

Partial Differential Equations in Modelica

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Chapter 1

Modelica extension for PDE

1.1 Requests on language extension and possible approaches

Space & coordinates

What should be specified

- Dimension of the problem (1,2 or 3D)
- ?? Coordinates (cartesian, cylindrical, spherical ...) – where this information will be used (if at all):
 - in differential operators as grad, div, rot etc.
 - in visualization of results
 - ?? in computation – perhaps equations should be transformed and the calculation would be performed in cartesian coordinates
- Names of independent (coordinate) variables ($x, y, z, r, \varphi, \theta...$)

Perhaps all these should be specified within the domain definition.

Dimension can be inferred from number of return values of shape-function or different properties of the domain in other cases.

The base coordinates would be cartesian and they would be always implicitly defined in any domain. Besides that other coordinate systems could be defined also.

Names of independent variables in cartesian coordinates should be fixed $x, (x,y), (x,y,z)$ in 1D, 2D and 3D domains respectively.

Domain & boundary

What should be specified

- the domain where we perform the computation and where equations hold
- boundary and its subsets where particular boundary conditions hold
- normal vector of the boundary

Possible approaches

Parametrization of the domain with shape function and intervals – from
The Book (Principles of ...), section 8.5.2

Example from the book:

```
model HeatCircular2D
    import DifferentialOperators2D.*;
    parameter DomainCircular2DGrid omega;
    FieldCircular2DGrid u(domain=omega, FieldValueType=SI.Temperature);
equation
    der (u) = pder (u,D.x2)+ pder (u,D . y 2 )      in omega.interior;
    nder(u) = 0                                     in omega.boundary;
end HeatCircular2D;

record DomainCircular2DGrid "Domain being a circular region"
    parameter Real radius = 1;
    parameter Integer nx = 100;
    parameter Integer ny = 100;
    replaceable function shapeFunc = circular2Dfunc "2D circular region";
    DomainGe2D interior(shape=shapeFunc,interval={{O,radius},{O,1}},geom= ..
    DomainGe2D boundary (shape=shapeFunc, interval={{radius, radius), { 0,1}
    function shapeFunc = circular2Dfunc "Function spanning circular region";
end DomainCircular2DGrid;

function circular2Dfunc "Spanned circular region for v in interval 0..1"
    input Real r,v;
    output Real x,y;
algorithm
    x := r*cos (2*PI*v);
    y := r*sin(2*PI*v);
end circular2Dfunc;

record FieldCircular2DGrid
    parameter DomainCircular2DGrid domain;
```

```

        replaceable type FieldValueType = Real;
        replaceable type FieldType = Real[domain.nx, domain.ny, domain.nz];
        parameter FieldType start = zeros(domain.nx, domain.ny, domain.nz.);
        FieldType Val;
    end FieldCircularZDGrid;

```

And modified version, where all numerical stuff (grid, number of points – this should be configured using simulation setup or annotations) omitted, modified `pder` operator, `Field` as Modelica build-in type:

```

model HeatCircular2D
    parameter DomainCircular2D omega(radius=2);
    field Real u(domain=omega, start = 0, FieldValueType=SI.Temperature);
equation
    pder(u,time) = pder(u,x)+ pder(u,y) in omega.interior;
    pder(u,omega.boundary.n) = 0 in omega.boundary;
end HeatCircular2D;

record DomainCircular2D
    parameter Real radius = 1;
    parameter Real cx = 0;
    parameter Real cy = 0;
    function shapeFunc
        input Real r,v;
        output Real x,y;
    algorithm
        x := cx + radius*r * cos(2 * C.pi * v);
        y := cy + radius*r * sin(2 * C.pi * v);
    end shapeFunc;
    Region2D interior(shape = shapeFunc, interval = {{0,1},{0,1}});
    Region1D boundary(shape = shapeFunc, interval = {{1,1},{0,1}});
end DomainCircular2D;

```

Description by the boundary Domain is defined by closed boundary curve, which may be composed of several connected curves. Needs new operator *interior* and type *Domain2d* (and *Domain1D* and *Domain3d*). (similarly used in FlexPDE – <http://www.pdesolutions.com/>.)

```

package BoundaryRepresentation
    partial function cur
        input Real u;
        output Real x;
        output Real y;
    end cur;
    function arc

