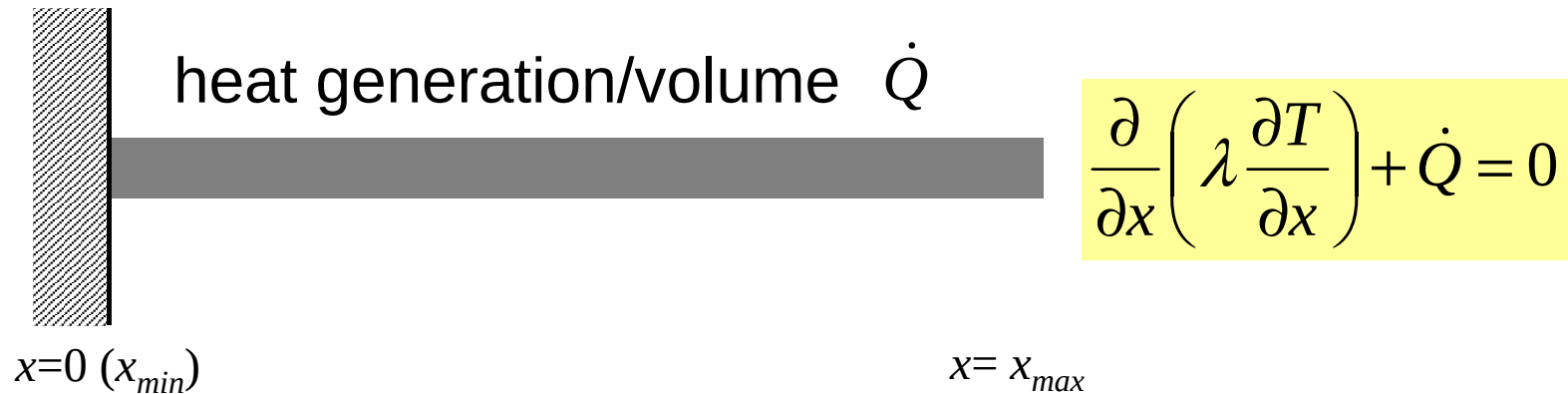
Programming for Parallel Computing (616-2057)
Seminar on Advanced Computing (616-4009)

- 1D-code for Steady State Heat Conduction Problems by Galerkin FEM
- Sparse Linear Solver
 - Conjugate Gradient Method
 - Preconditioning
- Storage of Sparse Matrices
- Program

Keywords

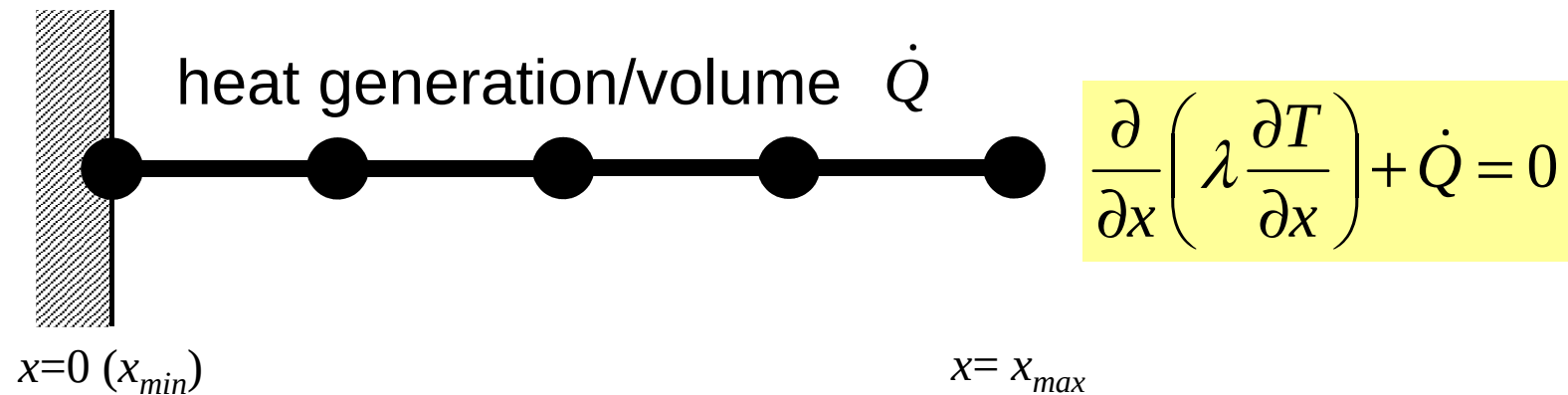
- 1D Steady State Heat Conduction Problems
- Galerkin Method
- Linear Element
- Preconditioned Conjugate Gradient Method

1D Steady State Heat Conduction



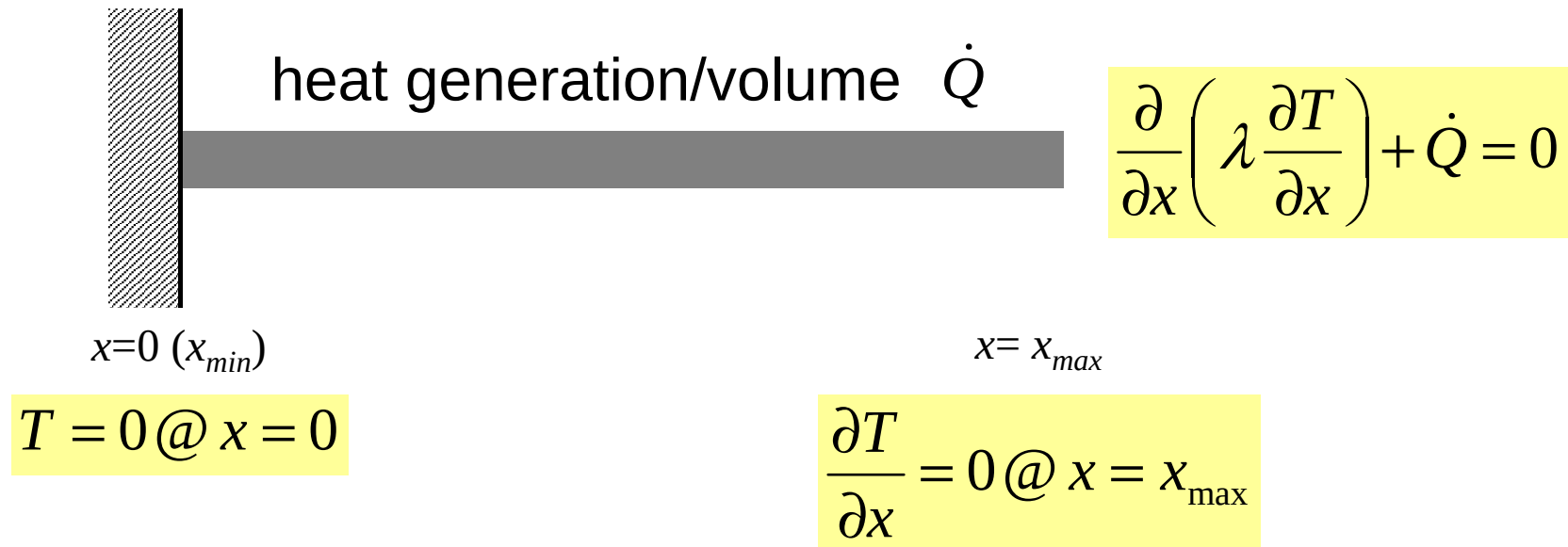
- Uniform: Sectional Area: A , Thermal Conductivity: λ
- Heat Generation Rate/Volume/Time [$QL^{-3}T^{-1}$] $\dot{Q}$
- Boundary Conditions
 - $x=0$: $T=0$ (Fixed Temperature)
 - $x=x_{max}$: $\frac{\partial T}{\partial x} = 0$ (Insulated)

1D Steady State Heat Conduction



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Analytical Solution



$$\lambda T'' = -\dot{Q}$$

$$\lambda T' = -\dot{Q}x + C_1 \Rightarrow C_1 = \dot{Q}x_{max}, \quad T' = 0 @ x = x_{max}$$

$$\lambda T = -\frac{1}{2}\dot{Q}x^2 + C_1x + C_2 \Rightarrow C_2 = 0, \quad T = 0 @ x = 0$$

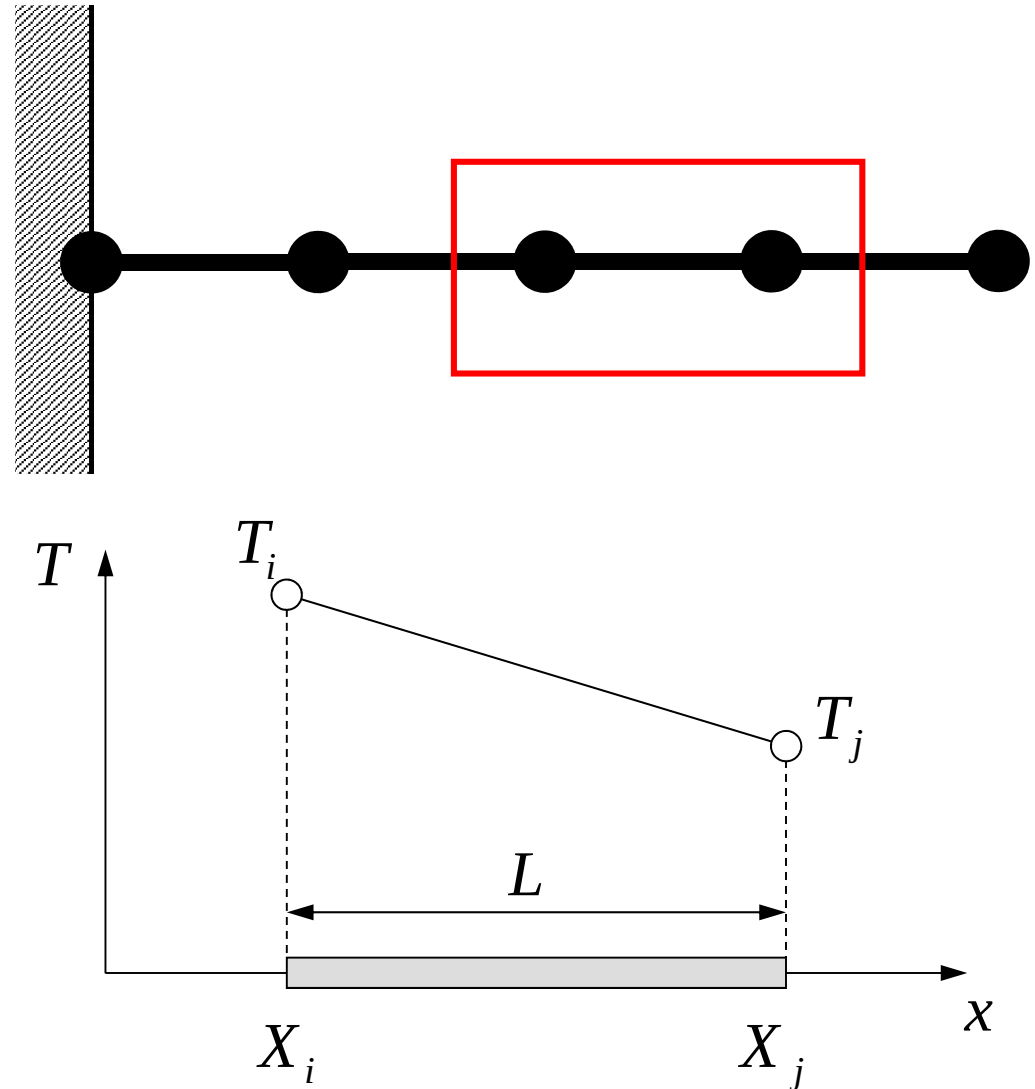
$$\therefore T = -\frac{1}{2\lambda}\dot{Q}x^2 + \frac{\dot{Q}x_{max}}{\lambda}x$$

1D Linear Element (1/4)

一次元線形要素

- 1D Linear Element
 - Length = L
 - Node (Vertex) (節点)
 - Element (要素)
 - T_i Temperature at i
 - T_j Temperature at j
 - Temperature T on each element is linear function of x (Piecewise Linear):

$$T = \alpha_1 + \alpha_2 x$$

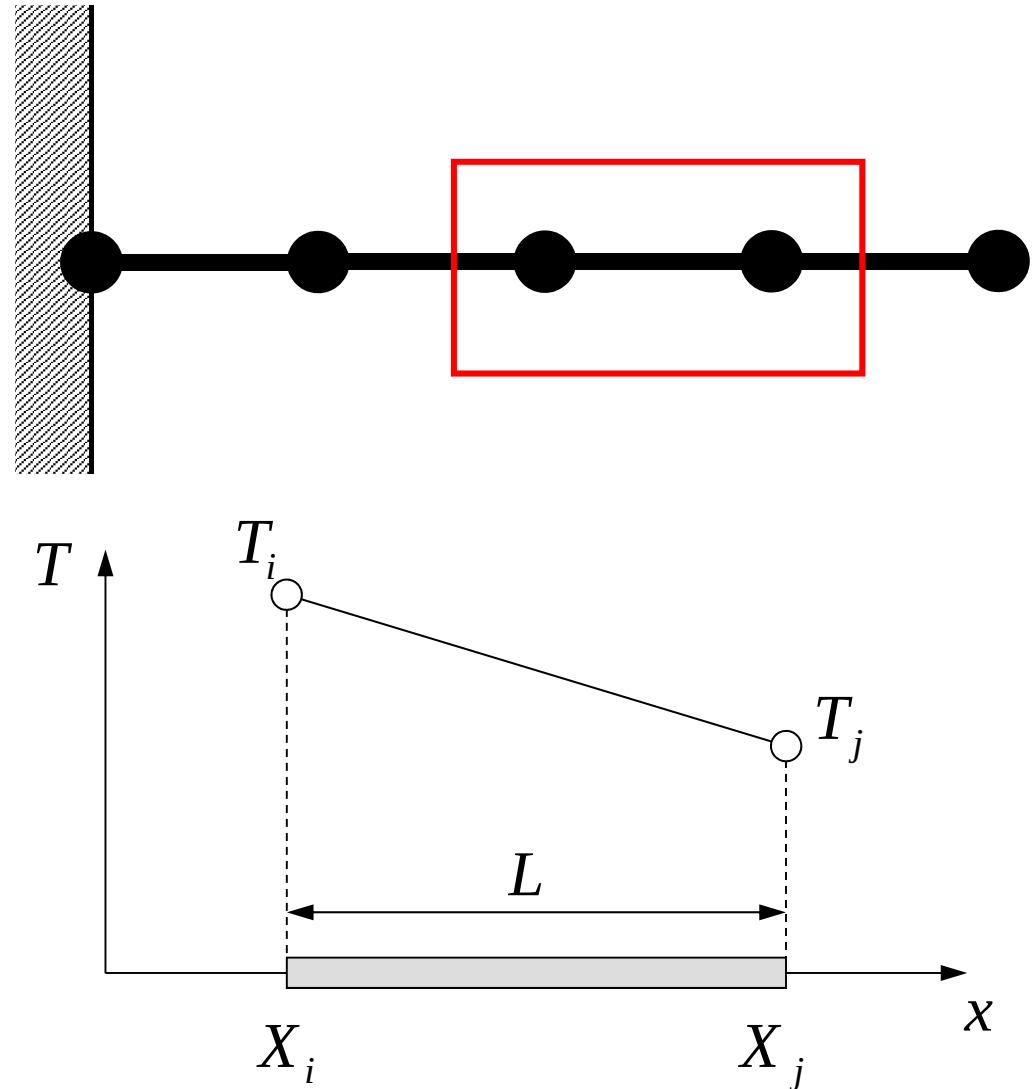


1D Linear Element (1/4)

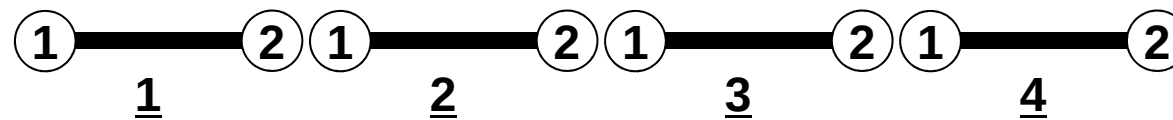
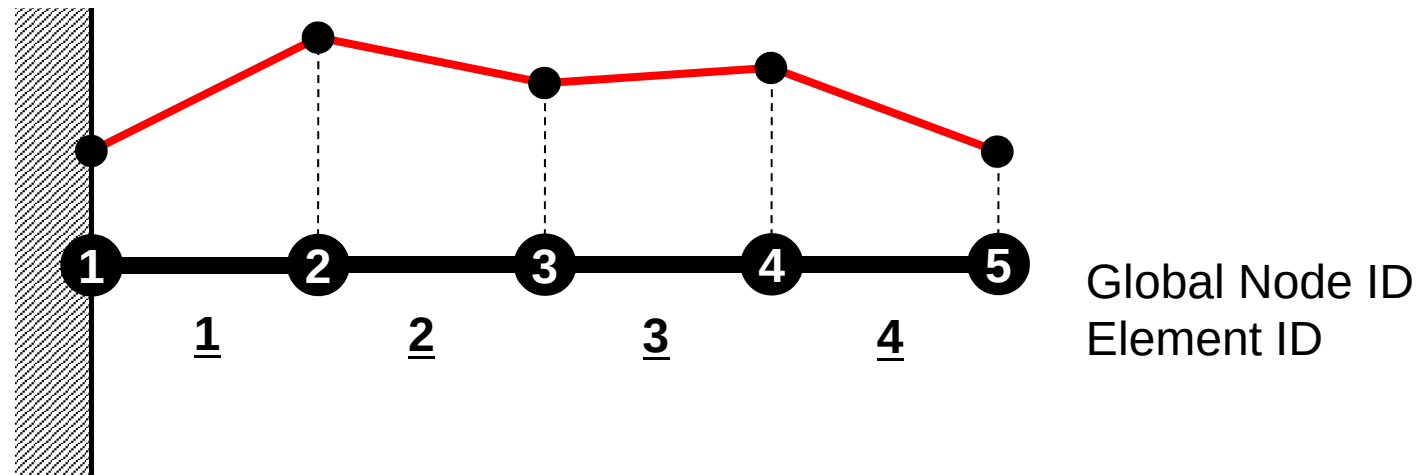
一次元線形要素

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Piecewise Linear



Gradient of temperature is constant in each element (might be discontinuous at each “node”)

1D Linear Elem.: Shape Function (2/4)

- Coef's are calculated based on info. at each node

$$T = T_i @ x = X_i, \quad T = T_j @ x = X_j$$

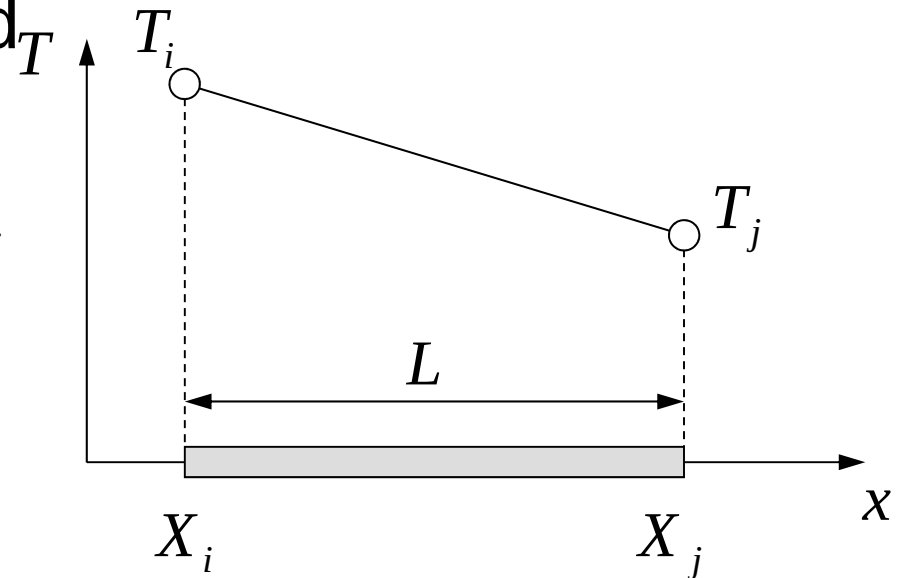
$$T_i = \alpha_1 + \alpha_2 X_i, \quad T_j = \alpha_1 + \alpha_2 X_j$$

- Coefficients:

$$\alpha_1 = \frac{T_i X_j - T_j X_i}{L}, \quad \alpha_2 = \frac{T_j - T_i}{L}$$

- T can be written as follows, according to T_i and T_j :

$$T = \underbrace{\left(\frac{X_j - x}{L} \right)}_{N_i} T_i + \underbrace{\left(\frac{x - X_i}{L} \right)}_{N_j} T_j$$



N_i, N_j

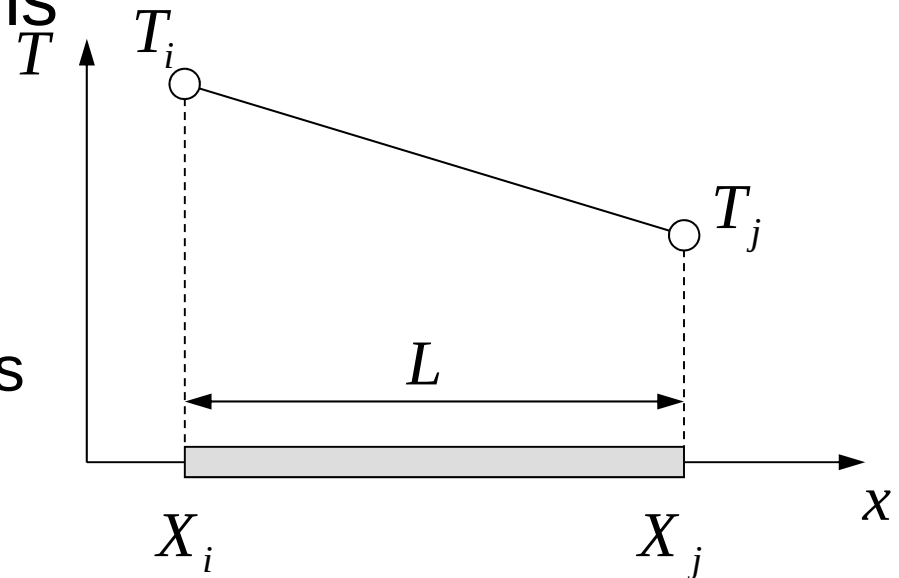
Shape Function (形状関数) or
Interpolation Function (内挿関数), function of x (only)

1D Linear Elem.: Shape Function (3/4)

- Number of Shape Functions = Number of Vertices of Each Element

- N_i : Function of Position
- A kind of Test/Trial Functions

$$N_i = \left(\frac{X_j - x}{L} \right), \quad N_j = \left(\frac{x - X_i}{L} \right)$$



- Linear combination of shape functions provides temperature “in” each element
 - Coef’s (unknowns): Temperature at each node

$$T = N_i T_i + N_j T_j \longleftrightarrow$$

$$T_M = \sum_{i=1}^M a_i \Psi_i$$

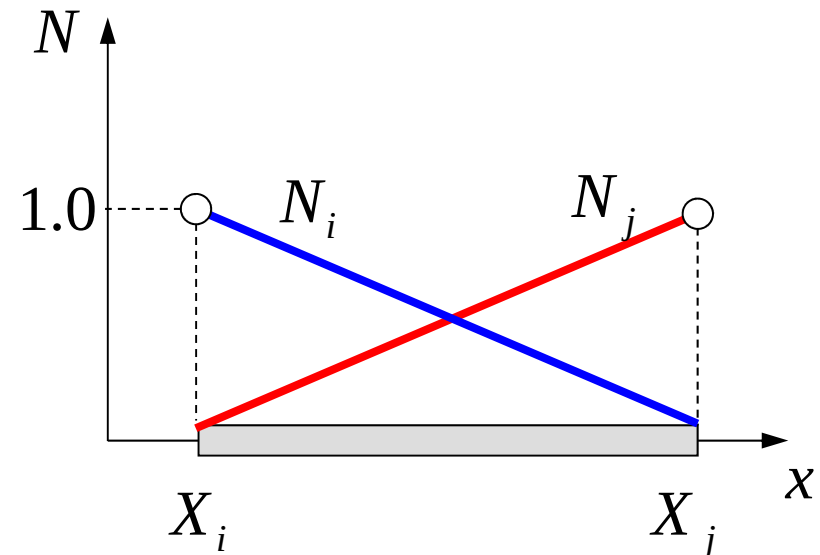
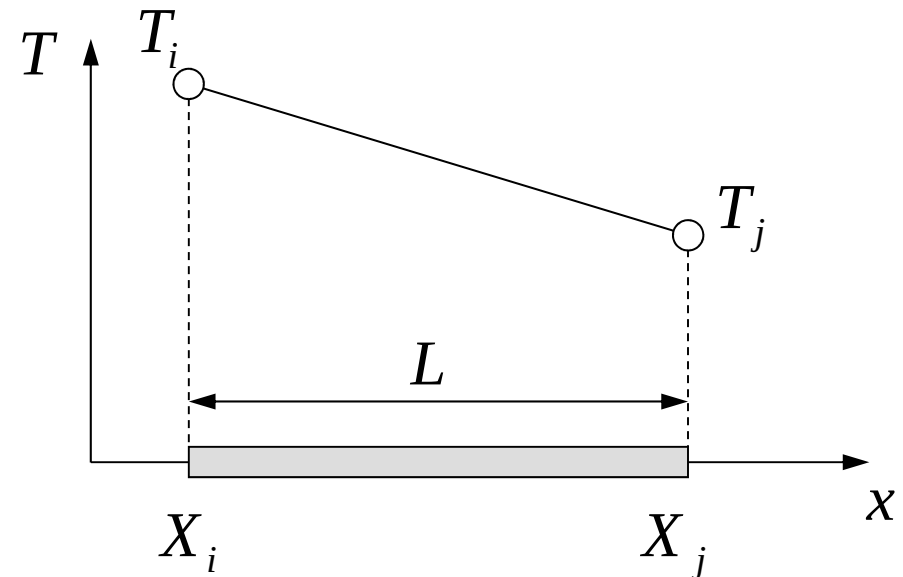
Ψ_i Trial/Test Function (known function of position, defined in domain and at boundary. “Basis” in linear algebra.

a_i Coefficients (unknown)

1D Linear Elem.: Shape Function (4/4)

- Value of N_i
 - =1 at one of the nodes in element
 - =0 on other nodes

$$N_i = \left(\frac{X_j - x}{L} \right), \quad N_j = \left(\frac{x - X_i}{L} \right)$$



Galerkin Method (1/4)

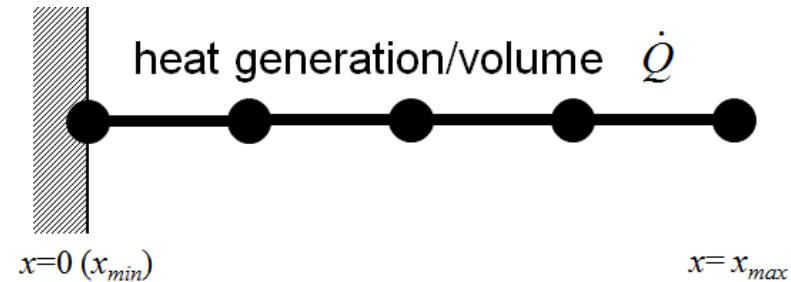
- Governing Equation for 1D Steady State Heat Conduction Problems (Uniform λ):

$$\lambda \left(\frac{d^2 T}{dx^2} \right) + \dot{Q} = 0$$

$T = [N]\{\phi\}$ Distribution of temperature in each element (matrix form), ϕ : Temperature at each node

- Following integral equation is obtained at each element by Galerkin method, where $[N]$'s are also weighting functions:

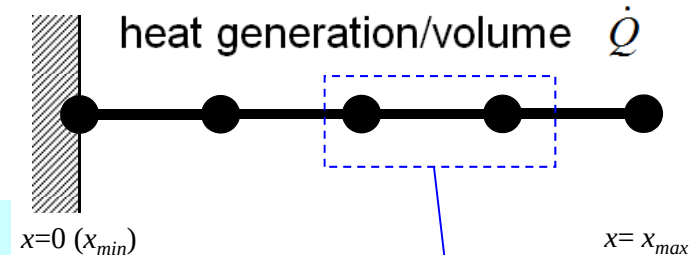
$$\int_V [N]^T \left\{ \lambda \left(\frac{d^2 T}{dx^2} \right) + \dot{Q} \right\} dV = 0$$



Galerkin Method (2/4)

- Green's Theorem (1D)

$$\int_V A \left(\frac{d^2 B}{dx^2} \right) dV = \int_S A \frac{dB}{dx} dS - \int_V \left(\frac{dA}{dx} \frac{dB}{dx} \right) dV$$



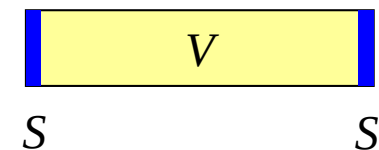
- Apply this to the 1st part of eqn with 2nd-order diff.:

$$\int_V \lambda [N]^T \left(\frac{d^2 T}{dx^2} \right) dV = - \int_V \lambda \left(\frac{d[N]^T}{dx} \frac{dT}{dx} \right) dV + \int_S \lambda [N]^T \frac{dT}{dx} dS$$

- Consider the following terms:

$$T = [N] \{\phi\}, \quad \frac{dT}{dx} = \frac{d[N]}{dx} \{\phi\} \quad \bar{q} = -\lambda \frac{dT}{dx}$$

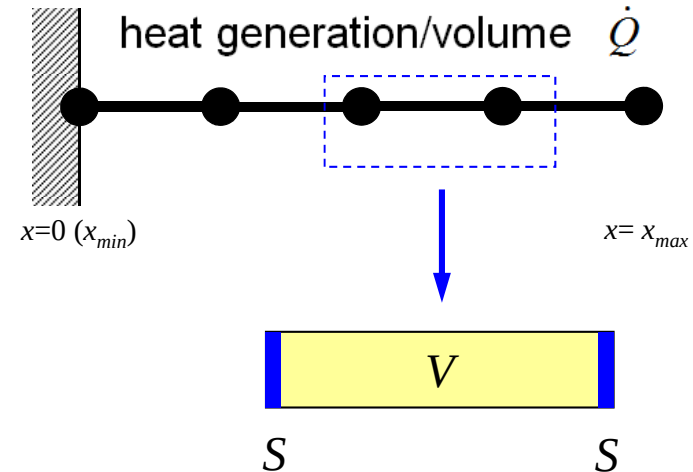
: Heat flux at element surface [$QL^{-2}T^{-1}$]



Galerkin Method (3/4)

- Finally, following eqn is obtained by considering heat generation term $\dot{Q}$:

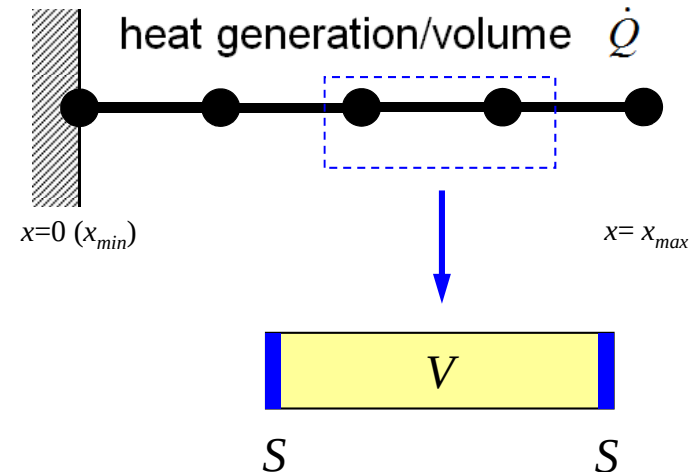
$$-\int_V \lambda \left(\frac{d[N]^T}{dx} \frac{d[N]}{dx} \right) dV \cdot \{\phi\} - \int_S \bar{q} [N]^T dS + \int_V Q [N]^T dV = 0$$



- This is called “weak form (弱形式)”. Original PDE consists of terms with 2nd-order diff., but this “weak form” only includes 1st-order diff by Green’s theorem.
 - Requirements for shape functions are “weaker” in “weak form”. Linear functions can describe effects of 2nd-order differentiation.

Galerkin Method (4/4)

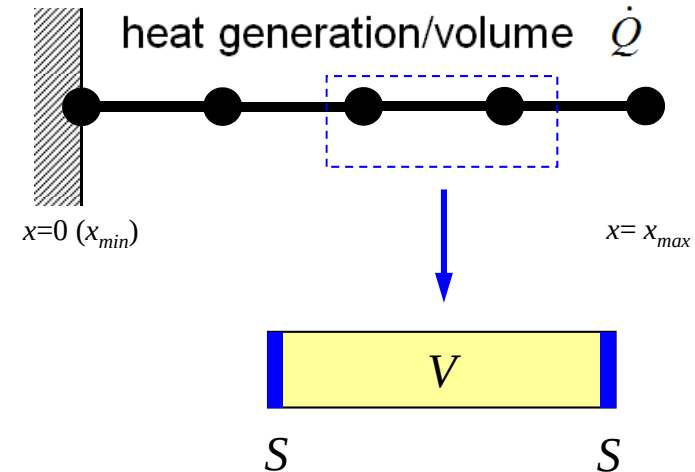
$$\begin{aligned}
 & - \int_V \lambda \left(\frac{d[N]^T}{dx} \frac{d[N]}{dx} \right) dV \cdot \{\phi\} \\
 & \boxed{- \int_S \bar{q} [N]^T dS} + \int_V \dot{Q} [N]^T dV = 0
 \end{aligned}$$



- These terms coincide at element boundaries and disappear. Finally, only terms on the domain boundaries remain.

Weak Form and Boundary Conditions

- Value of dependent variable is defined (Dirichlet)
 - Weighting Function = 0
 - Principal B.C. (Boundary Condition) (第一種境界条件)
 - Essential B.C. (基本境界条件)
- Derivatives of Unknowns (Neumann)
 - Naturally satisfied in weak form
 - Secondary B.C. (第二種境界条件)
 - Natural B.C (自然境界条件)



$$\begin{aligned}
 & - \int_V \lambda \left(\frac{d[N]^T}{dx} \frac{d[N]}{dx} \right) dV \cdot \{\phi\} \\
 & - \int_S \bar{q} [N]^T dS + \int_V \dot{Q} [N]^T dV = 0
 \end{aligned}$$

where $\bar{q} = -\lambda \frac{dT}{dx}$

Weak Form with B.C.: on each elem.

$$[k]^{(e)} \{\phi\}^{(e)} = \{f\}^{(e)}$$

$$[k]^{(e)} = \int_V \lambda \left(\frac{d[N]^T}{dx} \frac{d[N]}{dx} \right) dV$$

$$[f]^{(e)} = \int_V \dot{Q}[N]^T dV - \int_S \bar{q}[N]^T dS$$

Integration over Each Element: $[k]$

$$N_i = \left(\frac{X_j - x}{L} \right), \quad N_j = \left(\frac{x - X_i}{L} \right)$$

$$\frac{dN_i}{dx} = \left(\frac{-1}{L} \right), \quad \frac{dN_j}{dx} = \left(\frac{1}{L} \right)$$

$$\int_V \lambda \left(\frac{d[N]^T}{dx} \frac{d[N]}{dx} \right) dV$$

$$= \lambda \int_0^L \begin{bmatrix} -1/L \\ 1/L \end{bmatrix} \begin{bmatrix} -1/L & 1/L \end{bmatrix} A dx$$

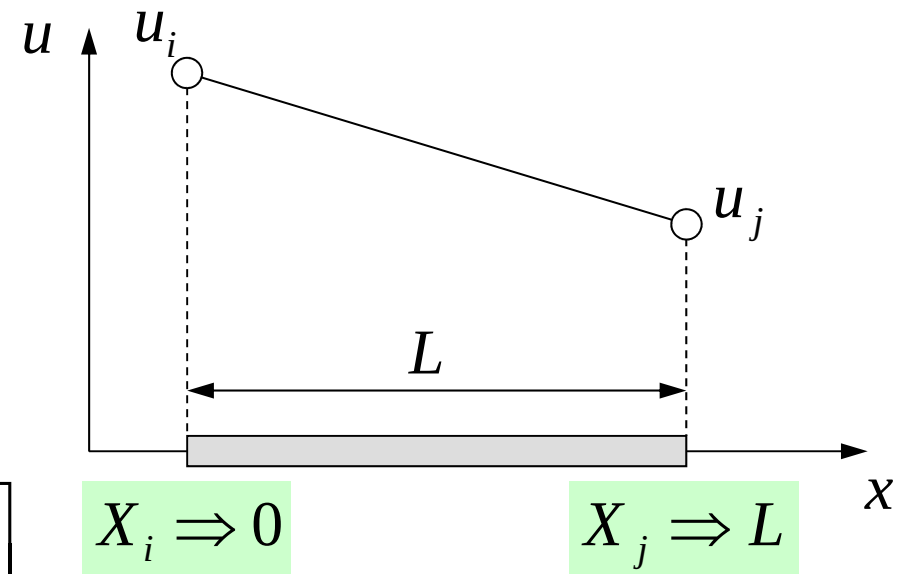
2x1 matrix

1x2 matrix

$$= \frac{\lambda A}{L^2} \int_0^L \begin{bmatrix} +1 & -1 \\ -1 & +1 \end{bmatrix} dx = \frac{\lambda A}{L} \begin{bmatrix} +1 & -1 \\ -1 & +1 \end{bmatrix}$$

A: Sectional Area

L: Length



$$N_i = \left(1 - \frac{x}{L} \right), \quad N_j = \left(\frac{x}{L} \right)$$

Integration over Each Element: $\{f\}$ (1/2)

$$N_i = \left(\frac{X_j - x}{L} \right), \quad N_j = \left(\frac{x - X_i}{L} \right) \quad \frac{dN_i}{dx} = \left(\frac{-1}{L} \right), \quad \frac{dN_j}{dx} = \left(\frac{1}{L} \right)$$

$$N_i = \left(1 - \frac{x}{L} \right), \quad N_j = \left(\frac{x}{L} \right)$$

$$\int_V \dot{Q} [N]^T dV = \dot{Q} A \int_0^L \begin{bmatrix} 1 - x/L \\ x/L \end{bmatrix} dx = \frac{\dot{Q} A L}{2} \begin{Bmatrix} 1 \\ 1 \end{Bmatrix}$$

Heat Generation
(Volume)



A: Sectional Area

L: Length

Integration over Each Element: $\{f\}$ (2/2)

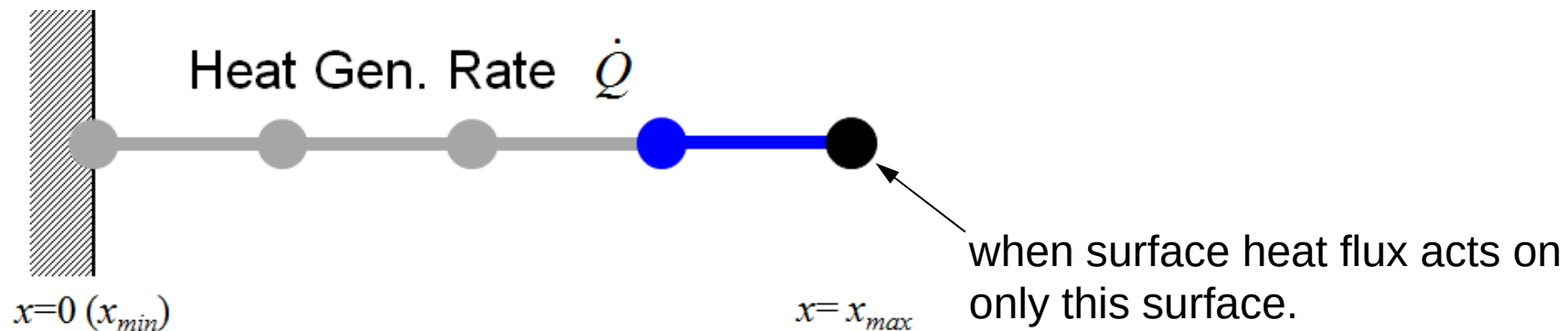
$$N_i = \left(\frac{X_j - x}{L} \right), \quad N_j = \left(\frac{x - X_i}{L} \right) \quad \frac{dN_i}{dx} = \left(\frac{-1}{L} \right), \quad \frac{dN_j}{dx} = \left(\frac{1}{L} \right)$$

$$\int_V \dot{Q} [N]^T dV = \dot{Q} A \int_0^L \begin{bmatrix} 1 - x/L \\ x/L \end{bmatrix} dx = \frac{\dot{Q} A L}{2} \begin{Bmatrix} 1 \\ 1 \end{Bmatrix}$$

Heat Generation
(Volume)

$$\int_S \bar{q} [N]^T dS = \bar{q} A \Big|_{x=L} = \bar{q} A \begin{Bmatrix} 0 \\ 1 \end{Bmatrix}, \quad \bar{q} = -\lambda \frac{dT}{dx}$$

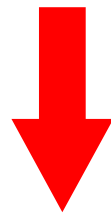
Surface Heat Flux



Global Equations

- Accumulate Element Equations:

$$[k]^{(e)} \{\phi\}^{(e)} = \{f\}^{(e)} \quad \text{Element Matrix, Element Equations}$$



$$[K] \cdot \{\Phi\} = \{F\} \quad \text{Global Matrix, Global Equations}$$

$$[K] = \sum [k], \quad \{F\} = \sum \{f\}$$

$$\{\Phi\}: \text{global vector of } \{\phi\}$$

This is the final linear equations
(global equations) to be solved.

ECSS2016 System

Creating Directory

```
>$ cd Documents  
>$ mkdir 2016Winter your favorite name  
>$ cd 2016Winter
```

This is your “top” directory, and is called **<\$E-TOP>** in this class.