```

```

    extends cur;
    parameter Real r;
    parameter Real cx;
    parameter Real cy;
algorithm
    x:=cx + r * cos(u);
    y:=cy + r * sin(u);
end arc;
function line
    extends cur;
    parameter Real x1;
    parameter Real y1;
    parameter Real x2;
    parameter Real y2;
algorithm
    x:=x1 + (x2 - x1) * u;
    y:=y1 + (y2 - y1) * u;
end line;
function bezier3
    extends cur;
    //start-point
    parameter Real x1;
    parameter Real y1;
    //end-point
    parameter Real x2;
    parameter Real y2;
    //start-control-point
    parameter Real cx1;
    parameter Real cy1;
    //end-control-point
    parameter Real cx2;
    parameter Real cy2;
algorithm
    x:=(1 - u) ^ 3 * x1 + 3 * (1 - u) ^ 2 * u * cx1 + 3 *
        (1 - u) * u ^ 2 * cx2 + u ^ 3 * x2;
    y:=(1 - u) ^ 3 * y1 + 3 * (1 - u) ^ 2 * u * cy1 + 3 *
        (1 - u) * u ^ 2 * cy2 + u ^ 3 * y2;
end bezier3;
record Curve
    function curveFun = line;
    // to be replaced with another fun
    parameter Real uStart;
    parameter Real uEnd;
end Curve;
record Boundary
    constant Integer NCurves;

```

```

Curve curves[NCurves];
// for i in 1:(NCurves-1) loop
// assert(Curve[i].curveFun(Curve[i].uEnd) = Curve[i
+1].curveFun(Curve[i+1].uStart), String(i)+"th
curve and "+String(i+1)+"th curve are not
connected.", level = AssertionLevel.error);
// end for;
// assert(curves[NCurves].curveFun(curves[NCurves
].uEnd) =
// curves[1].curveFun(curves
[1].uStart),
// String(NCurves)+"th curve
and first curve are not connected.",
// level = AssertionLevel.
error);
end Boundary;
record DomainHalfCircle
constant Real pi = Modelica.Constants.pi;
arc myArcFun(cx = 0, cy = 0, r = 1);
Curve myArc(curveFun = myArcFun, uStart = pi / 2,
uEnd = (pi * 3) / 2);
line myLineFun(x1 = 0, y1 = -1, x2 = 0, y2 = 1);
Curve myLine(curveFun = myLineFun, uStart = 0, uEnd =
1);
line myLine2(curveFun = line(x1 = 0, y1 = -1, x2 = 0,
y2 = 1), uStart = pi / 2, uEnd = (pi * 3) / 2);
Boundary b(NCurves = 2, curves = {myArc, myLine});
//new externally defined type Domain2D and operator
interior:
Domain2D d = interior Boundary;
end DomainHalfCircle;
end BoundaryRepresentation;

```

Constructive solid geometry used in Matlab PDE toolbox, http://en.wikipedia.org/wiki/Constructive_sol

Domain is build from primitives (cuboids, cylinders, spheres, cones, user defined shapes ...) applying boolean operations *union*, *intersection* and *difference*.

How to describe boundaries?

Listing of points – export from CAD

Inequalities

Boundary representation (BRep) (NETGEN, STEP)

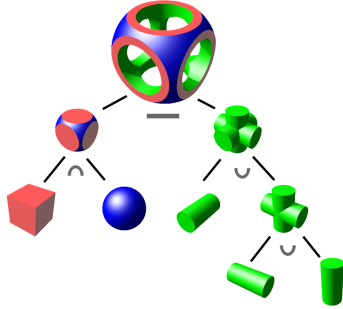


Figure 1.1: constructive solid geometry

Fields

Partial derivative

$\frac{\partial^2 u}{\partial x \partial y}$... `pder(u,x,y)`
directional derivative ... `pder(u,omega.boundary.n)`

Equations, boundary and initial conditions

Use the *in* operator to express where equations hold.

1.2 New concepts and language elements

domain records (or class) `DomainLineSegment1D`, `DomainRectangle2D`, `DomainCircular2D`, `DomainBlock3D` ... defined within PDE package, user can define his/her own domain records. Domain record contains at least one region member . During translation is treated in different way than usual records. Needs OMC modification.

interval to define parameter interval for a shape-function. E.g. `interval={{0,1},0}`.
Used in domain records. (Previously called range.)
New language element.

shape function one-to-one map of points in k-dimensional interval (usually cartesian product of intervals) to points in n-dimensional domain and thus define a coordinate system in domain .

region types and regions `Region0D`, `Region1D`, `Region2D`, `Region3D` used in domain records to define interior, boundaries and other regions where certain equations hold (e.g. connection of PDE and ODE). Two mandatory attributes are `shape` and `interval`. E.g. `Region2D left (shape =`

`shapeFunc, interval = {0,{0,1}}).`
 New language element.

normal vector implicitly defined for all N-1 dimensional regions in N dimensional domain. (e.g. `omega.left.n`) Used in boundary condition equations. How to write domain and region independently - perhaps `region.n`?
 New language element.

fields A variable whose value depends on space position, is called field. It is defined with keyword `field`. Field can be of any type. It can be defined also as a parameter. Field may be an array to represent vector field. Mandatory attribute is `domain`. Other attributes are same as for corresponding regular type (e.g. for `Real`: `start`, `fixed`, `nominal`, `min`, `max`, `unit`, `displayUnit`, `quantity`, `stateSelect`. (Not shure about `fixed` and `stateSelect`.) Attribute `start` can be assigned constant value or function of type $(\text{Real} \times \text{Real} \times \dots) \rightarrow \text{typeOfField}$, number of arguments is same as dimension of fields domain. Start attribute can be treated as scalar or as array to assign initial values also to derivatives. E.g.
`field Real x(domain = omega, start[0] = xInit, start[1] = 0)`
 see 3.2.2
 New language element.

operations and functions on fields All operators (`=`, `:=`, `+`, `-`, `*`, `/`, `^`, `<`, `<=`, `>`, `>=`, `==`, `<>`) and functions can be applied on fields. The result is also a field. If a binary operator or function of more arguments is applied on two (or more) fields, these fields must be defined within the same domain.