1D Code for Steady-State Heat Conduction Problems

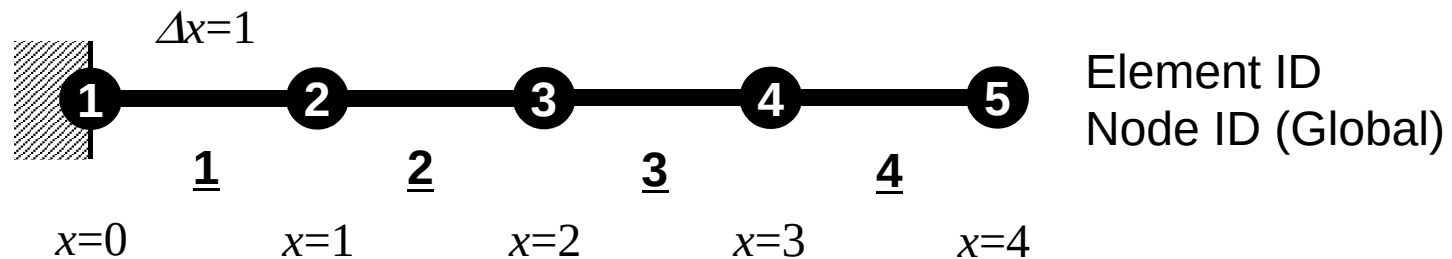
```
>$ cd <$E-TOP>  
>$ (download http://nkl.cc.u-tokyo.ac.jp/files/fem-f.tar)  
>$ tar xvf fem-f.tar  
>$ cd fem-f/1d
```

Compile & GO !

```
>$ cd <$E-TOP>/fem-f/1d
>$ gfortran -O 1d.f
>$ ./a.out
```

Control Data input.dat

4	NE (Number of Elements)
1.0 1.0 1.0 1.0	Δx (Length of Each Elem.: $\mathbf{L}$) , $\mathbf{Q}$, $\mathbf{A}$, λ
100	Number of MAX. Iterations for CG Solver
1.e-8	Convergence Criteria for CG Solver



Results

```
>$ ./a.out
```

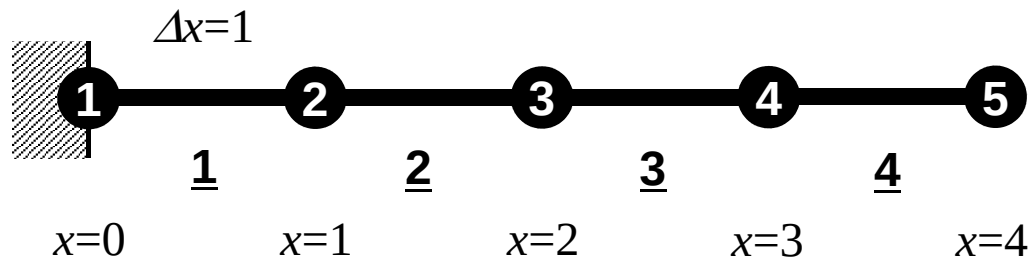
```
4 iters, RESID= 4.154074e-17
```

```
### TEMPERATURE
```

1	0.000000E+00	0.000000E+00
2	3.500000E+00	3.500000E+00
3	6.000000E+00	6.000000E+00
4	7.500000E+00	7.500000E+00
5	8.000000E+00	8.000000E+00

Computational

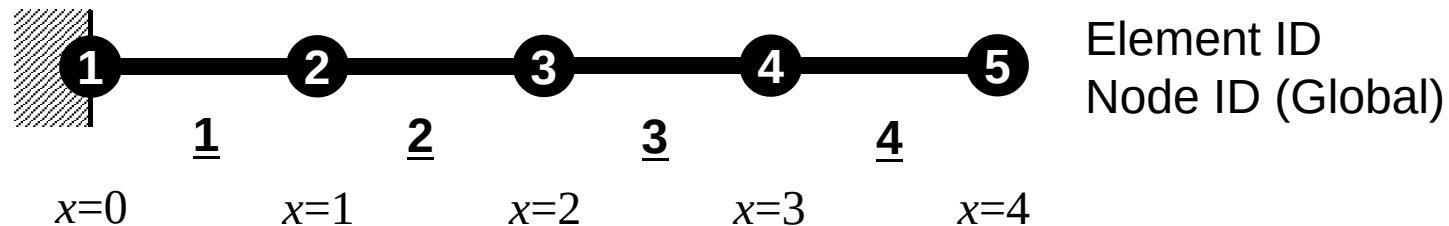
Analytical



Element ID
Node ID (Global)

Element Eqn's/Accumulation (1/3)

- 4 elements, 5 nodes



- $[k]$ and $\{f\}$ of Element-1:

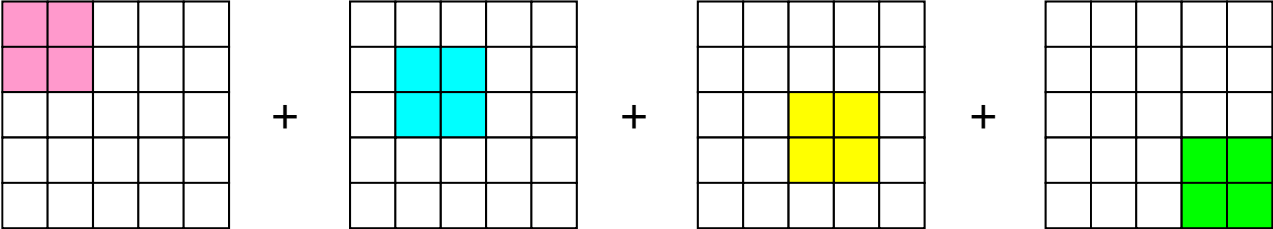
$$[k]^{(1)} = \frac{\lambda A}{L} \begin{bmatrix} +1 & -1 \\ -1 & +1 \end{bmatrix} \quad \{f\}^{(1)} = \frac{\dot{Q}AL}{2} \begin{Bmatrix} 1 \\ 1 \end{Bmatrix}$$

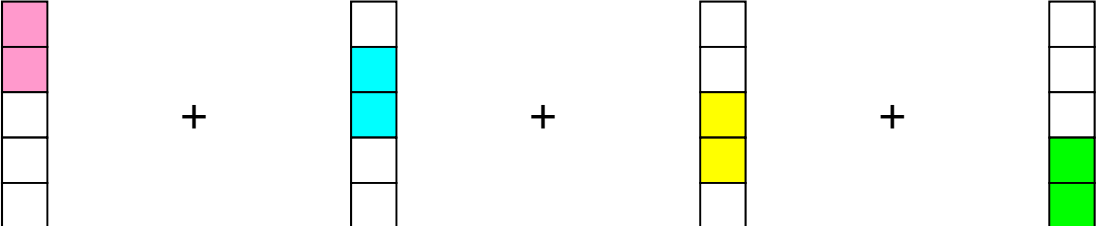
- As for Element-4:

$$[k]^{(4)} = \frac{\lambda A}{L} \begin{bmatrix} +1 & -1 \\ -1 & +1 \end{bmatrix} \quad \{f\}^{(4)} = \frac{\dot{Q}AL}{2} \begin{Bmatrix} 1 \\ 1 \end{Bmatrix}$$

Element Eqn's/Accumulation (2/3)

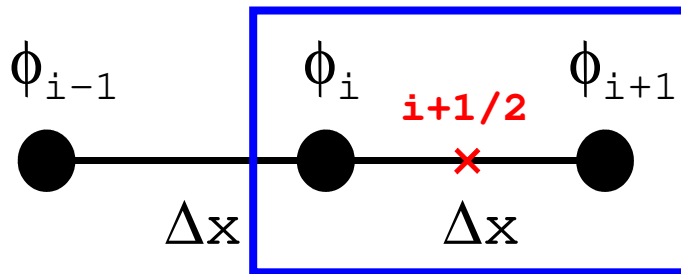
- Element-by-Element Accumulation:

$$[K] = \sum_{e=1}^4 [k]^{(e)} =$$


$$\{F\} = \sum_{e=1}^4 \{f\}^{(e)} =$$


2nd –Order Differentiation in FDM

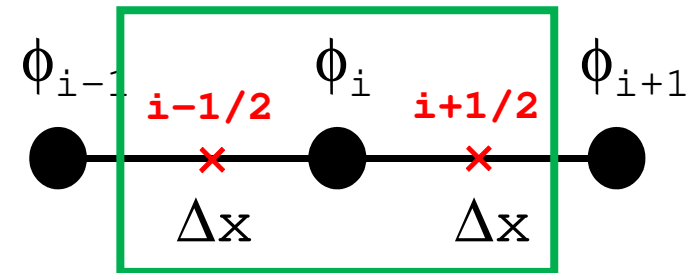
- Approximate Derivative at $\times$ (center of i and $i+1$)



$$\left(\frac{d\phi}{dx} \right)_{i+1/2} \approx \frac{\phi_{i+1} - \phi_i}{\Delta x}$$

$\Delta x \rightarrow 0$: Real Derivative

- 2nd-Order Diff. at i



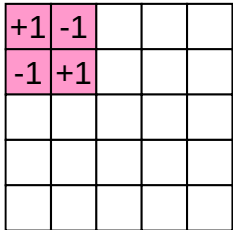
$$\left(\frac{d^2\phi}{dx^2} \right)_i \approx \frac{\left(\frac{d\phi}{dx} \right)_{i+1/2} - \left(\frac{d\phi}{dx} \right)_{i-1/2}}{\Delta x} = \frac{\frac{\phi_{i+1} - \phi_i}{\Delta x} - \frac{\phi_i - \phi_{i-1}}{\Delta x}}{\Delta x} = \frac{\phi_{i+1} - 2\phi_i + \phi_{i-1}}{\Delta x^2}$$

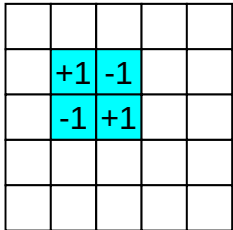
Element-by-Element Operation

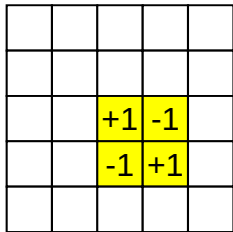
very flexible if each element has different material property, size, etc.

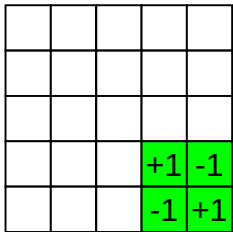
$$[k]^{(e)} = \frac{\lambda^{(e)} A^{(e)}}{L^{(e)}} \begin{bmatrix} +1 & -1 \\ -1 & +1 \end{bmatrix}$$

$$[K] = \sum_{e=1}^4 [k^{(e)}] =$$

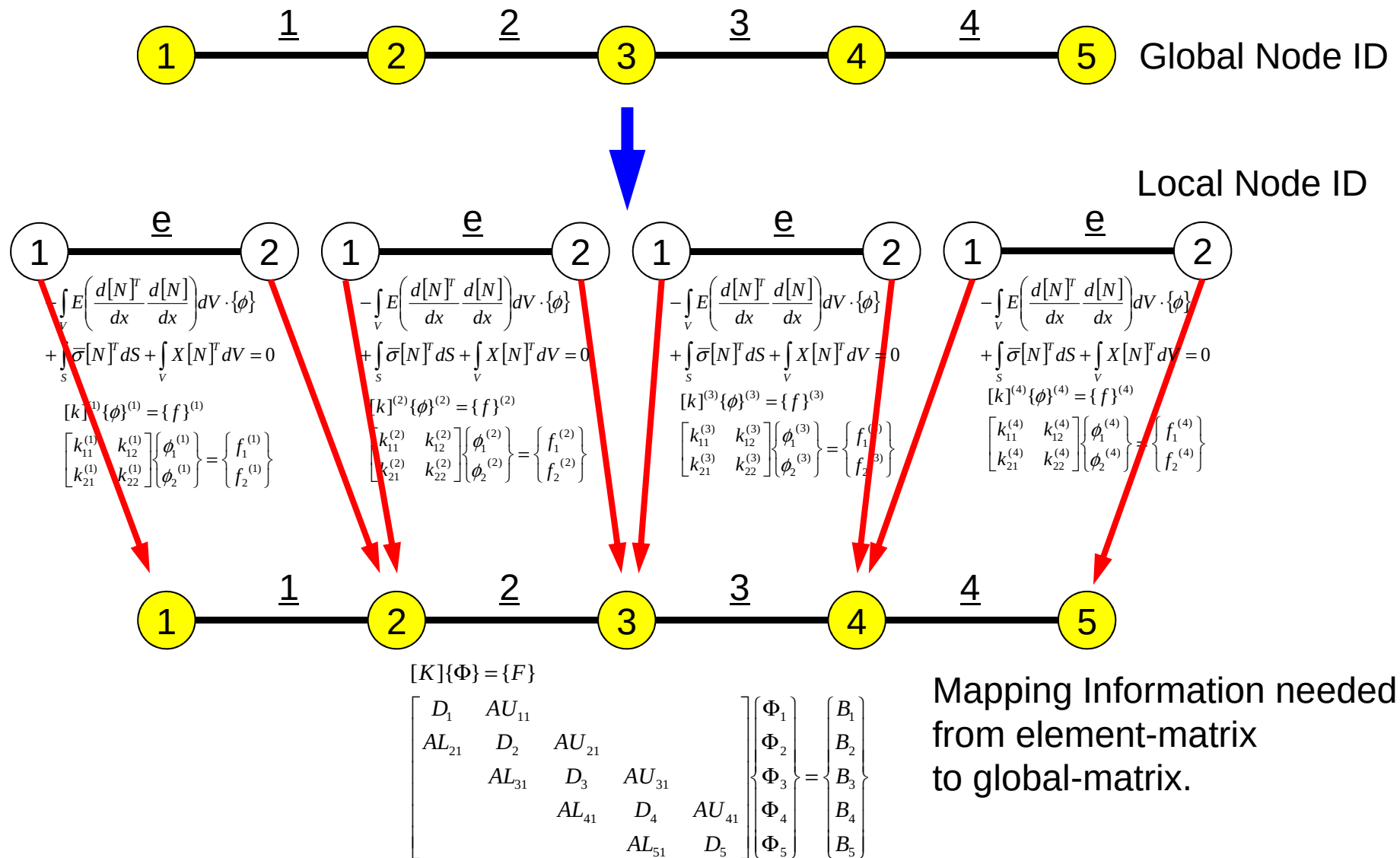

 $\times \frac{\lambda^{(1)} A^{(1)}}{L^{(1)}}$


 $\times \frac{\lambda^{(2)} A^{(2)}}{L^{(2)}}$

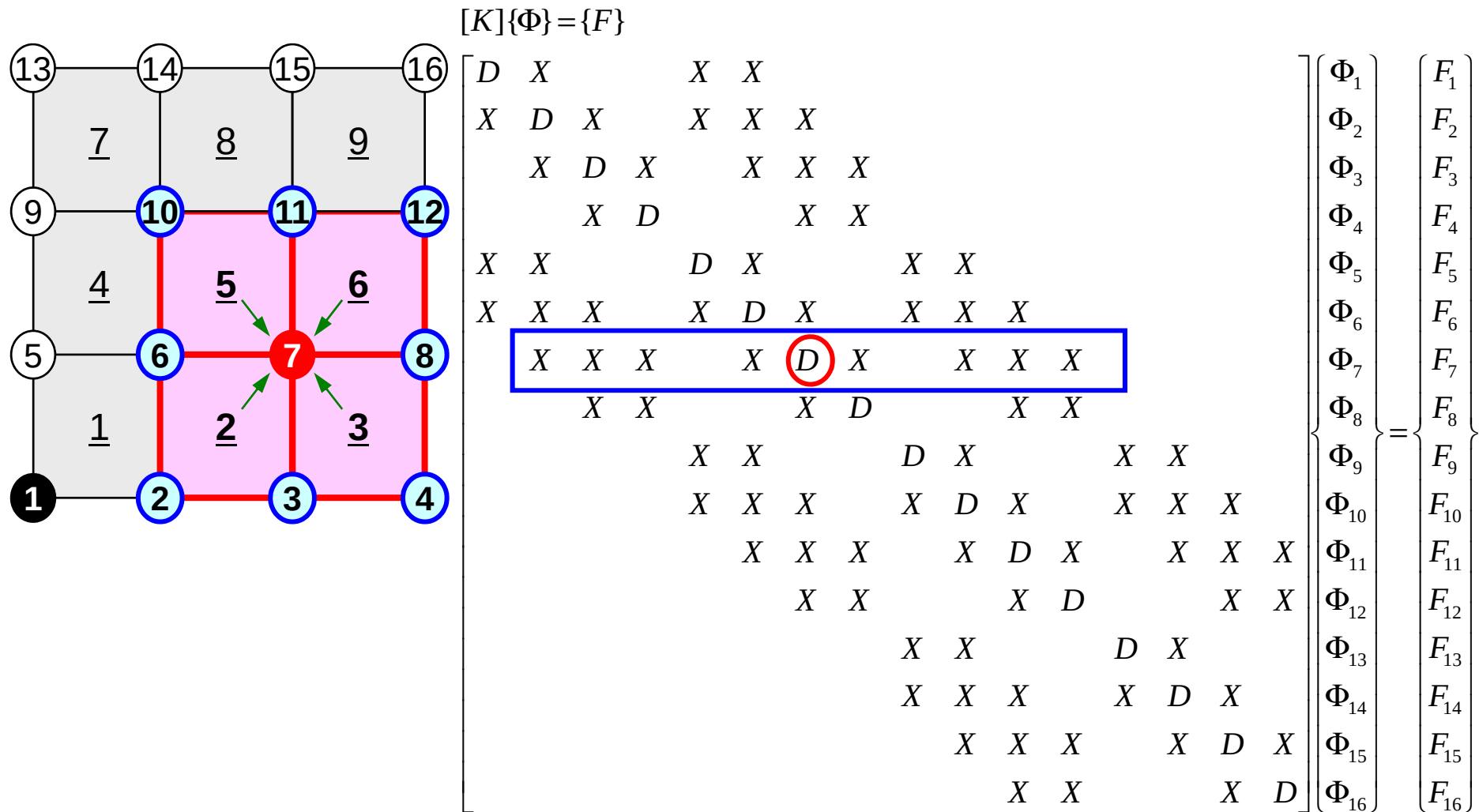

 $\times \frac{\lambda^{(3)} A^{(3)}}{L^{(3)}}$


 $\times \frac{\lambda^{(4)} A^{(4)}}{L^{(4)}}$

Element/Global Operations



Accumulation to Global/overall Matrix



- 1D-code for Static Linear-Elastic Problems by Galerkin FEM
- Sparse Linear Solver
 - Conjugate Gradient Method
 - Preconditioning
- Storage of Sparse Matrices
- Program

Large-Scale Linear Equations in Scientific Applications

- Solving large-scale linear equations $\mathbf{Ax}=\mathbf{b}$ is the most important and expensive part of various types of scientific computing.
 - for both linear and nonlinear applications
- Various types of methods proposed & developed.
 - for dense and sparse matrices
 - classified into direct and iterative methods
- Dense Matrices: 密行列: Globally Coupled Problems
 - BEM, Spectral Methods, MO/MD (gas, liquid)
- Sparse Matrices: 疎行列: Locally Defined Problems
 - **FEM**, FDM, DEM, MD (solid), BEM w/FMM

Direct Method

直接法

- Gaussian Elimination/LU Factorization.
 - compute $\mathbf{A}^{-1}$ directly.

Good

- Robust for wide range of applications.
- Good for both dense and sparse matrices

Bad

- More expensive than iterative methods (memory, CPU)
 - not scalable

What is Iterative Method ?

反復法

Linear Equations
連立一次方程式

$$\begin{pmatrix} a_{11} & a_{12} & \cdots & a_{1n} \\ a_{21} & a_{22} & \cdots & a_{2n} \\ \vdots & \vdots & \ddots & \vdots \\ a_{n1} & a_{n2} & \cdots & a_{nn} \end{pmatrix} \begin{pmatrix} x_1 \\ x_2 \\ \vdots \\ x_n \end{pmatrix} = \begin{pmatrix} b_1 \\ b_2 \\ \vdots \\ b_n \end{pmatrix}$$

A **x** **b**

Initial Solution
初期解

$$\mathbf{x}^{(0)} = \begin{pmatrix} x_1^{(0)} \\ x_2^{(0)} \\ \vdots \\ x_n^{(0)} \end{pmatrix}$$

Starting from a initial vector $\mathbf{x}^{(0)}$, iterative method obtains the final converged solutions by iterations

$$\mathbf{x}^{(1)}, \mathbf{x}^{(2)}, \cdots$$

Iterative Method

反復法

- Stationary Method
 - Only $\mathbf{x}$ (solution vector) changes during iterations.
 - SOR, Gauss-Seidel, Jacobi
 - Generally slow, impractical

$$\mathbf{Ax} = \mathbf{b} \Rightarrow$$

$$\mathbf{x}^{(k+1)} = \mathbf{M}\mathbf{x}^{(k)} + \mathbf{N}\mathbf{b}$$

- Non-Stationary Method
 - With restriction/optimization conditions
 - Krylov-Subspace
 - CG: Conjugate Gradient
 - BiCGSTAB: Bi-Conjugate Gradient Stabilized
 - GMRES: Generalized Minimal Residual

Iterative Method (cont.)

Good

- Less expensive than direct methods, especially in memory.
- Suitable for parallel and vector computing.

Bad

- Convergence strongly depends on problems, boundary conditions (condition number etc.)
- Preconditioning is required : Key Technology for Parallel FEM

Non-Stationary/Krylov Subspace Method (1/2)

非定常法・クリロフ部分空間法

$$\mathbf{Ax} = \mathbf{b} \Rightarrow \mathbf{x} = \mathbf{b} + (\mathbf{I} - \mathbf{A})\mathbf{x}$$

Compute $\mathbf{x}_0, \mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_k$ by the following iterative procedures:

$$\begin{aligned}\mathbf{x}_k &= \mathbf{b} + (\mathbf{I} - \mathbf{A})\mathbf{x}_{k-1} \\ &= (\mathbf{b} - \mathbf{Ax}_{k-1}) + \mathbf{x}_{k-1} \\ &= \mathbf{r}_{k-1} + \mathbf{x}_{k-1}\end{aligned}$$

where $\mathbf{r}_k = \mathbf{b} - \mathbf{Ax}_k$: residual



$$\mathbf{x}_k = \mathbf{x}_0 + \sum_{i=0}^{k-1} \mathbf{r}_i$$

$$\begin{aligned}\mathbf{r}_k &= \mathbf{b} - \mathbf{Ax}_k = \mathbf{b} - \mathbf{A}(\mathbf{r}_{k-1} + \mathbf{x}_{k-1}) \\ &= (\mathbf{b} - \mathbf{Ax}_{k-1}) - \mathbf{Ar}_{k-1} = \mathbf{r}_{k-1} - \mathbf{Ar}_{k-1} = (\mathbf{I} - \mathbf{A})\mathbf{r}_{k-1}\end{aligned}$$

Non-Stationary/Krylov Subspace Method (2/2)

非定常法・クリロフ部分空間法

$$\mathbf{x}_k = \mathbf{x}_0 + \sum_{i=0}^{k-1} \mathbf{r}_i = \mathbf{x}_0 + \mathbf{r}_0 + \sum_{i=0}^{k-2} (\mathbf{I} - \mathbf{A})\mathbf{r}_i = \mathbf{x}_0 + \mathbf{r}_0 + \sum_{i=1}^{k-1} (\mathbf{I} - \mathbf{A})^i \mathbf{r}_0$$

$$\mathbf{z}_k = \mathbf{r}_0 + \sum_{i=1}^{k-1} (\mathbf{I} - \mathbf{A})^i \mathbf{r}_0 = \left[\mathbf{I} + \sum_{i=1}^{k-1} (\mathbf{I} - \mathbf{A})^i \right] \mathbf{r}_0$$



$\mathbf{z}_k$ is a vector which belongs to k^{th} Krylov Subspace (クリロフ部分空間), approximate solution vector $\mathbf{x}_k$ is derived by the Krylov Subspace:

$$\left[\mathbf{r}_0, \mathbf{A}\mathbf{r}_0, \mathbf{A}^2\mathbf{r}_0, \dots, \mathbf{A}^{k-1}\mathbf{r}_0 \right]$$

Conjugate Gradient Method

共役勾配法

- Conjugate Gradient: CG
 - Most popular “non-stationary” iterative method
- for Symmetric Positive Definite (SPD) Matrices
 - 対称正定
 - $\{x\}^T[A]\{x\} > 0$ for arbitrary $\{x\}$
 - All of diagonal components, eigenvalues and leading principal minors > 0 (主小行列式・首座行列式)
 - Matrices of Galerkin-based FEM: heat conduction, Poisson, static linear elastic problems