If some binary operator or function with more arguments is performed on field and regular variable (it means a variable that is not a field), the operation is performed as if the regular variable is field that is constant in space.

`pder()` **operator** for partial and directional derivative of real field. Higher derivatives are allowed. E.g. `pder(u,omega.x,omega.x,omega.y)` means $\frac{\partial^3 u}{\partial x^2 y}$. Directional derivative: `pder(u,omega.left.n)`. Time derivative of field must be written also using `pder` operator not `der`.

in operator to define where PDE, boundary conditions and other equations hold. On left is an equation on right is a region where the equation hold.
 E.g. `x=0 in omega.left`
 New language element.

dom keyword stands for current domain specified with `in` operator.
`pder(u,dom.x)=0 in omega.interior;`
 is equivalent to
`pder(u,omega.x)=0 in omega.interior;`
 Is useful to write equations domain independent.
 New language element.

region keyword stands for current region specified with **in** operator.

```
v*region.n=0 in omega.left + omega.right;
```

is equivalent to

```
v*omega.left.n=0 in omega.left; v*omega.right.n=0 in omega.right;
```

Is useful to get normal vector in domain independent equations and on anonymous regions.

New language element. Not shure if it will be needed.

region addition **+** operator can be used to add regions. Can be used in domain record to form a new region, e.g.

```
boundaries = left + right;
```

or on right side of **in** operator, e.g.

```
x = 0 in omega.left + omega.right;
```

New language element. Not shure if it will be needed.

vector differential operators **grad**, **div**, **rot**

Coordinate type to define new coordinates - design of concept unfinished.

New language element.

Chapter 2

Numerics

Goals

1. advection equation in 1D and eulerian coordinate, dirichlet BC, explicit solver
2. numann BC
3. automatic dt
4. diffusion or mixed equation
5. implicit solver
6. systems of equations
7. 2D (rectangle), 3D (cube)
8. lagrangian coordinate
9. general domain

difference schemes separated from the rest of solver

Difussion eq:

$$u_t = \alpha u_{xx}$$

or

$$\begin{aligned} u_t &= -w_x \\ w &= -\alpha u_x. \end{aligned}$$

String eq:

$$y_{tt} = ky_{xx}$$

or

$$\begin{aligned} s_x &= kv_t \\ y_t &= v \\ y_x &= s \end{aligned}$$

The description without higher derivative is ugly, we need higher derivatives.

Representation

Explicit

$$u_t = f(u, u_x, t) \quad (2.1)$$

resp.

$$u_t = f(u, u_x, u_{xx}, \dots, t)$$

..

Implicit

$$F(u, u_t, u_x, t) = 0 \quad (2.2)$$

resp.

$$F(u, u_t, u_x, u_{xx}, \dots, t) = 0$$

Solvers

Difference schemes for explicit solver

U denotes discretized u

Time difference from Lax-Friedrichs in explicit form (i.e. with the u_m^{n+1} on LHS):

$$u_m^{n+1} = D_t^{exp}(v, U, n, m) = v\Delta t + \frac{1}{2}(u_{m+1}^n + u_{m-1}^n) \quad (2.3)$$

Space difference from Lax-Friedrichs:

$$D_x(U, n, m) = \frac{u_{i+1}^n - u_{i-1}^n}{2\Delta x} \quad (2.4)$$

Explicit solver Lax-Friedrichs

We solve equation (2.1) substituting space difference (2.4) and applying time difference in explicit form (2.3):

$$\begin{aligned} u_m^{n+1} &= D_t^{exp}(f((u_m^n, D_x(U, n, m), t^n))) = \\ &= \Delta t \cdot f(u, \frac{u_{i+1}^n - u_{i-1}^n}{2\Delta x}, t) + \frac{1}{2}(u_{m+1}^n + u_{m-1}^n) \end{aligned}$$

Difference schemes for implicit solver space difference from Crank-Nicolson

$$\begin{aligned} D_x(u_{m-1}^n, u_m^n, u_{m+1}^n, u_{m-1}^{n+1}, u_m^{n+1}, u_{m+1}^{n+1}) &= \frac{1}{2} \left(\frac{u_{m+1}^{n+1} - u_{m-1}^{n+1}}{2\Delta x} + \frac{u_{m+1}^n - u_{m-1}^n}{2\Delta x} \right) \\ D_{xx}(u_{m-1}^n, u_m^n, u_{m+1}^n, u_{m-1}^{n+1}, u_m^{n+1}, u_{m+1}^{n+1}) &= \frac{1}{2} \left(\frac{u_{m+1}^{n+1} - 2u_m^{n+1} + u_{m-1}^{n+1}}{2(\Delta x)^2} + \frac{u_{m+1}^n - 2u_m^n + u_{m-1}^n}{2(\Delta x)^2} \right) \end{aligned} \quad (2.5)$$

time difference from Crank-Nicolson

$$D_t(u_{m-1}^n, u_m^n, u_{m+1}^n, u_{m-1}^{n+1}, u_m^{n+1}, u_{m+1}^{n+1}) = \frac{u_m^{n+1} - u_m^n}{\Delta t} \quad (2.6)$$

Implicit solver Crank-Nicolson

With nonlinear solver:

We solve equation (2.2) substituting space (2.5) and time (2.6) differences

$$F(u_m^n, D_t(u_{m-1}^n, \dots), D_x(u_{m-1}^n, \dots), t^n) = 0, \quad m \in \hat{M} \quad (2.7)$$

and than solving the whole system for all unknown u^{n+1} . System has 3-band Jacobian. If F is linear in u_x and u_t , system is also linear with 3-band matrix eventhou is given generaly. Is there any solver efficient in solving linear equations with banded matrix given implicitly? (I hope Newton-Raphson is.) As initial guess for the solution we can use extrapolated values. If solving fails we can try value from the node on left or right (this could help on shocks).

With linear solver:

If F is linear, we expres (2.7) as

$$\mathbf{A}\bar{\mathbf{u}}^{n+1} = \bar{\mathbf{b}}.$$

\mathbf{A} is $M \times M$ 3-diagonal. Functions for evaluation of \mathbf{M} and $\bar{\mathbf{b}}$ are generated during compilation. In runtime we solve just the linear system. In this aproach difference schema must be chosen before compilation of model.

Implicit solver and systems of PDE If we solve e.g. system with three variables u, v, w , se can sort difference equations in order

$$u_1, v_1, w_1, u_2, v_2, w_2, u_3, v_3, w_3, \dots$$

so that the system is stil banded.

Chapter 3

Example models

3.1 Package PDEDomains

Modelica code of domain definitions:

```
package PDEDomains
import C = Modelica.Constants;
record DomainLineSegment1D
  parameter Real l = 1;
  parameter Real a = 0;
  function shapeFunc
    input Real v;
    output Real x = l*v + a;
  end shapeFunc;
  Coordinate x (name = "cartesian");
  Region1D interior(shape = shapeFunc, interval =
    {0,1});
  Region0D left(shape = shapeFunc, interval = 0);
  Region0D right(shape = shapeFunc, interval = 1);
  Region0D boundary = left + right;
end DomainLineSegment1D;

class DomainRectangle2D
  parameter Real Lx = 1;
  parameter Real Ly = 1;
  parameter Real ax = 0;
  parameter Real ay = 0;
  function shapeFunc
    input Real v1, v2;
    output Real x = ax + Lx * v1, y = ay + Ly * v2;
  end shapeFunc;
```

```

Coordinate x (name = "cartesian");
Coordinate y (name = "cartesian");

Coordinate r (name = "polar");
Coordinate theta (name = "polar");

equation
  r = sqrt(x^2 + y^2);
  theta = arctg(y/x);

Region2D interior(shape = shapeFunc, interval =
  {{0,1},{0,1}});
Region1D right(shape = shapeFunc, interval =
  {1,{0,1}});
Region1D bottom(shape = shapeFunc, interval =
  {{0,1},0});
Region1D left(shape = shapeFunc, interval =
  {0,{0,1}});
Region1D top(shape = shapeFunc, interval = {{0,1},1})
;
Region1D boundary = right + bottom + left + top;
end DomainRectangle2D;

class DomainCircular2D
  parameter Real radius = 1;
  parameter Real cx = 0;
  parameter Real cy = 0;
  function shapeFunc
    input Real r,v;
    output Real x,y;
  algorithm
    x:=cx + radius * r * cos(2 * C.pi * v);
    y:=cy + radius * r * sin(2 * C.pi * v);
  end shapeFunc;

Coordinate x (name="cartesian");
Coordinate y (name="cartesian");

Coordinate r (name="polar");
Coordinate theta (name="polar");

equation
  r = sqrt(x^2 + y^2);
  theta = arctg(y/x);
end polar;