- Algorithm

- “Steepest Descent Method”
- $\mathbf{x}^{(i)} = \mathbf{x}^{(i-1)} + \alpha_i \mathbf{p}^{(i)}$
 - $\mathbf{x}^{(i)}$: solution, $\mathbf{p}^{(i)}$: search direction, α_i : coefficient
- Solution $\{x\}$ minimizes $\{x-y\}^T[A]\{x-y\}$, where $\{y\}$ is exact solution.

$$\det \begin{bmatrix} a_{11} & a_{12} & a_{13} & a_{14} & \cdots & a_{1n} \\ a_{21} & a_{22} & a_{23} & a_{24} & \cdots & a_{2n} \\ a_{31} & a_{32} & a_{33} & a_{34} & \cdots & a_{3n} \\ a_{41} & a_{42} & a_{43} & a_{44} & \cdots & a_{4n} \\ \vdots & \vdots & \vdots & \vdots & & \vdots \\ a_{n1} & a_{n2} & a_{n3} & a_{n4} & \cdots & a_{nn} \end{bmatrix}$$

Procedures of Conjugate Gradient

```

Compute  $r^{(0)} = b - [A]x^{(0)}$ 
for  $i = 1, 2, \dots$ 
     $z^{(i-1)} = r^{(i-1)}$ 
     $\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$ 
    if  $i = 1$ 
         $p^{(1)} = z^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
    endif
     $q^{(i)} = [A]p^{(i)}$ 
     $\alpha_i = \rho_{i-1} / p^{(i)} \cdot q^{(i)}$ 
     $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
     $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
    check convergence  $|r|$ 
end

```

- Mat-Vec. Multiplication
- Dot Products
- DAXPY (Double Precision: $a\{X\} + \{Y\}$)

$x^{(i)}$: Vector

α_i : Scalar

Procedures of Conjugate Gradient

```

Compute  $r^{(0)} = b - [A]x^{(0)}$ 
for i= 1, 2, ...
     $z^{(i-1)} = r^{(i-1)}$ 
     $\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$ 
    if i=1
         $p^{(1)} = z^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
    endif
     $q^{(i)} = [A]p^{(i)}$ 
     $\alpha_i = \rho_{i-1} / p^{(i)} \cdot q^{(i)}$ 
     $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
     $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
    check convergence  $|r|$ 
end

```

- Mat-Vec. Multiplication
- Dot Products
- DAXPY

$x^{(i)}$: Vector

α_i : Scalar

Procedures of Conjugate Gradient

```

Compute  $r^{(0)} = b - [A]x^{(0)}$ 
for  $i = 1, 2, \dots$ 
     $z^{(i-1)} = r^{(i-1)}$ 
     $\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$ 
    if  $i = 1$ 
         $p^{(1)} = z^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
    endif
     $q^{(i)} = [A]p^{(i)}$ 
     $\alpha_i = \rho_{i-1} / (p^{(i)} \cdot q^{(i)})$ 
     $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
     $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
    check convergence  $|r|$ 
end

```

- Mat-Vec. Multiplication
- Dot Products
- DAXPY

$x^{(i)}$: Vector

α_i : Scalar

Procedures of Conjugate Gradient

```

Compute  $r^{(0)} = b - [A]x^{(0)}$ 
for i= 1, 2, ...
     $z^{(i-1)} = r^{(i-1)}$ 
     $\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$ 
    if i=1
         $p^{(1)} = z^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
    endif
     $q^{(i)} = [A]p^{(i)}$ 
     $\alpha_i = \rho_{i-1} / p^{(i)} \cdot q^{(i)}$ 
     $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
     $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
    check convergence  $|r|$ 
end

```

- Mat-Vec. Multiplication
- Dot Products
- DAXPY
 - Double
 - $\{y\} = a\{x\} + \{y\}$

$x^{(i)}$: Vector

α_i : Scalar

Procedures of Conjugate Gradient

```

Compute  $r^{(0)} = b - [A]x^{(0)}$ 
for  $i = 1, 2, \dots$ 
     $z^{(i-1)} = r^{(i-1)}$ 
     $\rho_{i-1} = r^{(i-1)} z^{(i-1)}$ 
    if  $i = 1$ 
         $p^{(1)} = z^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
    endif
     $q^{(i)} = [A]p^{(i)}$ 
     $\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$ 
     $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
     $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
    check convergence  $|r|$ 
end

```

$x^{(i)}$: Vector

α_i : Scalar

Derivation of CG Algorithm (1/5)

Solution x minimizes the following equation if y is the exact solution ($Ay=b$)

$$(x-y)^T [A](x-y)$$

$$\begin{aligned} (x-y)^T [A](x-y) &= (x, Ax) - (y, Ax) - (x, Ay) + (y, Ay) \\ &= (x, Ax) - 2(x, Ay) + (y, Ay) = (x, Ax) - 2(x, b) + \underline{(y, b)} \quad \text{Const.} \end{aligned}$$

Therefore, the solution x minimizes the following $f(x)$:

$$f(x) = \frac{1}{2}(x, Ax) - (x, b)$$

$$f(x+h) = f(x) + (h, Ax-b) + \frac{1}{2}(h, Ah)$$

Arbitrary vector h

$$f(x) = \frac{1}{2}(x, Ax) - (x, b)$$

$$f(x+h) = f(x) + (h, Ax - b) + \frac{1}{2}(h, Ah)$$

Arbitrary vector h

$$\begin{aligned} f(x+h) &= \frac{1}{2}(x+h, A(x+h)) - (x+h, b) \\ &= \frac{1}{2}(x+h, Ax) + \frac{1}{2}(x+h, Ah) - (x, b) - (h, b) \\ &= \frac{1}{2}(x, Ax) + \frac{1}{2}(h, Ax) + \frac{1}{2}(x, Ah) + \frac{1}{2}(h, Ah) - (x, b) - (h, b) \\ &= \frac{1}{2}(x, Ax) - (x, b) + (h, Ax) - (h, b) + \frac{1}{2}(h, Ah) \\ &= f(x) + (h, Ax - b) + \frac{1}{2}(h, Ah) \end{aligned}$$

Derivation of CG Algorithm (2/5)

CG method minimizes $f(x)$ at each iteration.
 Assume that approximate solution: $x^{(0)}$, and
 search direction vector $p^{(k)}$ is defined at k -th iteration.

$$x^{(k+1)} = x^{(k)} + \alpha_k p^{(k)}$$

Minimization of $f(x^{(k+1)})$ is done as follows:

$$f(x^{(k)} + \alpha_k p^{(k)}) = \frac{1}{2} \alpha_k^2 (p^{(k)}, Ap^{(k)}) - \alpha_k (p^{(k)}, b - Ax^{(k)}) + f(x^{(k)})$$

$$\frac{\partial f(x^{(k)} + \alpha_k p^{(k)})}{\partial \alpha_k} = 0 \Rightarrow \alpha_k = \frac{(p^{(k)}, b - Ax^{(k)})}{(p^{(k)}, Ap^{(k)})} = \frac{(p^{(k)}, r^{(k)})}{(p^{(k)}, Ap^{(k)})} \quad \textcolor{red}{(1)}$$

$$r^{(k)} = b - Ax^{(k)} \text{ residual vector}$$

Derivation of CG Algorithm (3/5)

Residual vector at $(k+1)$ -th iteration: $r^{(k+1)} = b - Ax^{(k+1)}, r^{(k)} = b - Ax^{(k)}$

$$r^{(k+1)} = r^{(k)} - \alpha_k Ap^{(k)} \quad \textcolor{red}{(2)} \qquad r^{(k+1)} - r^{(k)} = Ax^{(k+1)} - Ax^{(k)} = \alpha_k Ap^{(k)}$$

Search direction vector p is defined by the following recurrence formula:

$$p^{(k+1)} = r^{(k+1)} + \beta_k p^{(k)}, p^{(0)} = r^{(0)} \quad \textcolor{red}{(3)}$$

It's lucky if we can get exact solution y at $(k+1)$ -th iteration:

$$y = x^{(k+1)} + \alpha_{k+1} p^{(k+1)}$$

Derivation of CG Algorithm (4/5)

BTW, we have the following (convenient) orthogonality relation:

$$\left(Ap^{(k)}, y - x^{(k+1)} \right) = 0$$

$$\begin{aligned} \left(Ap^{(k)}, y - x^{(k+1)} \right) &= \left(p^{(k)}, Ay - Ax^{(k+1)} \right) = \left(p^{(k)}, b - Ax^{(k+1)} \right) \\ &= \left(p^{(k)}, b - A[x^{(k)} + \alpha_k p^{(k)}] \right) = \left(p^{(k)}, b - Ax^{(k)} - \alpha_k Ap^{(k)} \right) \\ &= \left(p^{(k)}, r^{(k)} - \alpha_k Ap^{(k)} \right) = \left(p^{(k)}, r^{(k)} \right) - \alpha_k \left(p^{(k)}, Ap^{(k)} \right) = 0 \end{aligned}$$

$$\therefore \alpha_k = \frac{\left(p^{(k)}, r^{(k)} \right)}{\left(p^{(k)}, Ap^{(k)} \right)}$$

Thus, following relation is obtained:

$$\left(Ap^{(k)}, y - x^{(k+1)} \right) = \left(Ap^{(k)}, \alpha_{k+1} p^{(k+1)} \right) = 0 \Rightarrow \left(p^{(k+1)}, Ap^{(k)} \right) = 0$$

Derivation of CG Algorithm (5/5)

$$\begin{aligned} (p^{(k+1)}, Ap^{(k)}) &= (r^{(k+1)} + \beta_k p^{(k)}, Ap^{(k)}) = (r^{(k+1)}, Ap^{(k)}) + \beta_k (p^{(k)}, Ap^{(k)}) = 0 \\ \Rightarrow \beta_k &= \frac{-(r^{(k+1)}, Ap^{(k)})}{(p^{(k)}, Ap^{(k)})} \quad (4) \end{aligned}$$

$(p^{(k+1)}, Ap^{(k)}) = 0$ $p^{(k)}$ and $p^{(k+1)}$ are “conjugate (共役)” for matrix A

```

Compute  $p^{(0)} = r^{(0)} = b - [A]x^{(0)}$ 
for i= 1, 2, ...
    calc.  $\alpha_{i-1}$ 
     $x^{(i)} = x^{(i-1)} + \alpha_{i-1}p^{(i-1)}$ 
     $r^{(i)} = r^{(i-1)} - \alpha_{i-1}[A]p^{(i-1)}$ 

    check convergence  $|r|$ 
    (if not converged)
    calc.  $\beta_{i-1}$ 
     $p^{(i)} = r^{(i)} + \beta_{i-1}p^{(i-1)}$ 
end
  
```

$$\alpha_{i-1} = \frac{(p^{(i-1)}, r^{(i-1)})}{(p^{(i-1)}, Ap^{(i-1)})}$$

$$\beta_{i-1} = \frac{-(r^{(i)}, Ap^{(i-1)})}{(p^{(i-1)}, Ap^{(i-1)})}$$

Properties of CG Algorithm

Following “conjugate (共役)” relationship is obtained for arbitrary (i,j) :

$$(p^{(i)}, Ap^{(j)}) = 0 \quad (i \neq j)$$

Following relationships are also obtained for $p^{(k)}$ and $r^{(k)}$:

$$(r^{(i)}, r^{(j)}) = 0 \quad (i \neq j), \quad (p^{(k)}, r^{(k)}) = (r^{(k)}, r^{(k)})$$

In N-dimensional space, only N sets of orthogonal and linearly independent residual vector $r^{(k)}$. This means CG method converges after N iterations if number of unknowns is N. Actually, round-off error sometimes affects convergence.

Proof (1/3)

Mathematical Induction

数学的帰納法

$$\begin{aligned} (r^{(i)}, r^{(j)}) &= 0 \quad (i \neq j) \\ (p^{(i)}, Ap^{(j)}) &= 0 \quad (i \neq j) \end{aligned}$$

$$(1) \quad \alpha_k = \frac{(p^{(k)}, r^{(k)})}{(p^{(k)}, Ap^{(k)})}$$

$$(2) \quad r^{(k+1)} = r^{(k)} - \alpha_k Ap^{(k)}$$

$$(3) \quad p^{(k+1)} = r^{(k+1)} + \beta_k p^{(k)}, \quad r^{(0)} = p^{(0)}$$

$$(4) \quad \beta_k = \frac{-(r^{(k+1)}, Ap^{(k)})}{(p^{(k)}, Ap^{(k)})}$$

Proof (2/3)

Mathematical Induction

数学的帰納法

$$\begin{aligned} (r^{(i)}, r^{(j)}) &= 0 \quad (i \neq j) \\ (p^{(i)}, Ap^{(j)}) &= 0 \quad (i \neq j) \end{aligned} \quad (*)$$

$(*)$ is satisfied for $i \leq k, j \leq k$ where $i \neq j$

$$\begin{aligned} \text{if } i < k \quad (r^{(k+1)}, r^{(i)}) &= (r^{(i)}, r^{(k+1)}) \stackrel{(2)}{=} (r^{(i)}, r^{(k)} - \alpha_k Ap^{(k)}) \\ &\stackrel{(*)}{=} -\alpha_k (r^{(i)}, Ap^{(k)}) \stackrel{(4)}{=} -\alpha_k (p^{(i)} - \beta_{i-1} p^{(i-1)}, Ap^{(k)}) \\ &= -\alpha_k (p^{(i)}, Ap^{(k)}) + \alpha_k \beta_{i-1} (p^{(i-1)}, Ap^{(k)}) \stackrel{(*)}{=} 0 \end{aligned}$$

$$\text{if } i = k \quad (r^{(k+1)}, r^{(k)}) \stackrel{(2)}{=} (r^{(k)}, r^{(k)}) - (r^{(k)}, \alpha_k Ap^{(k)})$$

$$(1) \alpha_k = \frac{(p^{(k)}, r^{(k)})}{(p^{(k)}, Ap^{(k)})}$$

$$(2) r^{(k+1)} = r^{(k)} - \alpha_k Ap^{(k)}$$

$$(3) p^{(k+1)} = r^{(k+1)} + \beta_k p^{(k)}$$

$$(4) \beta_k = \frac{-(r^{(k+1)}, Ap^{(k)})}{(p^{(k)}, Ap^{(k)})}$$

$$\stackrel{(3)}{=} (r^{(k)}, r^{(k)}) - (p^{(k)} - \beta_{k-1} p^{(k-1)}, \alpha_k Ap^{(k)})$$

$$\stackrel{(*)}{=} (r^{(k)}, r^{(k)}) - \alpha_k (p^{(k)}, Ap^{(k)}) \stackrel{(1)}{=} (r^{(k)}, r^{(k)}) - (p^{(k)}, r^{(k)})$$

$$\stackrel{(3)}{=} (r^{(k)}, r^{(k)}) - (\beta_{k-1} p^{(k-1)} + r^{(k)}, r^{(k)})$$

$$= -\beta_{k-1} (p^{(k-1)}, r^{(k)}) \stackrel{(2)}{=} -\beta_{k-1} (p^{(k-1)}, r^{(k-1)} - \alpha_{k-1} Ap^{(k-1)})$$

$$= -\beta_{k-1} \{ (p^{(k-1)}, r^{(k-1)}) - \alpha_{k-1} (p^{(k-1)}, Ap^{(k-1)}) \} \stackrel{(1)}{=} 0$$

Proof (3/3)

Mathematical Induction

数学的帰納法

$$\begin{aligned} (r^{(i)}, r^{(j)}) &= 0 \quad (i \neq j) \\ (p^{(i)}, Ap^{(j)}) &= 0 \quad (i \neq j) \end{aligned} \quad (*)$$

$(*)$ is satisfied for $i \leq k, j \leq k$ where $i \neq j$

$$\begin{aligned} \text{if } i < k \quad (p^{(k+1)}, Ap^{(i)}) &\stackrel{(3)}{=} (r^{(k+1)} + \beta_k p^{(k)}, Ap^{(i)}) \\ &\stackrel{(*)}{=} (r^{(k+1)}, Ap^{(i)}) \end{aligned}$$

$$\stackrel{(2)}{=} \frac{1}{\alpha_k} (r^{(k+1)}, r^{(i)} - r^{(i-1)}) = 0$$

$$\begin{aligned} \text{if } i = k \quad (p^{(k+1)}, Ap^{(k)}) &\stackrel{(3)}{=} (r^{(k+1)}, Ap^{(k)}) + \beta_k (p^{(k)}, Ap^{(k)}) \\ &\stackrel{(4)}{=} 0 \end{aligned}$$

$$(1) \alpha_k = \frac{(p^{(k)}, r^{(k)})}{(p^{(k)}, Ap^{(k)})}$$

$$(2) r^{(k+1)} = r^{(k)} - \alpha_k Ap^{(k)}$$

$$(3) p^{(k+1)} = r^{(k+1)} + \beta_k p^{(k)}$$

$$(4) \beta_k = \frac{-(r^{(k+1)}, Ap^{(k)})}{(p^{(k)}, Ap^{(k)})}$$

$$\begin{aligned}
\left(r^{(k+1)}, r^{(k)}\right) &= 0 \\
\left(r^{(k+1)}, r^{(k)}\right) &\stackrel{(2)}{=} \left(r^{(k)}, r^{(k)}\right) - \left(r^{(k)}, \alpha_k A p^{(k)}\right) \\
&\stackrel{(3)}{=} \left(r^{(k)}, r^{(k)}\right) - \left(p^{(k)} - \beta_{k-1} p^{(k-1)}, \alpha_k A p^{(k)}\right) \\
&\stackrel{(*)}{=} \left(r^{(k)}, r^{(k)}\right) - \alpha_k \left(p^{(k)}, A p^{(k)}\right) \stackrel{(1)}{=} \left(r^{(k)}, r^{(k)}\right) - \left(p^{(k)}, r^{(k)}\right) = 0
\end{aligned}$$

$$\therefore \left(r^{(k)}, r^{(k)}\right) = \left(p^{(k)}, r^{(k)}\right)$$

$$(1) \alpha_k = \frac{\left(p^{(k)}, r^{(k)}\right)}{\left(p^{(k)}, A p^{(k)}\right)}$$

$$(2) r^{(k+1)} = r^{(k)} - \alpha_k A p^{(k)}$$

$$(3) p^{(k+1)} = r^{(k+1)} + \beta_k p^{(k)}$$

$$(4) \beta_k = \frac{-\left(r^{(k+1)}, A p^{(k)}\right)}{\left(p^{(k)}, A p^{(k)}\right)}$$

$$\alpha_k, \beta_k$$

Usually, we use simpler definitions of α_k, β_k as follows:

$$\alpha_k = \frac{(p^{(k)}, b - Ax^{(k)})}{(p^{(k)}, Ap^{(k)})} = \frac{(p^{(k)}, r^{(k)})}{(p^{(k)}, Ap^{(k)})} = \frac{(r^{(k)}, r^{(k)})}{(p^{(k)}, Ap^{(k)})}$$

$$\because (p^{(k)}, r^{(k)}) = (r^{(k)}, r^{(k)})$$

$$\beta_k = \frac{-(r^{(k+1)}, Ap^{(k)})}{(p^{(k)}, Ap^{(k)})} = \frac{(r^{(k+1)}, r^{(k+1)})}{(r^{(k)}, r^{(k)})}$$

$$\because (r^{(k+1)}, Ap^{(k)}) = \frac{(r^{(k+1)}, r^{(k)} - r^{(k+1)})}{\alpha_k} = -\frac{(r^{(k+1)}, r^{(k+1)})}{\alpha_k}$$

Procedures of Conjugate Gradient

```

Compute  $r^{(0)} = b - [A]x^{(0)}$ 
for i = 1, 2, ...
     $z^{(i-1)} = r^{(i-1)}$ 
     $\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$ 
    if i = 1
         $p^{(1)} = z^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
    endif
     $q^{(i)} = [A]p^{(i)}$ 
     $\alpha_i = \rho_{i-1} / p^{(i)} \cdot q^{(i)}$ 
     $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
     $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
    check convergence  $|r|$ 
end

```

$x^{(i)}$: Vector

α_i : Scalar

$$\beta_{i-1} = \frac{(r^{(i-1)}, r^{(i-1)})}{(r^{(i-2)}, r^{(i-2)})} \quad (= \rho_{i-1})$$

$$\alpha_i = \frac{(r^{(i-1)}, r^{(i-1)})}{(p^{(i)}, Ap^{(i)})} \quad (= \rho_{i-1})$$

Preconditioning for Iterative Solvers

- Convergence rate of iterative solvers strongly depends on the spectral properties (eigenvalue distribution) of the coefficient matrix $\mathbf{A}$.
 - Eigenvalue distribution is small, eigenvalues are close to 1
 - In “ill-conditioned” problems, “condition number” (ratio of max/min eigenvalue if $\mathbf{A}$ is symmetric) is large (条件数) .
- A preconditioner $\mathbf{M}$ (whose properties are similar to those of $\mathbf{A}$) transforms the linear system into one with more favorable spectral properties (前处理)
 - $\mathbf{M}$ transforms $\mathbf{Ax}=\mathbf{b}$ into $\mathbf{A}'\mathbf{x}=\mathbf{b}'$ where $\mathbf{A}'=\mathbf{M}^{-1}\mathbf{A}$, $\mathbf{b}'=\mathbf{M}^{-1}\mathbf{b}$
 - If $\mathbf{M}\sim\mathbf{A}$, $\mathbf{M}^{-1}\mathbf{A}$ is close to identity matrix.
 - If $\mathbf{M}^{-1}=\mathbf{A}^{-1}$, this is the best preconditioner (Gaussian Elim.)
 - Generally, $\mathbf{A}'\mathbf{x}'=\mathbf{b}'$ where $\mathbf{A}'=\mathbf{M}_L^{-1}\mathbf{A}\mathbf{M}_R^{-1}$, $\mathbf{b}'=\mathbf{M}_L^{-1}\mathbf{b}$, $\mathbf{x}'=\mathbf{M}_R\mathbf{x}$
 - $\mathbf{M}_L/\mathbf{M}_R$: Left/Right Preconditioning (左／右前处理)