```

```

    Region2D interior(shape = shapeFunc, interval = {{0
    ,1},{0,1}});
    Region1D boundary(shape = shapeFunc, interval =
    {1,{0,1}});
end DomainCircular2D;

record DomainBlock3D
    parameter Real Lx = 1, Ly = 1, Lz = 1;
    parameter Real ax = 0, ay = 0, az = 0;
    function shapeFunc
        input Real vx, vy, vz;
        output Real x = ax + Lx * vx, y = ay + Ly * vy, z =
            az + Lz * vz;
    end shapeFunc;
    Coordinate x (name="cartesian");
    Coordinate y (name="cartesian");
    Coordinate z (name="cartesian");
    Region3D interior(shape = shapeFunc, interval =
        {{0,1},{0,1},{0,1}});
    Region2D right(shape = shapeFunc, interval =
        {1,{0,1},{0,1}});
    Region2D bottom(shape = shapeFunc, interval =
        {{0,1},{0,1},1});
    Region2D left(shape = shapeFunc, interval =
        {0,{0,1},{0,1}});
    Region2D top(shape = shapeFunc, interval =
        {{0,1},{0,1},1});
    Region2D front(shape = shapeFunc, interval =
        {{0,1},0,{0,1}});
    Region2D rear(shape = shapeFunc, interval =
        {{0,1},1,{0,1}});
    Region2D boundary = right + bottom + left + top +
        front + rear;
end DomainBlock3D;
//and others ...
end PDEDomains;

```

Listing 3.1: Standard domains definitions: 1D – Line segment, 2D – Rectangle, Circle, 3D – Block

3.2 Simple models

3.2.1 Advection equation (1D)[13]

$L \dots$ length

c .. constant, assume $c > 0$
 $u \in \langle 0, L \rangle \times \langle 0, T \rangle \rightarrow \mathbb{R}$

equation

$$\frac{\partial u}{\partial t} + c \frac{\partial u}{\partial x} = 0$$

initial conditions

$$u(x, 0) = 1$$

boundary conditions

$$u(0, t) = \cos(2\pi t)$$

Modelica code

```

model advection "advection equation"
  import C = Modelica.Constants;
  parameter Real L = 1; // length
  parameter Real c = 1;
  parameter DomainLineSegment1D omega(length = L);
  field Real u(domain = omega, start = 1);
equation
  pder(u, time) + c*pder(u, dom.x) = 0 in omega.
    interior;
  u = cos(2*C.pi*time) in omega.left;
end advection;

```

Listing 3.2: Advection equation in Modelica

Flat model

```

/*TODO: finish it!!*/

model advection_flat "advection equation"
  import C = Modelica.Constants;
  parameter Real L = 1; // length
  parameter Real c = 1;
  // parameter DomainLineSegment1D omega(length = L
  );
  parameter Real DomainLineSegment1D.l = L;
  parameter Real DomainLineSegment1D.a = 0;
  function DomainLineSegment1D.shapeFunc

```

```

    input Real v;
    output Real x = 1*v + a;
end DomainLineSegment1D.shapeFunc;
Domain1DInterior DomainLineSegment1D.interior(
    shape = shapeFunc, range = {0,1});
Domain1DBoundary DomainLineSegment1D.left(shape
    = shapeFunc, range = {0,0});
Domain1DBoundary DomainLineSegment1D.right(shape
    = shapeFunc, range = {1,1});

field Real u(domain = omega, start = 1);
equation
    pder(u,time) + c*pder(u,x) = 0 in omega.
    interior;
    u = cos(2*pi*time) in omega.left;
end advection_flat;

```

Listing 3.3: Advection equation – flat model

Generated C code

```

#include <math.h>
#include "model_data.h"
#include "PDESolver.h"
#include "model.h"

// #define _USE_MATH_DEFINES
// #include <math.h>
double pi = 3.14159265358979323846;

int setupArrayDimensions(struct MODEL_DATA* mData)
{
    mData->nStateFields = 1;
    mData->nAlgebraicFields = 0;
    mData->nParameterFields = 0;
    mData->nParameters = 4;
    mData->nDomainSegments = 3;
    return 0;
}

int setupModelParameters(struct MODEL_DATA* mData)

```

```

{
    /* interior: */
    mData->domainRange[0].v0 = 0;
    mData->domainRange[0].v1 = 1;
    /* left */
    mData->domainRange[1].v0 = 0;
    mData->domainRange[1].v1 = 0;
    /* right */
    mData->domainRange[2].v0 = 1;
    mData->domainRange[2].v1 = 1;
    /* advection.L */ mData->parameters[0] = 1;
    /* advection.c */ mData->parameters[1] = 1;
    /* DomainLineSegment1D.l */ mData->parameters[2]
        = 1;
    /* DomainLineSegment1D.a */ mData->parameters[3]
        = 0;
    mData->isBc[mData->nStateFields*0 + 0] = 1;
    mData->isBc[mData->nStateFields*0 + 1] = 0;
    return 0;
}

int setupInitialState(struct MODEL_DATA* mData){
    int i;
    for (i=0; i<mData->M; i++){
        //TODO: should be done generally, with
        some kind of stateInitial(x) function
        called from static code.
        mData->stateFields[mData->M*0 + i] = 1;
    }
    return 0;
}

double shapeFunction(struct MODEL_DATA *mData,
    double v)
{
    return /*DomainLineSegment1D.l */ mData->
        parameters[2]*v + /*DomainLineSegment1D.a */
        mData->parameters[3];
}

int functionPDE(struct MODEL_DATA *mData, int
    dScheme)
{
    int M = mData->M;
    int i;