Preconditioned CG Solver

```

Compute  $\mathbf{r}^{(0)} = \mathbf{b} - [\mathbf{A}]\mathbf{x}^{(0)}$ 
for  $i = 1, 2, \dots$ 
    solve  $[\mathbf{M}]\mathbf{z}^{(i-1)} = \mathbf{r}^{(i-1)}$ 
     $\rho_{i-1} = \mathbf{r}^{(i-1)} \cdot \mathbf{z}^{(i-1)}$ 
    if  $i = 1$ 
         $\mathbf{p}^{(1)} = \mathbf{z}^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $\mathbf{p}^{(i)} = \mathbf{z}^{(i-1)} + \beta_{i-1} \mathbf{p}^{(i-1)}$ 
    endif
     $\mathbf{q}^{(i)} = [\mathbf{A}]\mathbf{p}^{(i)}$ 
     $\alpha_i = \rho_{i-1} / \mathbf{p}^{(i)} \cdot \mathbf{q}^{(i)}$ 
     $\mathbf{x}^{(i)} = \mathbf{x}^{(i-1)} + \alpha_i \mathbf{p}^{(i)}$ 
     $\mathbf{r}^{(i)} = \mathbf{r}^{(i-1)} - \alpha_i \mathbf{q}^{(i)}$ 
    check convergence  $|\mathbf{r}|$ 
end

```

$$[\mathbf{M}] = [\mathbf{M}_1][\mathbf{M}_2]$$

$$[\mathbf{A}']\mathbf{x}' = \mathbf{b}'$$

$$[\mathbf{A}'] = [\mathbf{M}_1]^{-1}[\mathbf{A}][\mathbf{M}_2]^{-1}$$

$$\mathbf{x}' = [\mathbf{M}_2]\mathbf{x}, \quad \mathbf{b}' = [\mathbf{M}_1]^{-1}\mathbf{b}$$

$$\mathbf{p}' \Rightarrow [\mathbf{M}_2]\mathbf{p}, \quad \mathbf{r}' \Rightarrow [\mathbf{M}_1]^{-1}\mathbf{r}$$

$$\mathbf{p}'^{(i)} = \mathbf{r}'^{(i-1)} + \beta'_{i-1} \mathbf{p}'^{(i-1)}$$

$$[\mathbf{M}_2]\mathbf{p}^{(i)} = [\mathbf{M}_1]^{-1}\mathbf{r}^{(i-1)} + \beta'_{i-1} [\mathbf{M}_2]\mathbf{p}^{(i-1)}$$

$$\mathbf{p}^{(i)} = [\mathbf{M}_2]^{-1}[\mathbf{M}_1]^{-1}\mathbf{r}^{(i-1)} + \beta'_{i-1} \mathbf{p}^{(i-1)}$$

$$\mathbf{p}^{(i)} = [\mathbf{M}]^{-1}\mathbf{r}^{(i-1)} + \beta'_{i-1} \mathbf{p}^{(i-1)}$$

$$\beta'_{i-1} = \frac{([\mathbf{M}]^{-1}\mathbf{r}^{(i-1)}, \mathbf{r}^{(i-1)})}{([\mathbf{M}]^{-1}\mathbf{r}^{(i-2)}, \mathbf{r}^{(i-2)})}$$

$$\alpha'_{i-1} = \frac{([\mathbf{M}]^{-1}\mathbf{r}^{(i-1)}, \mathbf{r}^{(i-1)})}{(\mathbf{p}^{(i-1)}, [\mathbf{A}]\mathbf{p}^{(i-1)})}$$

Preconditioning in PCG

```

Compute  $r^{(0)} = b - [A]x^{(0)}$ 
for  $i = 1, 2, \dots$ 
    solve  $[M]z^{(i-1)} = r^{(i-1)}$ 
     $\rho_{i-1} = r^{(i-1)} \cdot z^{(i-1)}$ 
    if  $i = 1$ 
         $p^{(1)} = z^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
    endif
     $q^{(i)} = [A]p^{(i)}$ 
     $\alpha_i = \rho_{i-1} / p^{(i)} \cdot q^{(i)}$ 
     $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
     $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
    check convergence  $|r|$ 
end

```

Solving the following equation:

$$\{z\} = [M]^{-1} \{r\}$$

“Approximate Inverse Matrix”

$$[M]^{-1} \approx [A]^{-1}, \quad [M] \approx [A]$$

Ultimate Preconditioning:

Inverse Matrix

$$[M]^{-1} = [A]^{-1}, \quad [M] = [A]$$

Diagonal Scaling: Simple but weak

$$[M]^{-1} = [D]^{-1}, \quad [M] = [D]$$

ILU(0), IC(0)

- Widely used Preconditioners for Sparse Matrices
 - Incomplete LU Factorization (不完全LU分解)
 - Incomplete Cholesky Factorization (for Symmetric Matrices) (不完全コレスキー分解)
- Incomplete Direct Method
 - Even if original matrix is sparse, inverse matrix is not necessarily sparse.
 - fill-in
 - ILU(0)/IC(0) without fill-in have same non-zero pattern with the original (sparse) matrices

Diagonal Scaling, Point-Jacobi

$$[M] = \begin{bmatrix} D_1 & 0 & \dots & 0 & 0 \\ 0 & D_2 & & 0 & 0 \\ \dots & & \dots & & \dots \\ 0 & 0 & & D_{N-1} & 0 \\ 0 & 0 & \dots & 0 & D_N \end{bmatrix}$$

- **solve $[M] \mathbf{z}^{(i-1)} = \mathbf{r}^{(i-1)}$** is very easy.
- Provides fast convergence for simple problems.
- 1d.f, 1d.c

- 1D-code for Static Linear-Elastic Problems by Galerkin FEM
- Sparse Linear Solver
 - Conjugate Gradient Method
 - Preconditioning
- Storage of Sparse Matrices
- Program

Coef. Matrix derived from FEM

- Sparse Matrix
 - Many “0”s
- Storing all components (e.g. $A(i,j)$) is not efficient for sparse matrices
 - $A(i,j)$ is suitable for dense matrices

$$\begin{bmatrix}
 D & X & & & & & & & & & & & & & & & \\
 X & D & X & & & & & & & & & & & & & & \\
 & & X & D & X & & & & & & & & & & & & \\
 & & & X & D & & & & & & & & & & & & \\
 X & X & & & D & X & & & & X & X & & & & & & \\
 X & X & X & & X & D & X & & & X & X & X & & & & & \\
 & & X & X & X & & X & D & X & & & X & X & X & & & \\
 & & & X & X & & & X & D & & & X & X & & & & \\
 & & & & X & X & & & D & X & & & X & X & & & \\
 & & & & X & X & X & & X & D & X & & X & X & X & & \\
 & & & & & X & X & & & X & D & & X & X & & & \\
 & & & & & & X & X & & & D & X & & & & & \\
 & & & & & & X & X & X & & X & D & X & & & & \\
 & & & & & & & X & X & & & X & D & X & & & \\
 & & & & & & & & X & X & & & X & D & & & \\
 & & & & & & & & & X & X & & & X & D & & \\
 & & & & & & & & & & X & X & & & X & D & \\
 & & & & & & & & & & & X & X & & & X & D
 \end{bmatrix}
 \begin{Bmatrix}
 \Phi_1 \\ \Phi_2 \\ \Phi_3 \\ \Phi_4 \\ \Phi_5 \\ \Phi_6 \\ \Phi_7 \\ \Phi_8 \\ \Phi_9 \\ \Phi_{10} \\ \Phi_{11} \\ \Phi_{12} \\ \Phi_{13} \\ \Phi_{14} \\ \Phi_{15} \\ \Phi_{16}
 \end{Bmatrix}
 =
 \begin{Bmatrix}
 F_1 \\ F_2 \\ F_3 \\ F_4 \\ F_5 \\ F_6 \\ F_7 \\ F_8 \\ F_9 \\ F_{10} \\ F_{11} \\ F_{12} \\ F_{13} \\ F_{14} \\ F_{15} \\ F_{16}
 \end{Bmatrix}$$

- Number of non-zero off-diagonal components is $O(100)$ in FEM
 - If number of unknowns is 10^8 :
 - $A(i,j)$: $O(10^{16})$ words
 - Actual Non-zero Components: $O(10^{10})$ words
- Only (really) non-zero off-diag. components should be stored on memory

Variables/Arrays in 1d.f, 1d.c related to coefficient matrix

name	type	size	description
N	I	–	# Unknowns
NPLU	I	–	# Non-Zero Off-Diagonal Components
Diag(:)	R	N	Diagonal Components
U(:)	R	N	Unknown Vector
Rhs(:)	R	N	RHS Vector
Index(:)	I	0:N N+1	Off-Diagonal Components (Number of Non-Zero Off-Diagonals at Each ROW)
Item(:)	I	NPLU	Off-Diagonal Components (Corresponding Column ID)
AMat(:)	R	NPLU	Off-Diagonal Components (Value)

Only non-zero components are stored according to “Compressed Row Storage”.

Mat-Vec. Multiplication for Sparse Matrix

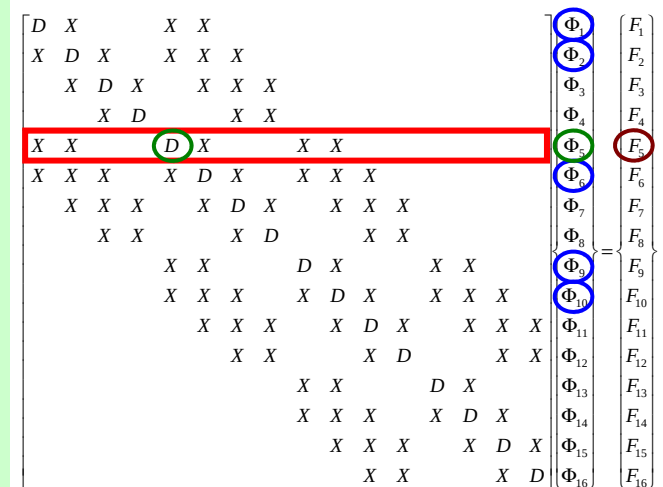
Compressed Row Storage (CRS)

Diag (i) Diagonal Components (REAL, i=1~N)
Index (i) Number of Non-Zero Off-Diagonals at Each ROW (INT, i=0~N)
Item (k) Off-Diagonal Components (Corresponding Column ID)
 (INT, k=1, index(N))
AMat (k) Off-Diagonal Components (Value)
 (REAL, k=1, index(N))

$$\{Y\} = [A] \{X\}$$

```

do i= 1, N
  Y(i)= Diag(i)*X(i)
  do k= Index(i-1)+1, Index(i)
    Y(i)= Y(i) + Amat(k)*X(Item(k))
  enddo
enddo
  
```



CRS or CSR ? for Compressed Row Storage

- In Japan and USA, “CRS” is very general for abbreviation of “Compressed Row Storage”, but they usually use “CSR” in Europe (especially in France).
- “CRS” in France
 - Compagnie Républicaine de Sécurité
 - Republic Security Company of France
- French scientists may feel uncomfortable when we use “CRS” in technical papers and/or presentations.



Mat-Vec. Multiplication for Sparse Matrix

Compressed Row Storage (CRS)

```
{Q}=[A] {P}
```

```
for (i=0; i<N; i++) {  
    W[Q][i] = Diag[i] * W[P][i];  
    for (k=Index[i]; k<Index[i+1]; k++) {  
        W[Q][i] += AMat[k]*W[P][Item[k]];  
    }  
}
```

Mat-Vec. Multiplication for Dense Matrix

Very Easy, Straightforward

$$\begin{bmatrix} a_{11} & a_{12} & \dots & a_{1,N-1} & a_{1,N} \\ a_{21} & a_{22} & & a_{2,N-1} & a_{2,N} \\ \dots & & \dots & & \dots \\ a_{N-1,1} & a_{N-1,2} & & a_{N-1,N-1} & a_{N-1,N} \\ a_{N,1} & a_{N,2} & \dots & a_{N,N-1} & a_{N,N} \end{bmatrix} \begin{Bmatrix} x_1 \\ x_2 \\ \vdots \\ x_{N-1} \\ x_N \end{Bmatrix} = \begin{Bmatrix} y_1 \\ y_2 \\ \vdots \\ y_{N-1} \\ y_N \end{Bmatrix}$$

$$\{Y\} = [A] \{X\}$$

```
do j= 1, N
  Y(j)= 0. d0
  do i= 1, N
    Y(j)= Y(j) + A(i, j)*X(i)
  enddo
enddo
```

Compressed Row Storage (CRS)

	1	2	3	4	5	6	7	8
1	1.1	2.4	0	0	3.2	0	0	0
2	4.3	3.6	0	2.5	0	3.7	0	9.1
3	0	0	5.7	0	1.5	0	3.1	0
4	0	4.1	0	9.8	2.5	2.7	0	0
5	3.1	9.5	10.4	0	11.5	0	4.3	0
6	0	0	6.5	0	0	12.4	9.5	0
7	0	6.4	2.5	0	0	1.4	23.1	13.1
8	0	9.5	1.3	9.6	0	3.1	0	51.3

Compressed Row Storage (CRS): Fortran

	①	②	③	④	⑤	⑥	⑦	⑧
①	1.1 ①	2.4 ②			3.2 ⑤			
②	4.3 ①	3.6 ②		2.5 ④		3.7 ⑥		9.1 ⑧
③			5.7 ③		1.5 ⑤		3.1 ⑦	
④		4.1 ②		9.8 ④	2.5 ⑤	2.7 ⑥		
⑤	3.1 ①	9.5 ②	10.4 ③		11.5 ⑤		4.3 ⑦	
⑥			6.5 ③			12.4 ⑥	9.5 ⑦	
⑦		6.4 ②	2.5 ③			1.4 ⑥	23.1 ⑦	13.1 ⑧
⑧		9.5 ②	1.3 ③	9.6 ④		3.1 ⑥		51.3 ⑧

N= 8

対角成分

Diag(1) = 1.1
 Diag(2) = 3.6
 Diag(3) = 5.7
 Diag(4) = 9.8
 Diag(5) = 11.5
 Diag(6) = 12.4
 Diag(7) = 23.1
 Diag(8) = 51.3

Compressed Row Storage (CRS)

	①	②	③	④	⑤	⑥	⑦	⑧
①	1.1 ①		2.4 ②			3.2 ⑤		
②	3.6 ②	4.3 ①			2.5 ④		3.7 ⑥	9.1 ⑧
③	5.7 ③					1.5 ⑤		3.1 ⑦
④	9.8 ④		4.1 ②			2.5 ⑤	2.7 ⑥	
⑤	11.5 ⑤	3.1 ①	9.5 ②	10.4 ③				4.3 ⑦
⑥	12.4 ⑥			6.5 ③				9.5 ⑦
⑦	23.1 ⑦		6.4 ②	2.5 ③			1.4 ⑥	13.1 ⑧
⑧	51.3 ⑧		9.5 ②	1.3 ③	9.6 ④		3.1 ⑥	

Compressed Row Storage (CRS)

						# Non-Zero Off-Diag.	
1	1.1 ①	2.4 ②	3.2 ⑤			2	$\text{index}(0) = 0$
2	3.6 ②	4.3 ①	2.5 ④	3.7 ⑥	9.1 ⑧	4	$\text{index}(1) = 2$
3	5.7 ③	1.5 ⑤	3.1 ⑦			2	$\text{index}(2) = 6$
4	9.8 ④	4.1 ②	2.5 ⑤	2.7 ⑥		3	$\text{index}(3) = 8$
5	11.5 ⑤	3.1 ①	9.5 ②	10.4 ③	4.3 ⑦	4	$\text{index}(4) = 11$
6	12.4 ⑥	6.5 ③	9.5 ⑦			2	$\text{index}(5) = 15$
7	23.1 ⑦	6.4 ②	2.5 ③	1.4 ⑥	13.1 ⑧	4	$\text{index}(6) = 17$
8	51.3 ⑧	9.5 ②	1.3 ③	9.6 ④	3.1 ⑥	4	$\text{index}(7) = 21$
							$\text{index}(8) = 25$

NPLU= 25
(=index(N))

$\text{index}(i-1) + 1^{\text{th}} \sim \text{index}(i)^{\text{th}}$
Non-Zero Off-Diag. Components corresponding to i -th row

Compressed Row Storage (CRS)

		# Non-Zero Off-Diag.	
1	1.1 ①	2	$\text{index}(0) = 0$
	2.4 ②,1		
	3.2 ⑤,2		
2	3.6 ②	4	$\text{index}(1) = 2$
	4.3 ①,3		
	2.5 ④,4		
	3.7 ⑥,5		
	9.1 ⑧,6		
3	5.7 ③	2	<u>$\text{index}(2) = 6$</u>
	1.5 ⑤,7		
	3.1 ⑦,8		
4	9.8 ④	3	<u>$\text{index}(3) = 8$</u>
	4.1 ②,9		
	2.5 ⑤,10		
	2.7 ⑥,11		<u>$\text{index}(4) = 11$</u>
5	11.5 ⑤	4	$\text{index}(5) = 15$
	3.1 ①,12		
	9.5 ②,13		
	10.4 ③,14		
	4.3 ⑦,15		
6	12.4 ⑥	2	$\text{index}(6) = 17$
	6.5 ③,16		
	9.5 ⑦,17		
7	23.1 ⑦	4	$\text{index}(7) = 21$
	6.4 ②,18		
	2.5 ③,19		
	1.4 ⑥,20		
	13.1 ⑧,21		
8	51.3 ⑧	4	$\text{index}(8) = 25$
	9.5 ②,22		
	1.3 ③,23		
	9.6 ④,24		
	3.1 ⑥,25		

$\text{index}(i-1) + 1^{\text{th}} \sim \text{index}(i)^{\text{th}}$
Non-Zero Off-Diag. Components corresponding to i -th row

NPLU = 25
(=index(N))

Compressed Row Storage (CRS)

1	1.1 ①	2.4 ②,1	3.2 ⑤,2		
2	3.6 ②	4.3 ①,3	2.5 ④,4	3.7 ⑥,5	9.1 ⑧,6
3	5.7 ③	1.5 ⑤,7	3.1 ⑦,8		
4	9.8 ④	4.1 ②,9	2.5 ⑤,10	2.7 ⑥,11	
5	11.5 ⑤	3.1 ①,12	9.5 ②,13	10.4 ③,14	4.3 ⑦,15
6	12.4 ⑥	6.5 ③,16	9.5 ⑦,17		
7	23.1 ⑦	6.4 ②,18	2.5 ③,19	1.4 ⑥,20	13.1 ⑧,21
8	51.3 ⑧	9.5 ②,22	1.3 ③,23	9.6 ④,24	3.1 ⑥,25

Example:

`item( 7) = 5, AMAT( 7) = 1.5`

`item(19) = 3, AMAT(19) = 2.5`

Compressed Row Storage (CRS)

1	1.1 ①	2.4 ②,1	3.2 ⑤,2		
2	3.6 ②	4.3 ①,3	2.5 ④,4	3.7 ⑥,5	9.1 ⑧,6
3	5.7 ③	1.5 ⑤,7	3.1 ⑦,8		
4	9.8 ④	4.1 ②,9	2.5 ⑤,10	2.7 ⑥,11	
5	11.5 ⑤	3.1 ①,12	9.5 ②,13	10.4 ③,14	4.3 ⑦,15
6	12.4 ⑥	6.5 ③,16	9.5 ⑦,17		
7	23.1 ⑦	6.4 ②,18	2.5 ③,19	1.4 ⑥,20	13.1 ⑧,21
8	51.3 ⑧	9.5 ②,22	1.3 ③,23	9.6 ④,24	3.1 ⑥,25

Diag (i) Diagonal Components (REAL, i=1~N)
Index (i) Number of Non-Zero Off-Diagonals at Each ROW (INT, i=0~N)
Item (k) Off-Diagonal Components (Corresponding Column ID) (INT, k=1, index(N))
AMat (k) Off-Diagonal Components (Value) (REAL, k=1, index(N))

$$\{Y\} = [A] \{X\}$$

```

do i= 1, N
  Y(i)= D(i)*X(i)
  do k= index(i-1)+1, index(i)
    Y(i)= Y(i) + AMAT(k)*X(item(k))
  enddo
enddo

```

- 1D-code for Static Linear-Elastic Problems by Galerkin FEM
- Sparse Linear Solver
 - Conjugate Gradient Method
 - Preconditioning
- Storage of Sparse Matrices
- Program

Finite Element Procedures

- Initialization
 - Control Data
 - Node, Connectivity of Elements (N: Node#, NE: Elem#)
 - Initialization of Arrays (Global/Element Matrices)
 - Element-Global Matrix Mapping (Index, Item)
- Generation of Matrix
 - Element-by-Element Operations (do icel= 1, NE)
 - Element matrices
 - Accumulation to global matrix
 - Boundary Conditions
- Linear Solver
 - Conjugate Gradient Method

Program: 1d.f (1/6)

variables and arrays

```
!C
!C 1D Steady-State Heat Transfer
!C FEM with Piece-wise Linear Elements
!C CG (Conjugate Gradient) Method
!C
!C  $d/dx(CdT/dx) + Q = 0$ 
!C  $T=0@x=0$ 
!C
  program heat1D
    implicit REAL*8 (A-H, O-Z)

    integer :: N, NPLU, ITERmax
    integer :: R, Z, P, Q, DD

    real(kind=8) :: dX, RESID, EPS
    real(kind=8) :: AREA, QV, COND
    real(kind=8), dimension(:), allocatable :: PHI, RHS, X
    real(kind=8), dimension(:), allocatable :: DIAG, AMAT
    real(kind=8), dimension(:, :), allocatable :: W

    real(kind=8), dimension(2, 2) :: KMAT, EMAT

    integer, dimension(:), allocatable :: ICELNOD
    integer, dimension(:), allocatable :: INDEX, ITEM
```

Variable/Arrays (1/2)

Name	Type	Size	I/O	Definition
NE	I		I	# Element
N	I		O	# Node
NPLU	I		O	# Non-Zero Off-Diag. Components
IterMax	I		I	MAX Iteration Number for CG
errno	I		O	ERROR flag
R, Z, Q, P, DD	I		O	Name of Vectors in CG
dX	R		I	Length of Each Element
Resid	R		O	Residual for CG
Eps	R		I	Convergence Criteria for CG
Area	R		I	Sectional Area of Element
QV	R		I	Heat Generation Rate/Volume/Time $\dot{Q}$
COND	R		I	Thermal Conductivity

Variable/Arrays (2/2)

Name	Type	Size	I/O	Definition
X	R	N	○	Location of Each Node
PHI	R	N	○	Temperature of Each Node
Rhs	R	N	○	RHS Vector
Diag	R	N	○	Diagonal Components
W	R	(N, 4)	○	Work Array for CG
Amat	R	NPLU	○	Off-Diagonal Components (Value)
Index	I	0:N	○	Number of Non-Zero Off-Diagonals at Each ROW
Item	I	NPLU	○	Off-Diagonal Components (Corresponding Column ID)
Icelnod	I	2*NE	○	Node ID for Each Element
Kmat	R	(2, 2)	○	Element Matrix [k]
Emat	R	(2, 2)	○	Element Matrix

Program: 1d.f (2/6)

Initialization, Allocation of Arrays

```
!C
!C +-----+
!C |  INIT.  |
!C +-----+
!C==
```

```
open (11, file='input.dat', status='unknown')
read (11,*) NE
read (11,*) dX, QV, AREA, COND
read (11,*) ITERmax
read (11,*) EPS
close (11)
```

Control Data input.dat

4	NE (Number of Elements)
1.0 1.0 1.0 1.0	Δx (Length of Each Elem.: L), Q , A , λ
100	Number of MAX. Iterations for CG Solver
1.e-8	Convergence Criteria for CG Solver

```
N= NE + 1
allocate (PHI(N), DIAG(N), AMAT(2*N-2), RHS(N))
allocate (ICELNOD(2*NE), X(N))
allocate (INDEX(0:N), ITEM(2*N-2), W(N,4))
```

```
PHI = 0. d0
AMAT= 0. d0
DIAG= 0. d0
RHS= 0. d0
X= 0. d0
```



NE: # Element
N : # Node (NE+1)

Program: 1d.f (2/6)

Initialization, Allocation of Arrays

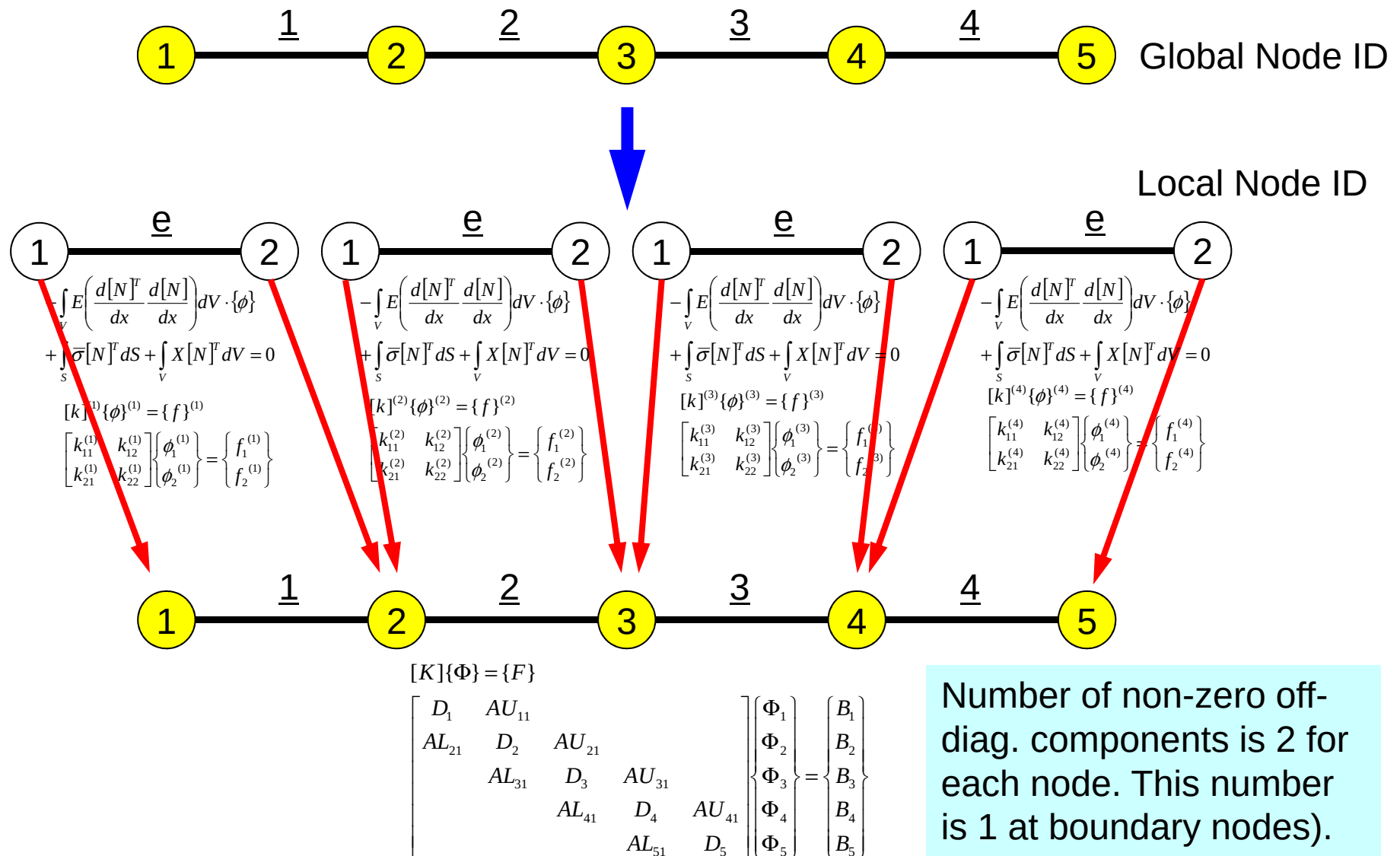
```
!C
!C +-----+
!C |  INIT.  |
!C +-----+
!C===
      open (11, file='input.dat', status='unknown')
      read (11,*) NE
      read (11,*) dX, QV, AREA, COND
      read (11,*) ITERmax
      read (11,*) EPS
      close (11)
```

```
      N= NE + 1
      allocate (PHI(N), DIAG(N), AMAT(2*N-2), RHS(N))
      allocate (ICELNOD(2*NE), X(N))
      allocate (INDEX(0:N), ITEM(2*N-2), W(N,4))
```

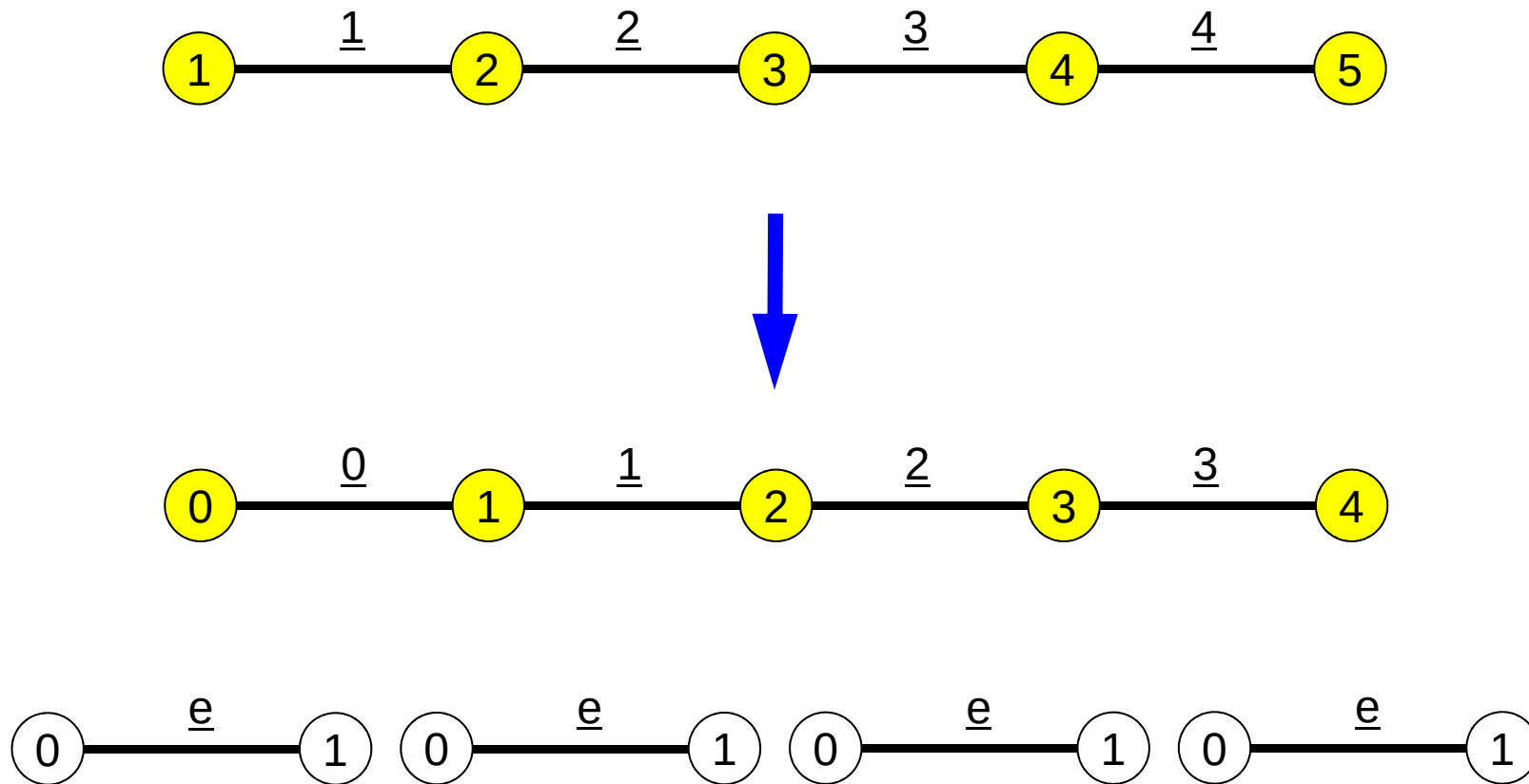
```
      PHI = 0. d0
      AMAT= 0. d0
      DIAG= 0. d0
      RHS= 0. d0
      X= 0. d0
```

Amat: Non-Zero Off-Diag. Comp.
Item: Corresponding Column ID

Element/Global Operations



Attention: In C program, node and element ID's start from 0.



Program: 1d.f (2/6)

Initialization, Allocation of Array

```

!C
!C +-----+
!C |  INIT.  |
!C +-----+
!C===
      open  (11, file='input.dat', status='unknown')
      read  (11,*) NE
      read  (11,*) dX, QV, AREA, COND
      read  (11,*) ITERmax
      read  (11,*) EPS
      close (11)

```

```

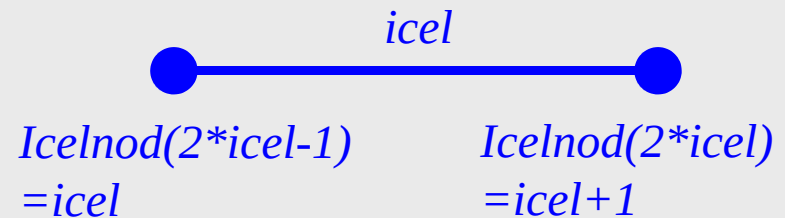
N= NE + 1
allocate (PHI(N), DIAG(N), AMAT(2*N-2), RHS(N))
allocate (ICELNOD(2*NE), X(N))
allocate (INDEX(0:N), ITEM(2*N-2), W(N,4))

```

```

PHI = 0. d0
AMAT= 0. d0
DIAG= 0. d0
RHS= 0. d0
X= 0. d0

```



Amat: Non-Zero Off-Diag. Comp.

Item: Corresponding Column ID

Number of non-zero off-diag. components is 2 for each node. This number is 1 at boundary nodes).

Total Number of Non-Zero Off-Diag. Components:

$$2 * (N-2) + 1 + 1 = 2 * N - 2$$

Program: 1d.f (3/6)

Initialization, Allocation of Arrays (cont.)

```
do i= 1, N
  X(i)= dfloat(i-1)*dX
enddo

do icel= 1, NE
  ICELNOD(2*icel-1)= icel
  ICELNOD(2*icel )= icel + 1
enddo

KMAT (1, 1)= +1. d0
KMAT (1, 2)= -1. d0
KMAT (2, 1)= -1. d0
KMAT (2, 2)= +1. d0
```

x: X-coordinate
component of each node

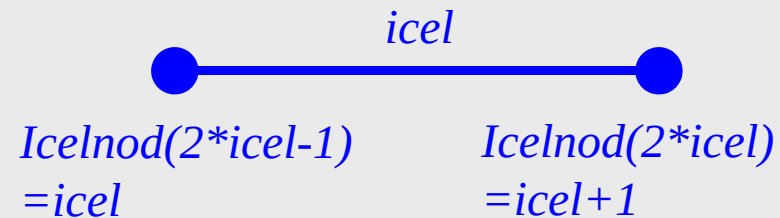
Program: 1d.f (3/6)

Initialization, Allocation of Arrays (cont.)

```
do i= 1, N  
  X(i)= dfloat(i-1)*dX  
enddo
```

```
do icel= 1, NE  
  ICELNOD(2*icel-1)= icel  
  ICELNOD(2*icel )= icel + 1  
enddo
```

```
KMAT (1, 1)= +1. d0  
KMAT (1, 2)= -1. d0  
KMAT (2, 1)= -1. d0  
KMAT (2, 2)= +1. d0
```



Program: 1d.f (3/6)

Initialization, Allocation of Arrays (cont.)

```
do i= 1, N
  X(i)= dfloat(i-1)*dX
enddo

do icel= 1, NE
  ICELNOD(2*icel-1)= icel
  ICELNOD(2*icel )= icel + 1
enddo
```

```
KMAT (1, 1)= +1. d0
KMAT (1, 2)= -1. d0
KMAT (2, 1)= -1. d0
KMAT (2, 2)= +1. d0
```

$$[k]^{(e)} = \int_V \lambda \left(\frac{d[N]^T}{dx} \frac{d[N]}{dx} \right) dV = \frac{\lambda A}{L} \begin{bmatrix} +1 & -1 \\ -1 & +1 \end{bmatrix}$$

[Kmat]

Program: 1d.f (4/6)

Global Matrix: Column ID for Non-Zero Off-Diag's

```
!C
!C +-----+
!C | CONNECTIVITY |
!C +-----+
!C==
```

INDEX = 2

INDEX(0) = 0

INDEX(1) = 1

INDEX(N) = 1

```
do i= 1, N
  INDEX(i)= INDEX(i) + INDEX(i-1)
enddo
```

NPLU= INDEX(N)

```
do i= 1, N
  js= INDEX(i-1)
  if (i.eq.1) then
    ITEM(js+1)= i+1
  else if
&    (i.eq.N) then
    ITEM(js+1)= i-1
  else
    ITEM(js+1)= i-1
    ITEM(js+2)= i+1
  endif
enddo
```

```
!C==
```

Number of non-zero off-diag. components is 2 for each node. This number is 1 at boundary nodes).

Total Number of Non-Zero Off-Diag. Components:
 $2*(N-2)+1+1= 2*N-2= NPLU= \text{Index}[N]$

						# Non-Zero Off-Diag.	index(0) = 0
1	1.1 ①	2.4 ②	3.2 ⑤			2	index(1) = 2
2	3.6 ②	4.3 ①	2.5 ④	3.7 ⑥	9.1 ⑧	4	index(2) = 6
3	5.7 ③	1.5 ⑤	3.1 ⑦			2	index(3) = 8
4	9.8 ④	4.1 ②	2.5 ⑤	2.7 ⑥		3	index(4) = 11
5	11.5 ⑤	3.1 ①	9.5 ②	10.4 ③	4.3 ⑦	4	index(5) = 15
6	12.4 ⑥	6.5 ③	9.5 ⑦			2	index(6) = 17
7	23.1 ⑦	6.4 ②	2.5 ③	1.4 ⑥	13.1 ⑧	4	index(7) = 21
8	51.3 ⑧	9.5 ②	1.3 ③	9.6 ④	3.1 ⑥	4	index(8) = 25

$\text{index}(i-1)+1^{\text{th}} \sim \text{index}(i)^{\text{th}}$
 Non-Zero Off-Diag. Components corresponding to i -th row

Program: 1d.f (4/6)

Global Matrix: Column ID for Non-Zero Off-Diag's

```
!C
!C +-----+
!C | CONNECTIVITY |
!C +-----+
!C===
```

```
INDEX = 2
```

```
INDEX(0) = 0
```

```
INDEX(1) = 1
```

```
INDEX(N) = 1
```

```
do i= 1, N
  INDEX(i) = INDEX(i) + INDEX(i-1)
enddo
```

```
NPLU= INDEX(N)
```

```
do i= 1, N
  jS= INDEX(i-1)
  if (i.eq.1) then
    ITEM(jS+1)= i+1
  else if
&    (i.eq.N) then
    ITEM(jS+1)= i-1
  else
    ITEM(jS+1)= i-1
    ITEM(jS+2)= i+1
  endif
enddo
```

```
!C===
```



						# Non-Zero Off-Diag.	index(0) = 0
1	1.1 ①	2.4 ②	3.2 ⑤			2	index(1) = 2
2	3.6 ②	4.3 ①	2.5 ④	3.7 ⑥	9.1 ⑧	4	index(2) = 6
3	5.7 ③	1.5 ⑤	3.1 ⑦			2	index(3) = 8
4	9.8 ④	4.1 ②	2.5 ⑤	2.7 ⑥		3	index(4) = 11
5	11.5 ⑤	3.1 ①	9.5 ②	10.4 ③	4.3 ⑦	4	index(5) = 15
6	12.4 ⑥	6.5 ③	9.5 ⑦			2	index(6) = 17
7	23.1 ⑦	6.4 ②	2.5 ③	1.4 ⑥	13.1 ⑧	4	index(7) = 21
8	51.3 ⑧	9.5 ②	1.3 ③	9.6 ④	3.1 ⑥	4	index(8) = 25

$\text{index}(i-1) + 1^{\text{th}} \sim \text{index}(i)^{\text{th}}$
Non-Zero Off-Diag. Components corresponding to i -th row

Program: 1d.f (5/6)

Element Matrix ~ Global Matrix

```
!C +-----+
!C | MATRIX ASSEMBLE |
!C +-----+
!C==
```

```
do icel= 1, NE
  in1= ICELNOD(2*icel-1)
  in2= ICELNOD(2*icel )
  X1 = X(in1)
  X2 = X(in2)
  DL = dabs(X2-X1)
```

```
  cK= AREA*COND/DL
  EMAT(1,1)= Ck*KMAT(1,1)
  EMAT(1,2)= Ck*KMAT(1,2)
  EMAT(2,1)= Ck*KMAT(2,1)
  EMAT(2,2)= Ck*KMAT(2,2)
```

```
  DIAG(in1)= DIAG(in1) + EMAT(1,1)
  DIAG(in2)= DIAG(in2) + EMAT(2,2)
```

```
  if (icel.eq.1) then
    k1= INDEX(in1-1) + 1
  else
    k1= INDEX(in1-1) + 2
  endif
  k2= INDEX(in2-1) + 1
```

```
  AMAT(k1)= AMAT(k1) + EMAT(1,2)
  AMAT(k2)= AMAT(k2) + EMAT(2,1)
```

```
  QN= 0.50d0*QV*AREA*DL
  RHS(in1)= RHS(in1) + QN
  RHS(in2)= RHS(in2) + QN
```

```
enddo
!C==
```



Program: 1d.f (5/6)

Element Matrix ~ Global Matrix

```
!C +-----+
!C | MATRIX ASSEMBLE |
!C +-----+
!C==
```

```
do icel= 1, NE
  in1= ICENOD (2*icel-1)
  in2= ICENOD (2*icel )
  X1 = X(in1)
  X2 = X(in2)
  DL = dabs(X2-X1)
```

```
  cK= AREA*COND/DL
  EMAT (1, 1)= Ck*KMAT (1, 1)
  EMAT (1, 2)= Ck*KMAT (1, 2)
  EMAT (2, 1)= Ck*KMAT (2, 1)
  EMAT (2, 2)= Ck*KMAT (2, 2)
```

```
  DIAG(in1)= DIAG(in1) + EMAT (1, 1)
  DIAG(in2)= DIAG(in2) + EMAT (2, 2)
```

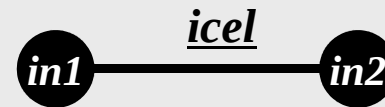
```
  if (icel.eq.1) then
    k1= INDEX(in1-1) + 1
  else
    k1= INDEX(in1-1) + 2
  endif
  k2= INDEX(in2-1) + 1
```

```
  AMAT(k1)= AMAT(k1) + EMAT (1, 2)
  AMAT(k2)= AMAT(k2) + EMAT (2, 1)
```

```
  QN= 0.50d0*QV*AREA*DL
  RHS(in1)= RHS(in1) + QN
  RHS(in2)= RHS(in2) + QN
```

```
enddo
```

```
!C==
```



$$[Emat] = [k]^{(e)} = \frac{\lambda A}{L} \begin{bmatrix} +1 & -1 \\ -1 & +1 \end{bmatrix} = \frac{\lambda A}{L} [Kmat]$$

Program: 1d.f (5/6)

Element Matrix ~ Global Matrix

```
!C +-----+
!C | MATRIX ASSEMBLE |
!C +-----+
!C==
```

```
do icel= 1, NE
  in1= ICELNOD(2*icel-1)
  in2= ICELNOD(2*icel)
  X1 = X(in1)
  X2 = X(in2)
  DL = dabs(X2-X1)
```

```
  cK= AREA*COND/DL
  EMAT(1,1)= Ck*KMAT(1,1)
  EMAT(1,2)= Ck*KMAT(1,2)
  EMAT(2,1)= Ck*KMAT(2,1)
  EMAT(2,2)= Ck*KMAT(2,2)
```

```
  DIAG(in1)= DIAG(in1) + EMAT(1,1)
  DIAG(in2)= DIAG(in2) + EMAT(2,2)
```

```
  if (icel.eq.1) then
    k1= INDEX(in1-1) + 1
  else
    k1= INDEX(in1-1) + 2
  endif
  k2= INDEX(in2-1) + 1
```

```
  AMAT(k1)= AMAT(k1) + EMAT(1,2)
  AMAT(k2)= AMAT(k2) + EMAT(2,1)
```

```
  QN= 0.50d0*QV*AREA*DL
  RHS(in1)= RHS(in1) + QN
  RHS(in2)= RHS(in2) + QN
```

```
enddo
!C==
```



$$[Emat] = [k]^{(e)} = \frac{EA}{L} \begin{bmatrix} +1 & -1 \\ -1 & +1 \end{bmatrix}$$

Program: 1d.f (5/6)