```

```

    for (i = 0; i < M; i++)
        /*u_t*/mData->stateFieldsDerTime[M*0 + i]
            = - /*c*/mData->parameters[1] * /*u_x*/
                mData->stateFieldsDerSpace[mData->M*0 +
                    i];
    return 0;
}

int functionBC(struct MODEL_DATA *mData)
{
    //should be written generally -- independent on
    //particular grid
    int M = mData->M;
    mData->stateFields[M*0 + 0] = cos(2*pi*mData->
        time);
    return 0;
}

/*double eqSystemMaxEigenVal(struct MODEL_DATA*
    mData){
    return cmData->parameters[1];
}*/

```

Listing 3.4: Advection equation – "generated" C code

3.2.2 String equation (1D)[19]

L .. length
 $u \in \langle 0, L \rangle \times \langle 0, T \rangle \rightarrow \mathbb{R}$ (string position)
 c .. constant
equation:

$$\frac{\partial^2 u}{\partial t^2} - c \frac{\partial^2 u}{\partial x^2} = 0$$

initial conditions

$$\begin{aligned} u(x, 0) &= \sin\left(\frac{4\pi}{L}x\right) \\ \dot{u}(x, 0) &= 0 \end{aligned}$$

boundary conditions

$$u(0, t) = 0, \quad u(L, t) = 0$$

Modelica code

```

model string "model of a vibrating string with fixed ends
"
    import C = Modelica.Constants;
    parameter Real L = 1; // length
    parameter Real c = 1; // tension/(linear density)
    parameter DomainLineSegment1D omega(length = L);
    function u0
        input Real x;
        output Real u0 := sin(4*C.pi/L*x);
    end u0;
    field Real u(domain = omega, start[0] = u0, start[1] =
        0);
equation
    pder(u,time,time) - c*pder(u,x,x) = 0    in omega.
        interior;
    u = 0;                                     in omega.left +
        omega.right;
end string;

```

Listing 3.5: String model in Modelica

Generated C code

```

#include <math.h>
#include "model_data.h"
#include "PDESolver.h"
#include "model.h"
#include "diff.h"

double pi = 3.14159265358979323846;

int setupArrayDimensions(struct MODEL_DATA* mData) {
    mData->nStateFields = 2;
    mData->nAlgebraicFields = 1;
    mData->nParameterFields = 0;
    mData->nParameters = 4;
    mData->nDomainSegments = 3;
    return 0;
}

int setupModelParameters(struct MODEL_DATA* mData)
{
    /* interior */
    mData->domainRange[0].v0 = 0;
    mData->domainRange[0].v1 = 1;
    /* left */

```

```

mData->domainRange[1].v0 = 0;
mData->domainRange[1].v1 = 0;
/* right */
mData->domainRange[2].v0 = 1;
mData->domainRange[2].v1 = 1;
/* string.L */ mData->parameters[0] = 1;
/* string.c */ mData->parameters[1] = 1;
/* DomainLineSegment1D.l */ mData->parameters[2] = 1;
/* DomainLineSegment1D.a */ mData->parameters[3] = 0;
mData->isBc[mData->nStateFields*0 + 0] = 1;
mData->isBc[mData->nStateFields*0 + 1] = 1;
return 0;
}

double /*u0*/function_0(struct MODEL_DATA* mData, double
x){
    return sin(4*pi/ /*string.L*/mData->parameters[0]*x);
}

int setupInitialState(struct MODEL_DATA* mData){
    int i;
    for (i=0; i<mData->M; i++){
        /*u*/mData->stateFields[mData->M*0 + i] =
            function_0(mData, mData->spaceField[mData->M*0
+ i]);
        /*u_t*/mData->stateFields[mData->M*1 + i] = 0;
    }
    return 0;
}

double shapeFunction(struct MODEL_DATA *mData, double v)
{
    return /*DomainLineSegment1D.l*/mData->parameters[2]*
        v + /*DomainLineSegment1D.a*/mData->parameters[3];
}

// pder(u, t) = u_t
// pder(u, x) = u_x
// pder(u_t, t) = c pder(u_x, x)

int functionPDE(struct MODEL_DATA *mData, int dScheme)
{
    // both states and algebraics have their specific
    array for space derivatives