Element Matrix ~ Global Matrix

```
!C +-----+
!C | MATRIX ASSEMBLE |
!C +-----+
!C===
```

```
do icel= 1, NE
  in1= ICELNOD(2*icel-1)
  in2= ICELNOD(2*icel )
  X1 = X(in1)
  X2 = X(in2)
  DL = dabs(X2-X1)
```

```
  cK= AREA*COND/DL
  EMAT(1,1)= Ck*KMAT(1,1)
  EMAT(1,2)= Ck*KMAT(1,2)
  EMAT(2,1)= Ck*KMAT(2,1)
  EMAT(2,2)= Ck*KMAT(2,2)
```

```
  DIAG(in1)= DIAG(in1) + EMAT(1,1)
  DIAG(in2)= DIAG(in2) + EMAT(2,2)
```

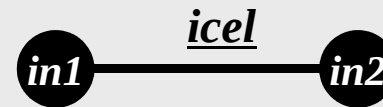
```
  if (icel.eq.1) then
    k1= INDEX(in1-1) + 1
  else
    k1= INDEX(in1-1) + 2
  endif
  k2= INDEX(in2-1) + 1
```

```
  AMAT(k1)= AMAT(k1) + EMAT(1,2)
  AMAT(k2)= AMAT(k2) + EMAT(2,1)
```

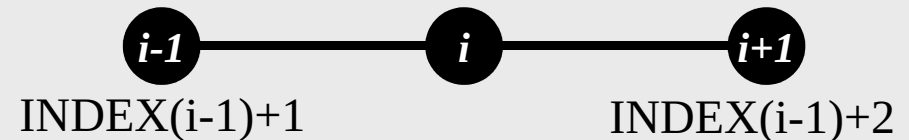
```
  QN= 0.5d0*QV*AREA*DL
  RHS(in1)= RHS(in1) + QN
  RHS(in2)= RHS(in2) + QN
```

```
enddo
```

```
!C===
```



Non-zero Off-Diag. at *i*-th row:
Index(*i*-1)+1, Index(*i*-1)+2



$$[Emat] = [k]^{(e)} = \frac{\lambda A}{L} \begin{bmatrix} +1 & \textcircled{-1} \\ \textcircled{-1} & +1 \end{bmatrix} \begin{matrix} k1 \\ k2 \end{matrix}$$

General Elements: k1

“in2” as a off-diag. component of “in1”

```
!C +-----+
!C | MATRIX ASSEMBLE |
!C +-----+
!C===
```

```
do icel= 1, NE
```

```
  in1= ICELNOD (2*icel-1)
  in2= ICELNOD (2*icel )
```

```
  X1 = X(in1)
```

```
  X2 = X(in2)
```

```
  DL = dabs(X2-X1)
```

```
  cK= AREA*COND/DL
```

```
  EMAT (1, 1)= Ck*KMAT (1, 1)
```

```
  EMAT (1, 2)= Ck*KMAT (1, 2)
```

```
  EMAT (2, 1)= Ck*KMAT (2, 1)
```

```
  EMAT (2, 2)= Ck*KMAT (2, 2)
```

```
  DIAG(in1)= DIAG(in1) + EMAT (1, 1)
```

```
  DIAG(in2)= DIAG(in2) + EMAT (2, 2)
```

```
  if (icel.eq.1) then
```

```
    k1= INDEX(in1-1) + 1
```

```
  else
```

```
    k1= INDEX(in1-1) + 2
```

```
  endif
```

```
  k2= INDEX(in2-1) + 1
```

```
  AMAT (k1)= AMAT (k1) + EMAT (1, 2)
```

```
  AMAT (k2)= AMAT (k2) + EMAT (2, 1)
```

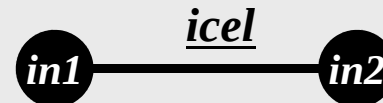
```
  QN= 0.5d0*QV*AREA*DL
```

```
  RHS(in1)= RHS(in1) + QN
```

```
  RHS(in2)= RHS(in2) + QN
```

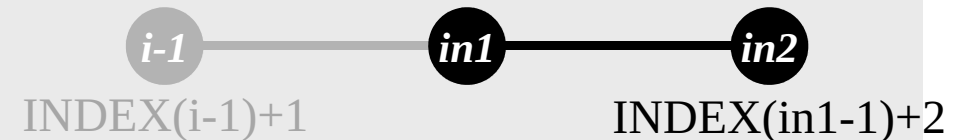
```
enddo
```

```
!C===
```



Non-zero Off-Diag. at *i*-th row:

Index(*i*-1)+1, Index(*i*-1)+2



$$[Emat] = [k]^{(e)} = \frac{\lambda A}{L} \begin{bmatrix} +1 & -1 \\ -1 & +1 \end{bmatrix} \quad k1$$

General Elements: k2

“in1” as a off-diag. component of “in2”

```

!C +-----+
!C | MATRIX ASSEMBLE |
!C +-----+
!C===
do icel= 1, NE
  in1= ICELNOD (2*icel-1)
  in2= ICELNOD (2*icel )
  X1 = X(in1)
  X2 = X(in2)
  DL = dabs(X2-X1)

  cK= AREA*COND/DL
  EMAT (1, 1)= Ck*KMAT (1, 1)
  EMAT (1, 2)= Ck*KMAT (1, 2)
  EMAT (2, 1)= Ck*KMAT (2, 1)
  EMAT (2, 2)= Ck*KMAT (2, 2)

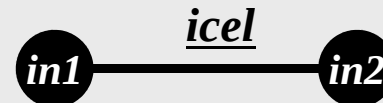
  DIAG(in1)= DIAG(in1) + EMAT (1, 1)
  DIAG(in2)= DIAG(in2) + EMAT (2, 2)

  if (icel.eq.1) then
    k1= INDEX(in1-1) + 1
  else
    k1= INDEX(in1-1) + 2
  endif
  k2= INDEX(in2-1) + 1

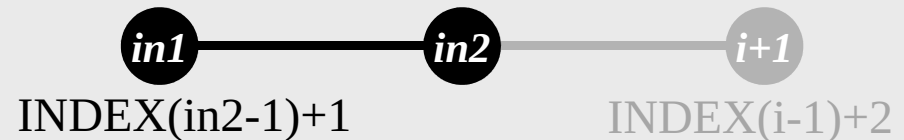
  AMAT (k1)= AMAT (k1) + EMAT (1, 2)
  AMAT (k2)= AMAT (k2) + EMAT (2, 1)

  QN= 0.50d0*QV*AREA*DL
  RHS(in1)= RHS(in1) + QN
  RHS(in2)= RHS(in2) + QN
enddo
!C===

```



Non-zero Off-Diag. at *i*-th row:
Index(*i*-1)+1, Index(*i*-1)+2



$$[Emat] = [k]^{(e)} = \frac{\lambda A}{L} \begin{bmatrix} +1 & -1 \\ \textcircled{-1} & +1 \end{bmatrix}$$

k2

0-th Element: k1

“in2” as a off-diag. component of “in1”

```
!C +-----+
!C | MATRIX ASSEMBLE |
!C +-----+
!C==
```

```
do icel= 1, NE
  in1= ICELNOD(2*icel-1)
  in2= ICELNOD(2*icel )
  X1 = X(in1)
  X2 = X(in2)
  DL = dabs(X2-X1)
```

```
  cK= AREA*COND/DL
  EMAT(1,1)= Ck*KMAT(1,1)
  EMAT(1,2)= Ck*KMAT(1,2)
  EMAT(2,1)= Ck*KMAT(2,1)
  EMAT(2,2)= Ck*KMAT(2,2)
```

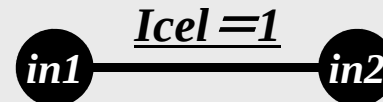
```
  DIAG(in1)= DIAG(in1) + EMAT(1,1)
  DIAG(in2)= DIAG(in2) + EMAT(2,2)
```

```
  if (icel.eq.1) then
    k1= INDEX(in1-1) + 1
  else
    k1= INDEX(in1-1) + 2
  endif
  k2= INDEX(in2-1) + 1
```

```
  AMAT(k1)= AMAT(k1) + EMAT(1,2)
  AMAT(k2)= AMAT(k2) + EMAT(2,1)
```

```
  QN= 0.50d0*QV*AREA*DL
  RHS(in1)= RHS(in1) + QN
  RHS(in2)= RHS(in2) + QN
enddo
```

```
!C==
```



Non-zero Off-Diag. at i -th row:
Index(i-1)+1 only



$$[Emat] = [k]^{(e)} = \frac{\lambda A}{L} \begin{bmatrix} +1 & \textcircled{-1} \\ -1 & +1 \end{bmatrix} \quad k1$$

Program: 1d.f (5/6)

RHS: Heat Generation Term

```
!C +-----+
!C | MATRIX ASSEMBLE |
!C +-----+
!C==
```

```
do icel= 1, NE
  in1= ICELNOD(2*icel-1)
  in2= ICELNOD(2*icel )
  X1 = X(in1)
  X2 = X(in2)
  DL = dabs(X2-X1)
```

```
  cK= AREA*COND/DL
  EMAT(1,1)= Ck*KMAT(1,1)
  EMAT(1,2)= Ck*KMAT(1,2)
  EMAT(2,1)= Ck*KMAT(2,1)
  EMAT(2,2)= Ck*KMAT(2,2)
```

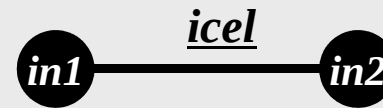
```
  DIAG(in1)= DIAG(in1) + EMAT(1,1)
  DIAG(in2)= DIAG(in2) + EMAT(2,2)
```

```
  if (icel.eq.1) then
    k1= INDEX(in1-1) + 1
  else
    k1= INDEX(in1-1) + 2
  endif
  k2= INDEX(in2-1) + 1
```

```
  AMAT(k1)= AMAT(k1) + EMAT(1,2)
  AMAT(k2)= AMAT(k2) + EMAT(2,1)
```

```
  QN= 0.5d0*QV*AREA*DL
  RHS(in1)= RHS(in1) + QN
  RHS(in2)= RHS(in2) + QN
```

```
enddo
!C==
```



$$\int_V \dot{Q} [N]^T dV = \dot{Q} A \int_0^L \begin{bmatrix} 1-x/L \\ x/L \end{bmatrix} dx = \frac{\dot{Q} A L}{2} \begin{Bmatrix} 1 \\ 1 \end{Bmatrix}$$

Program: 1d.f (6/6)

Dirichlet B.C. @ $X=0$

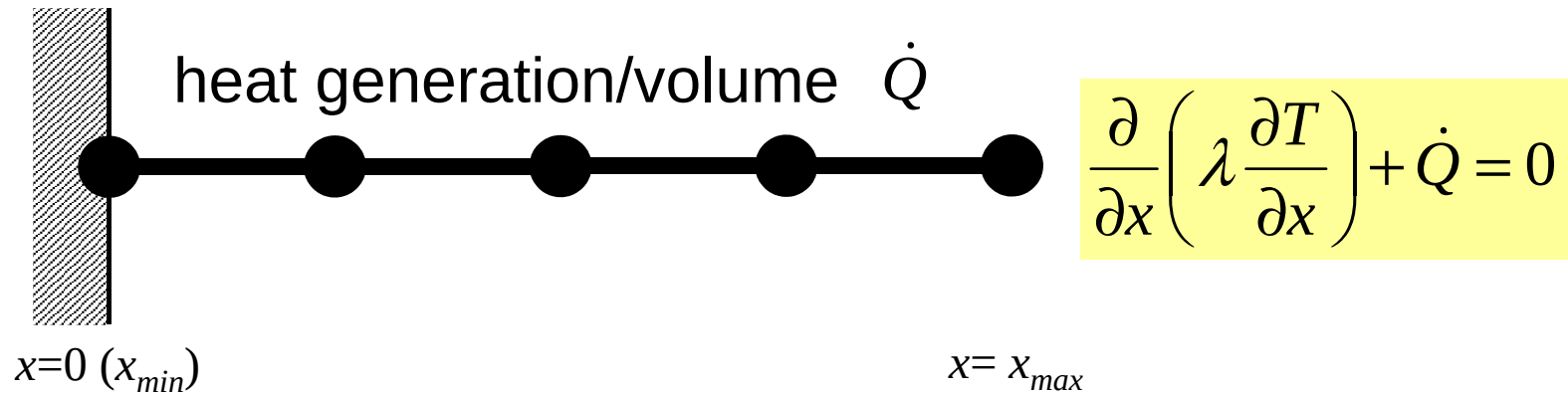
```
!C
!C +-----+
!C | BOUNDARY CONDITIONS |
!C +-----+
!C===

!C
!C-- X=Xmin
      i= 1
      jS= INDEX(i-1)

      AMAT(jS+1)= 0. d0
      DIAG(i)= 1. d0
      RHS (i)= 0. d0

      do k= 1, NPLU
        if (ITEM(k).eq. 1) AMAT(k)= 0. d0
      enddo
!C===
```

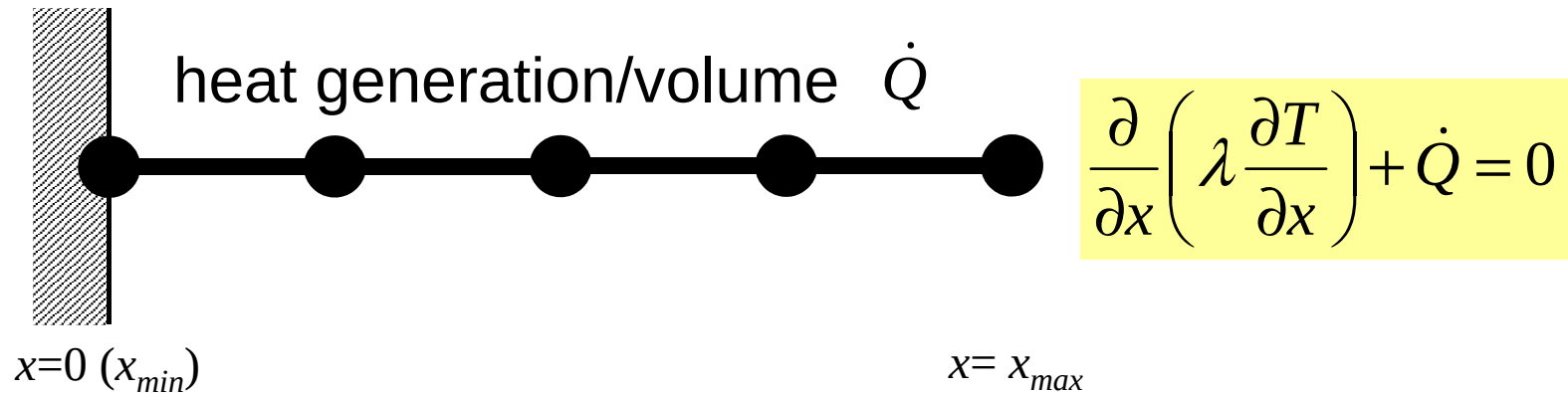
1D Steady State Heat Conduction



- **Uniform: Sectional Area: A , Thermal Conductivity: λ**
- Heat Generation Rate/Volume/Time [$QL^{-3}T^{-1}$] $\dot{Q}$
- Boundary Conditions
 - $x=0$: $T=0$ (Fixed Temperature)
 - $x=x_{max}$: $\frac{\partial T}{\partial x} = 0$ (Insulated)

(Linear) Equation at $x=0$

$$T_1 = 0 \text{ (or } T_0 = 0)$$



- Uniform: Sectional Area: A , Thermal Conductivity: λ
- Heat Generation Rate/Volume/Time [$QL^{-3}T^{-1}$] $\dot{Q}$
- Boundary Conditions
 - $x=0$: $T=0$ (Fixed Temperature)
 - $x=x_{max}$: $\frac{\partial T}{\partial x} = 0$ (Insulated)

Program: 1d.f (6/6)

Dirichlet B.C. @ X=0

```

!C
!C +-----+
!C | BOUNDARY CONDITIONS |
!C +-----+
!C===

!C
!C-- X=Xmin
      i= 1
      js= INDEX(i-1)

      AMAT (js+1)= 0. d0
      DIAG (i)= 1. d0
      RHS (i)= 0. d0

      do k= 1, NPLU
        if (ITEM(k).eq.1) AMAT(k)= 0. d0
      enddo
!C===

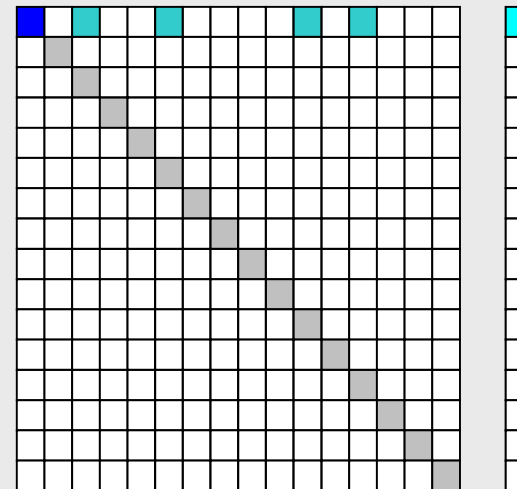
```

$$T_1=0$$

Diagonal Component=1

RHS=0

Off-Diagonal Components= 0.



Program: 1d.f (6/6)

Dirichlet B.C. @ X=0

```

!C
!C +-----+
!C | BOUNDARY CONDITIONS |
!C +-----+
!C===

!C
!C-- X=Xmin
      i= 1
      js= INDEX(i-1)

      AMAT (js+1)= 0. d0
      DIAG (i)= 1. d0
      RHS (i)= 0. d0

      do k= 1, NPLU
        if (ITEM(k).eq. 1) AMAT (k)= 0. d0
      enddo
!C===

```

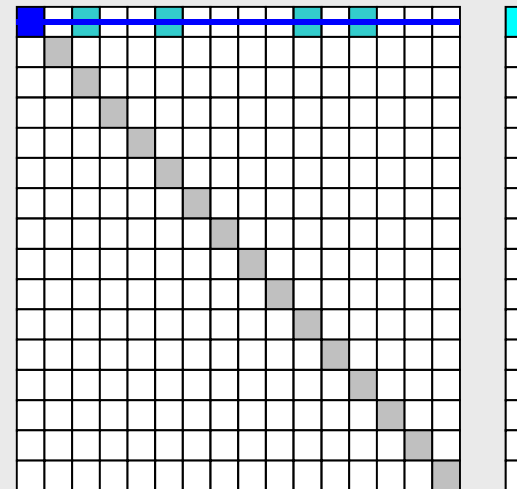
$$T_1=0$$

Diagonal Component=1

RHS=0

Off-Diagonal Components= 0.

Erase !



Program: 1d.f (6/6)

Dirichlet B.C. @ X=0

```
!C
!C +-----+
!C | BOUNDARY CONDITIONS |
!C +-----+
!C===

!C
!C-- X=Xmin
      i= 1
      js= INDEX(i-1)

      AMAT (js+1)= 0. d0
      DIAG (i)= 1. d0
      RHS (i)= 0. d0

      do k= 1, NPLU
        if (ITEM(k).eq. 1) AMAT (k)= 0. d0
      enddo
!C===
```

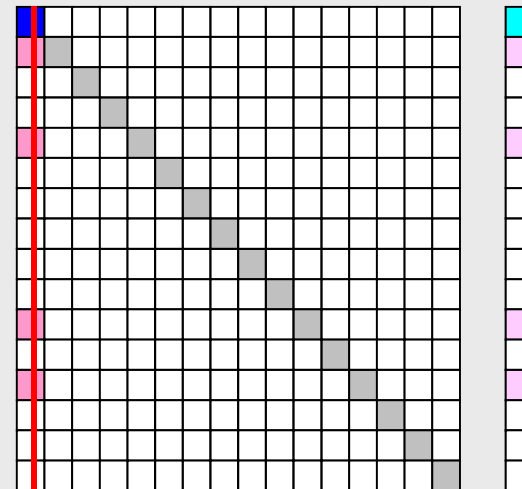
$$T_1=0$$

Diagonal Component=1

RHS=0

Off-Diagonal Components= 0.

Elimination and Erase



Column components of boundary nodes (Dirichlet B.C.) are moved to RHS and eliminated for keeping symmetrical feature of the matrix (in this case just erase off-diagonal components)

if $T_1 \neq 0$

```
!C
!C +-----+
!C | BOUNDARY CONDITIONS |
!C +-----+
!C===
```

Column components of boundary nodes (Dirichlet B.C.) are moved to RHS and eliminated for keeping symmetrical feature of the matrix.

```
!C
!C-- X=Xmin
      i= 1
      jS= INDEX(i-1)

      AMAT(jS+1)= 0. d0
      DIAG(i)= 1. d0
      RHS (i)= PHImin
```

$$Diag_j \phi_j + \sum_{k=Index[j]}^{Index[j+1]-1} Amat_k \phi_{Item[k]} = Rhs_j$$

```
do i= 1, N
  do k= INDEX(i-1)+1, INDEX(i)
    if (ITEM(k).eq.1) then
      RHS (i)= RHS(i) - AMAT(k)*PHImin
      AMAT(k)= 0. d0
    endif
  enddo
enddo
!C===
```

if $T_1 \neq 0$

```
!C
!C +-----+
!C | BOUNDARY CONDITIONS |
!C +-----+
!C===
```

```
!C
!C-- X=Xmin
      i= 1
      jS= INDEX(i-1)
```

```
      AMAT(jS+1)= 0. d0
      DIAG(i)= 1. d0
      RHS (i)= PHImin
```

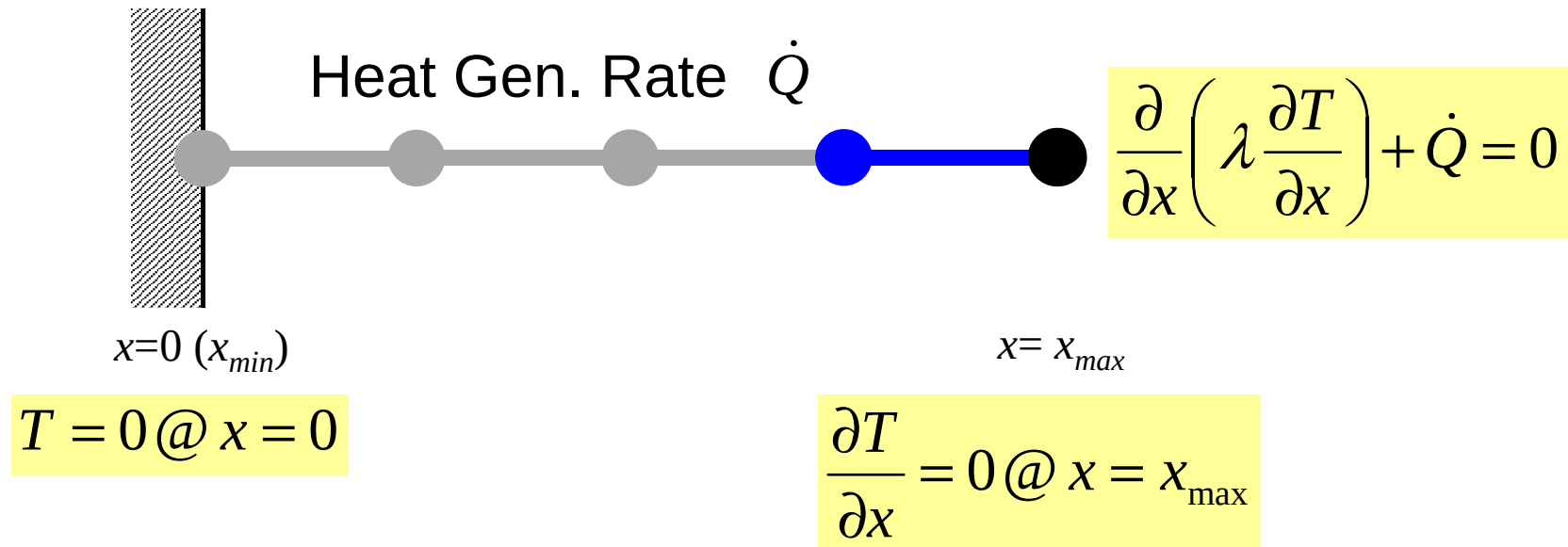
```
      do i= 1, N
        do k= INDEX(i-1)+1, INDEX(i)
          if (ITEM(k).eq.1) then
            RHS (i)= RHS(i) - AMAT(k)*PHImin
            AMAT(k)= 0. d0
          endif
        enddo
      enddo
```

```
!C===
```

$$\begin{aligned}
 & \text{Diag}_j \phi_j + \sum_{k=\text{Index}[j], k \neq k_s}^{\text{Index}[j+1]-1} \text{Amat}_k \phi_{\text{Item}[k]} \\
 &= \text{Rhs}_j - \text{Amat}_{k_s} \phi_{\text{Item}[k_s]} \\
 &= \text{Rhs}_j - \text{Amat}_{k_s} \phi_{\min} \quad \text{where } \text{Item}[k_s] = 1
 \end{aligned}$$

Column components of boundary nodes (Dirichlet B.C.) are moved to RHS and eliminated for keeping symmetrical feature of the matrix.

Secondary B.C. (Insulated)



$$\int_S \bar{q} [N]^T dS = \bar{q} A|_{x=L} = \bar{q} A \begin{Bmatrix} 0 \\ 1 \end{Bmatrix}, \quad \bar{q} = -\lambda \frac{dT}{dx} \quad \text{Surface Flux}$$



$$\frac{\partial T}{\partial x} = 0 @ x = x_{\max}$$

According to insulated B.C., $\bar{q} = 0$ is satisfied. No contribution by this term. Insulated B.C. is automatically satisfied without explicit operations
 -> Natural B.C.

Preconditioned CG Solver

```

Compute  $\mathbf{r}^{(0)} = \mathbf{b} - [\mathbf{A}]\mathbf{x}^{(0)}$ 
for i= 1, 2, ...
    solve  $[\mathbf{M}]\mathbf{z}^{(i-1)} = \mathbf{r}^{(i-1)}$ 
     $\rho_{i-1} = \mathbf{r}^{(i-1)} \cdot \mathbf{z}^{(i-1)}$ 
    if i=1
         $\mathbf{p}^{(1)} = \mathbf{z}^{(0)}$ 
    else
         $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
         $\mathbf{p}^{(i)} = \mathbf{z}^{(i-1)} + \beta_{i-1} \mathbf{p}^{(i-1)}$ 
    endif
     $\mathbf{q}^{(i)} = [\mathbf{A}]\mathbf{p}^{(i)}$ 
     $\alpha_i = \rho_{i-1} / \mathbf{p}^{(i)} \cdot \mathbf{q}^{(i)}$ 
     $\mathbf{x}^{(i)} = \mathbf{x}^{(i-1)} + \alpha_i \mathbf{p}^{(i)}$ 
     $\mathbf{r}^{(i)} = \mathbf{r}^{(i-1)} - \alpha_i \mathbf{q}^{(i)}$ 
    check convergence  $|\mathbf{r}|$ 
end

```

$$[\mathbf{M}] = \begin{bmatrix} D_1 & 0 & \dots & 0 & 0 \\ 0 & D_2 & & 0 & 0 \\ \dots & & \dots & & \dots \\ 0 & 0 & & D_{N-1} & 0 \\ 0 & 0 & \dots & 0 & D_N \end{bmatrix}$$

Diagonal Scaling, Point-Jacobi

$$[M] = \begin{bmatrix} D_1 & 0 & \dots & 0 & 0 \\ 0 & D_2 & & 0 & 0 \\ \dots & & \dots & & \dots \\ 0 & 0 & & D_{N-1} & 0 \\ 0 & 0 & \dots & 0 & D_N \end{bmatrix}$$

- **solve $[M] \mathbf{z}^{(i-1)} = \mathbf{r}^{(i-1)}$** is very easy.
- Provides fast convergence for simple problems.
- 1d.f, 1d.c

CG Solver (1/6)

```
!C
!C +-----+
!C | CG iterations |
!C +-----+
!C===
      R = 1
      Z = 2
      Q = 2
      P = 3
      DD= 4

      do i= 1, N
        W(i,DD)= 1.0D0 / DIAG(i)
      enddo
```

```
W(i, 1)= W(i, R)    ⇒ {r}
W(i, 2)= W(i, Z)    ⇒ {z}
W(i, 2)= W(i, Q)    ⇒ {q}
W(i, 3)= W(i, P)    ⇒ {p}
W(i, 4)= W(i, DD)   ⇒ 1/{D}
```

```
Compute  $r^{(0)} = b - [A]x^{(0)}$ 
for i= 1, 2, ...
  solve  $[M]z^{(i-1)} = r^{(i-1)}$ 
   $\rho_{i-1} = r^{(i-1)} z^{(i-1)}$ 
  if i=1
     $p^{(1)} = z^{(0)}$ 
  else
     $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
     $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
  endif
   $q^{(i)} = [A]p^{(i)}$ 
   $\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$ 
   $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
   $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
  check convergence |r|
end
```

CG Solver (1/6)

```
!C
!C +-----+
!C | CG iterations |
!C +-----+
!C===
```

```
R = 1
Z = 2
Q = 2
P = 3
DD= 4
```

```
do i= 1, N
  W(i,DD)= 1.0D0 / DIAG(i)
enddo
```

Reciprocal numbers (倒数) of diagonal components are stored in $W(i, DD)$. Computational cost for division is usually expensive.

```
W(i, 1)= W(i, R)   ⇒ {r}
W(i, 2)= W(i, Z)   ⇒ {z}
W(i, 2)= W(i, Q)   ⇒ {q}
W(i, 3)= W(i, P)   ⇒ {p}
W(i, 4)= W(i, DD)  ⇒ 1/{D}
```

CG Solver (2/6)

```
!C
!C-- {r0} = {b} - [A]{xini} |
```

!C 初期残差

```
do i= 1, N
  W(i,R) = DIAG(i)*PHI(i)
  do j= INDEX(i-1)+1, INDEX(i)
    W(i,R) = W(i,R) + AMAT(j)*PHI(ITEM(j))
  enddo
enddo
```

```
BNRM2= 0.0D0
```

```
do i= 1, N
  BNRM2 = BNRM2 + RHS(i) **2
  W(i,R)= RHS(i) - W(i,R)
enddo
```

BNRM2=|b|²
for convergence criteria
of CG solvers

Compute $\mathbf{r}^{(0)} = \mathbf{b} - [\mathbf{A}]\mathbf{x}^{(0)}$

```
for i= 1, 2, ...
  solve [M] z(i-1) = r(i-1)
  ρi-1 = r(i-1) z(i-1)
  if i=1
    p(1) = z(0)
  else
    βi-1 = ρi-1/ρi-2
    p(i) = z(i-1) + βi-1 p(i-1)
  endif
  q(i) = [A]p(i)
  αi = ρi-1/p(i)q(i)
  x(i) = x(i-1) + αip(i)
  r(i) = r(i-1) - αiq(i)
  check convergence |r|
end
```

CG Solver (3/6)

```

do iter= 1, ITERmax

!C
!C-- {z}= [Minv] {r}

do i= 1, N
  W(i, Z)= W(i, DD) * W(i, R)
enddo

!C
!C-- RHO= {r} {z}

RHO= 0. d0
do i= 1, N
  RHO= RHO + W(i, R)*W(i, Z)
enddo

```

```

Compute  $r^{(0)} = b - [A] x^{(0)}$ 
for i= 1, 2, ...
  solve  $[M] z^{(i-1)} = r^{(i-1)}$ 
   $\rho_{i-1} = r^{(i-1)} z^{(i-1)}$ 
  if i=1
     $p^{(1)} = z^{(0)}$ 
  else
     $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
     $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
  endif
   $q^{(i)} = [A] p^{(i)}$ 
   $\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$ 
   $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
   $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
  check convergence  $|r|$ 
end

```

CG Solver (4/6)

```

!C
!C-- {p} = {z} if      ITER=1
!C  BETA= RHO / RH01  otherwise

      if ( iter.eq.1 ) then
        do i= 1, N
          W(i,P)= W(i,Z)
        enddo
      else
        BETA= RHO / RH01
        do i= 1, N
          W(i,P)= W(i,Z) + BETA*W(i,P)
        enddo
      endif

!C
!C-- {q}= [A] {p}

      do i= 1, N
        W(i,Q) = DIAG(i)*W(i,P)
        do j= INDEX(i-1)+1, INDEX(i)
          W(i,Q) = W(i,Q) + AMAT(j)*W(ITEM(j),P)
        enddo
      enddo

```

```

Compute  $r^{(0)} = b - [A]x^{(0)}$ 
for i= 1, 2, ...
  solve  $[M]z^{(i-1)} = r^{(i-1)}$ 
   $\rho_{i-1} = r^{(i-1)} z^{(i-1)}$ 
  if i=1
     $p^{(1)} = z^{(0)}$ 
  else
     $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
     $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
  endif
   $q^{(i)} = [A]p^{(i)}$ 
   $\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$ 
   $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
   $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
  check convergence  $|r|$ 
end

```

CG Solver (5/6)

```

!C
!C-- ALPHA= RHO / {p} {q}

C1= 0.d0
do i= 1, N
  C1= C1 + W(i,P)*W(i,Q)
enddo
ALPHA= RHO / C1

!C
!C-- {x}= {x} + ALPHA*{p}
!C  {r}= {r} - ALPHA*{q}

do i= 1, N
  PHI(i)= PHI(i) + ALPHA * W(i,P)
  W(i,R)= W(i,R) - ALPHA * W(i,Q)
enddo

```

```

Compute  $r^{(0)} = b - [A] x^{(0)}$ 
for i= 1, 2, ...
  solve  $[M] z^{(i-1)} = r^{(i-1)}$ 
   $\rho_{i-1} = r^{(i-1)} z^{(i-1)}$ 
  if i=1
     $p^{(1)} = z^{(0)}$ 
  else
     $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
     $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
  endif
   $q^{(i)} = [A] p^{(i)}$ 
   $\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$ 
   $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
   $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
  check convergence  $|r|$ 
end

```

CG Solver (6/6)

```

DNRM2 = 0.0
do i= 1, N
  DNRM2= DNRM2 + W(i,R)**2
enddo

RESID= dsqrt (DNRM2/BNRM2)

  if ( RESID.le.EPS) goto 900
  RH01 = RHO  ρi-2

enddo
900 continue

```

$$\text{Resid} = \sqrt{\frac{\text{DNorm2}}{\text{BNorm2}}} = \frac{|r|}{|b|} = \frac{|Ax - b|}{|b|} \leq \text{Eps}$$

Control Data input.dat

```

4          NE (Number of Elements)
1.0 1.0 1.0 1.0 Δx (Length of Each Elem.: L), Q, A, λ
100        Number of MAX. Iterations for CG Solver
1.e-8      Convergence Criteria for CG Solver

```

```

Compute  $r^{(0)} = b - [A] x^{(0)}$ 
for i= 1, 2, ...
  solve  $[M] z^{(i-1)} = r^{(i-1)}$ 
   $\rho_{i-1} = r^{(i-1)} z^{(i-1)}$ 
  if i=1
     $p^{(1)} = z^{(0)}$ 
  else
     $\beta_{i-1} = \rho_{i-1} / \rho_{i-2}$ 
     $p^{(i)} = z^{(i-1)} + \beta_{i-1} p^{(i-1)}$ 
  endif
   $q^{(i)} = [A] p^{(i)}$ 
   $\alpha_i = \rho_{i-1} / p^{(i)} q^{(i)}$ 
   $x^{(i)} = x^{(i-1)} + \alpha_i p^{(i)}$ 
   $r^{(i)} = r^{(i-1)} - \alpha_i q^{(i)}$ 
  check convergence |r|
end

```

$$Ax = b \Rightarrow \alpha Ax = \alpha b$$

$$r = b - Ax \Rightarrow R = \alpha b - \alpha Ax = \alpha r$$

Finite Element Procedures

- Initialization
 - Control Data
 - Node, Connectivity of Elements (N: Node#, NE: Elem#)
 - Initialization of Arrays (Global/Element Matrices)
 - Element-Global Matrix Mapping (Index, Item)
- Generation of Matrix
 - Element-by-Element Operations (do icel= 1, NE)
 - Element matrices
 - Accumulation to global matrix
 - Boundary Conditions
- Linear Solver
 - Conjugate Gradient Method

Remedies for Higher Accuracy

- Finer Meshes

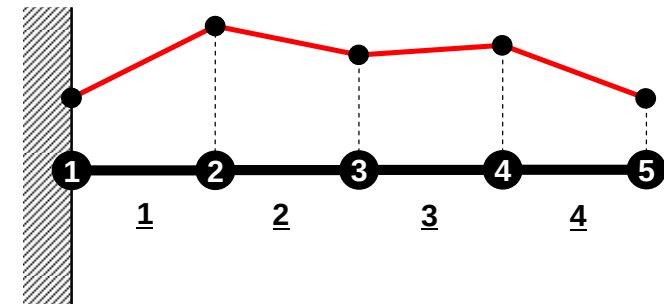
NE=8, dx=12.5

8 iters, RESID= 2.822910E-16 U(N)= 1.953586E-01

DISPLACEMENT

1	0.000000E+00	-0.000000E+00
2	1.101928E-02	1.103160E-02
3	2.348034E-02	2.351048E-02
4	3.781726E-02	3.787457E-02
5	5.469490E-02	5.479659E-02
6	7.520772E-02	7.538926E-02
7	1.013515E-01	1.016991E-01
8	1.373875E-01	1.381746E-01
9	1.953586E-01	1.980421E-01

$$\therefore u = \frac{F}{EA_1} [\log(A_1 x + A_2) - \log(A_2)]$$



NE=20, dx=5

20 iters, RESID= 5.707508E-15 U(N)= 1.975734E-01

DISPLACEMENT

1	0.000000E+00	-0.000000E+00
2	4.259851E-03	4.260561E-03
3	8.719160E-03	8.720685E-03
4	1.339752E-02	1.339999E-02
.....		
17	1.145876E-01	1.146641E-01
18	1.295689E-01	1.296764E-01
19	1.473466E-01	1.475060E-01
20	1.692046E-01	1.694607E-01
21	1.975734E-01	1.980421E-01

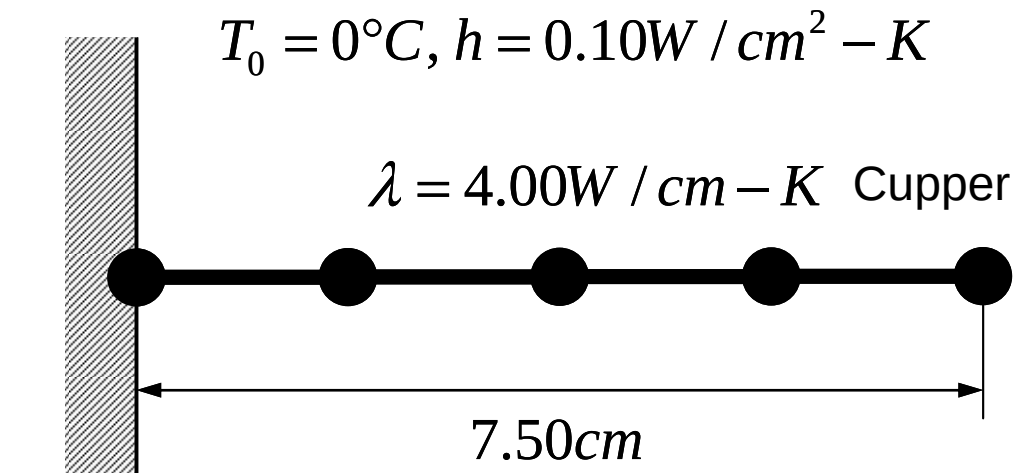
Remedies for Higher Accuracy

- Finer Meshes
- Higher Order Shape/Interpolation Function (高次補間関数・形状関数)
 - Higher-Order Element (高次要素)
 - Linear-Element, 1st-Order Element: Lower Order (低次要素)
- Formulation which assures continuity of n-th order derivatives
 - C^n Continuity (C^n 連続性)

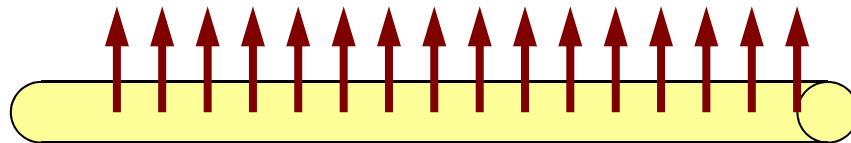
Remedies for Higher Accuracy

- Finer Meshes
- Higher Order Shape/Interpolation Function (高次補間関数・形状関数)
 - Higher-Order Element (高次要素)
 - Linear-Element, 1st-Order Element: Lower Order (低次要素)
- Formulation which assures continuity of n-th order derivatives
 - C^n Continuity (C^n 連続性)
- Linear Elements
 - Piecewise Linear
 - C^0 Continuity
 - Only dependent variables are continuous at element boundary

Example: 1D Heat Transfer (1/2)



$T_s = 150^\circ\text{C}$



Convective Heat Transfer on
Cylindrical Surface

- Temp. Thermal Fins
- Circular Sectional Area, $r=1\text{cm}$
- Boundary Condition
 - $x=0$: Fixed Temperature
 - $x=7.5$: Insulated
- Convective Heat Transfer on Cylindrical Surface
 - $q = h(T - T_0)$
 - q : Heat Flux
 - Heat Flow/Unit Surface Area/sec.

Example: 1D Heat Transfer (2/2)

RESULTS (linear interpolation)

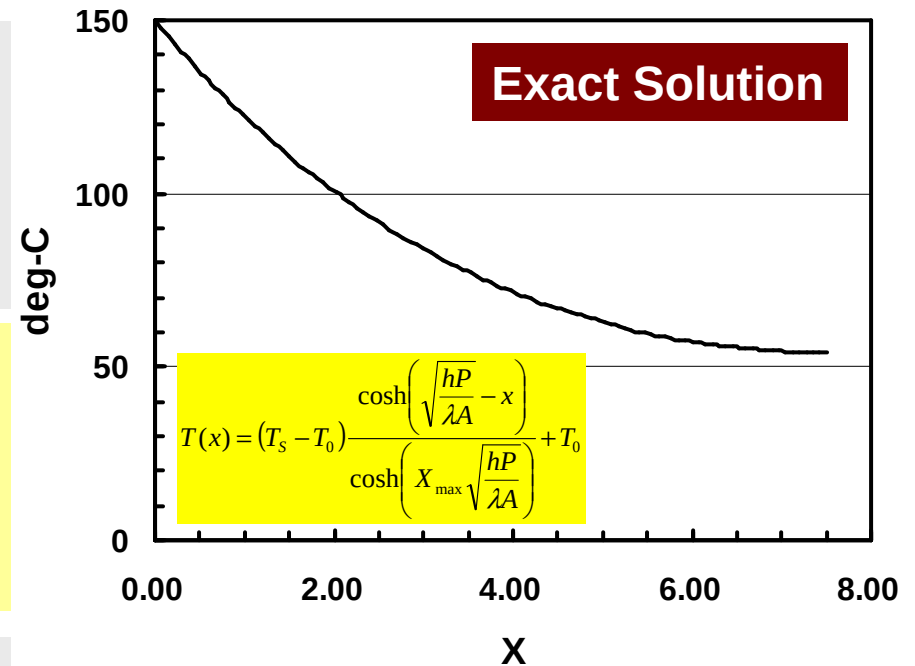
ID	X	FEM.	ANALYTICAL	ERR (%)
1	0.00000	150.00000	150.00000	0.00000
2	1.87500	102.62226	103.00165	0.25292
3	3.75000	73.82803	74.37583	0.36520
4	5.62500	58.40306	59.01653	0.40898
5	7.50000	53.55410	54.18409	0.41999

RESULTS (quadratic interpolation)

ID	X	FEM.	ANALYTICAL	ERR (%)
1	0.00000	150.00000	150.00000	0.00000
2	1.87500	102.98743	103.00165	0.00948
3	3.75000	74.40203	74.37583	0.01747
4	5.62500	59.02737	59.01653	0.00722
5	7.50000	54.21426	54.18409	0.02011

RESULTS (linear interpolation)

ID	X	FEM.	ANALYTICAL	ERR (%)
1	0.00000	150.00000	150.00000	0.00000
2	0.93750	123.71561	123.77127	0.03711
3	1.87500	102.90805	103.00165	0.06240
4	2.81250	86.65618	86.77507	0.07926
5	3.75000	74.24055	74.37583	0.09019
6	4.68750	65.11151	65.25705	0.09703
7	5.62500	58.86492	59.01653	0.10107
8	6.56250	55.22426	55.37903	0.10317
9	7.50000	54.02836	54.18409	0.10382



Quadratic interpolation provides more accurate solution, especially if X is close to 7.50cm.