```

```

// states u, u_t
// algebraics u_x

//we have u, u_t, pder(u,x), pder(u_t,x)

// u_x = pder(u,x)
// pder(u_x,x) = diff(u_x,x)
// pder(u,t) = u_t
// pder(u_t,t) = c pder(u_x,x)

//u      stateFields[M*0]
//u_t     stateFields[M*1]
//u_x     algebraicFields[M*0]

int M = mData->M;
int i;
for (i = 0; i < M; i++){
    /*u_x*/mData->algebraicFields[M*0 + i] = mData->
        stateFieldsDerSpace[M*0 + i];
}
differentiateX(/*u_x*/&(mData->algebraicFields[M*0]),
    /*pder(u_x,x)*/&(mData->algebraicFieldsDerSpace[M
*0]), mData, dScheme);
for (i = 0; i < M; i++){
    /*pder(u,t)*/mData->stateFieldsDerTime[M*0 + i] =
        /*u_t*/mData->stateFields[M*1 + i];
    /*pder(u_t,t)*/mData->stateFieldsDerTime[M*1 + i]
        = /*c*/mData->parameters[1] * /*pder(u_x,x)*/
        mData->algebraicFieldsDerSpace[M*0 + i];
}
return 0;
//in this approach some arrays for space derivatives
    might be unused (here pder(u_t,x))
}

//int functionPDE_2(struct MODEL_DATA *mData)
//{
//    // all space derivatives of states and algebraics
//    are stored as different algebraic fields
//    //-----
//    // TODO: pokracovat
//    //-----
//    // states u, u_t
//    // algebraics u_x

```

```

//
//      //we have u, u_t, pder(u,x), pder(u_t,x)
//
//      // u_x          = pder(u,x)
//      // pder(u_x,x)   = diff(u_x,x)
//      // pder(u,t)     = u_t
//      // pder(u_t,t)   = c pder(u_x,x)
//
//      //u      stateFields[M*0]
//      //u_t    stateFields[M*1]
//      //u_x    algebraicFields[M*0]
//
//
//      int M = mData->M;
//      int i;
//      for (i = 0; i < M; i++){
//          /*u_x*/mData->algebraicFields[M*0 + i] = mData
//          ->stateFieldsDerSpace[M*0 + i];
//      }
//      diffx(/*u_x*/mData->algebraicFields[M*0], /*der(u_x
//      ,x)*/mData->algebraicFieldsDerSpace[M*0]);
//      for (i = 0; i < M; i++){
//          /*pder(u,t)*/mData->stateFieldsDerTime[M*0 + i]
//          = /*u_t*/mData->stateFields[M*1 + i];
//          /*pder(u_t,t)*/mData->stateFieldsDerTime[M*1 +
//          i] = /*c*/mData->parameters[1]* /*pder(u_x,x)*/mData
//          ->algebraicFieldsDerSpace[M*0 + i];
//      }
//      return 0;
//      //this aproach is confusing as algebraics array is
//      used for various fields
//  }

int functionBC(struct MODEL_DATA *mData)
{
    int M = mData->M;
    mData->stateFields[mData->M*0 + 0] = 0;
    mData->stateFields[mData->M*0 + M-1] = 0;
    mData->stateFields[mData->M*1 + 0] = 0;
    mData->stateFields[mData->M*1 + M-1] = 0;

    return 0;
}

```

Listing 3.6: String equation – "generated" C code

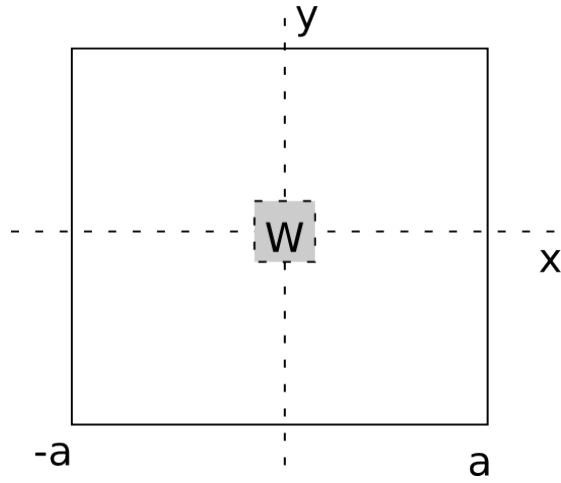


Figure 3.1: Heat eq.

3.2.3 Heat equation in square with sources (2D)

a .. domain square side hlaf length

c .. conductivity quocient

T .. temperature

$$W(x, y) = \begin{cases} 1 & \text{if } |x| < a/10 \text{ and } y < a/10 \\ 0 & \text{else} \end{cases}$$

equation

$$\frac{\partial T}{\partial t} - c \left(\frac{\partial^2 T}{\partial x^2} + \frac{\partial^2 T}{\partial y^2} \right) = W$$

initial conditions

$$T(x, y, 0) = 0$$

boundary conditions insulated walls (top, left, bottom)

$$\begin{aligned} \frac{\partial T}{\partial \bar{n}}(x, a, t) &= 0 \\ \frac{\partial T}{\partial \bar{n}}(-a, y, t) &= 0 \\ \frac{\partial T}{\partial \bar{n}}(x, -a, t) &= 0 \end{aligned}$$

fixed temperature (right)

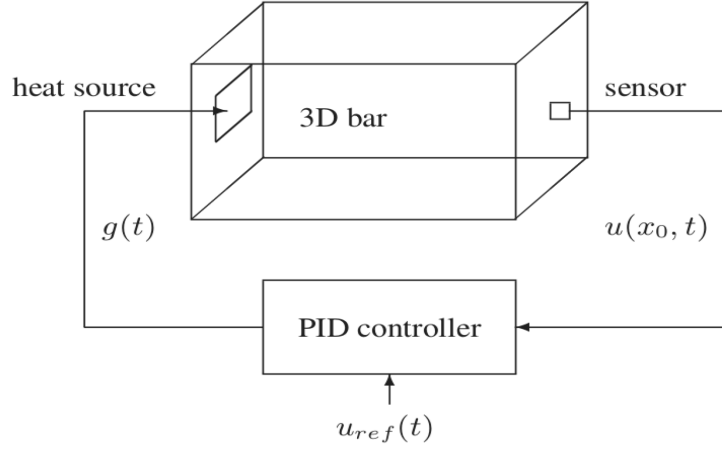


Figure 3.2: Heat transfer with source and PID controller

$$T(a, y, t) = 0$$

3.2.4 3D heat transfer with source and PID controller [10]

new problems:

- system of ODE and PDE
- `in` operator used to access field value in a concrete point (PID controller equation defining T_s).
- vector field
- differential operators `grad` and `diverg`

l_x, l_y, l_z .. room dimensions (6m, 4m, 3.2m)

T .. temperature (scalar field)

W .. thermal flux (vector field)

c .. specific heat capacity ($1012 J \cdot kg^{-1} \cdot K^{-1}$)

ρ .. density of air ($1.2041 kg \cdot m^{-3}$)

λ .. thermal conductivity ($0.0257 W \cdot m^{-1} K$)

T_{out} .. outside temperature ($0^\circ C$)

κ .. right wall heat transfer coefficient ($0.2 W \cdot m^{-2} \cdot K^{-1}$)

T_s .. temperature of the sensor placed in middle of the right wall

P .. power of heating

k_p, k_i, k_d .. coefficients of the PID controller (100, 200, 100)

T_d .. desired temperature ($20^\circ C$)

e .. difference between temperature of the sensor and desired temperature

heat equation

$$\begin{aligned}\frac{1}{c\rho}\nabla \cdot W &= -\frac{\partial T}{\partial t} \\ W &= -\lambda\nabla T\end{aligned}$$

boundary conditions left wall ($x = 0$) - heat flux given by heating power

$$W_x = \frac{P}{l_y l_z}$$

rear ($y = 0$) and front ($y = l_y$), resp. bottom ($y = 0$) and top ($z = l_z$) insulated walls

$$W_y = 0, \text{ resp. } W = 0$$

right wall ($x = l_x$) - not fully insulated

$$W_x = \kappa(T - T_{out})$$

PID controler

$$\begin{aligned}T_s &= T(l_x, \frac{l_y}{2}, \frac{l_z}{2}) \\ e &= T_d - T_s \\ P &= k_p e + k_i \int_0^t e(\tau) d\tau + k_d \frac{d}{dt} e\end{aligned}$$

Modelica code:

```
model heatPID
  record Room
    extends DomainBlock3D;
    Region0D sensorPosition(shape = shapeFunc, range =
      {{1,1},{0.5,0.5},{0.5,0.5}});
  end Room

  parameter Real lx = 1, ly = 1, lz = 1;
  Room room(Lx=lx, Ly=ly, Lz=lz);
  field Real T(domain = room, start = Tout);
  field Real[3] W(domain = room, start = {0,0,0});
  parameter Real c = 1012;
  parameter Real rho = 1.204;
  parameter Real lambda = 0.0257;
  parameter Real Tout = 0;
  parameter Real kappa = 0.2;
  Real Ts;
```

```

Real P;
parameter Real kp = 100, ki = 200, kd = 100;
parameter Real Td = 20;
Real eInt;
equation
  1/(c*rho)*diverg(W) = - pder(T,t)      in room.interior;
  W = -lambda*grad(T)                    in room.interior;
//TODO: use normal vector:
  W[1] = P/(lx*ly)                       in room.left;
  W[2] = 0                                in room.front +
    room.rear;
  W[3] = 0                                in room.top + room
    .bottom;
  W[1] = kappa*(T - Tout)                 in room.right;
  Ts = T                                  in room.
    sensorPosition;
  e = Td - Ts;
  der(eInt) = e;
  P = kp*e + ki*eInt + kd*der(e);
end heatPid;

```

Listing 3.7: heat equation with PID controller

3.3 More complex realistic models

3.3.1 Henleho klička - protiproudová výměna

$c_{in}(x, t)$.. koncentrace Na v sestupné části tubulu
 $\bar{c}_{in}(x, t)$.. koncentrace Na ve vzestupné části tubulu
 $c_{out}(x, t)$.. koncentrace Na v dřeni
 $Q(x, t)$.. tok vody v sestupné části tubulu
 $f_{H_2O}(x, t)$.. tok vody na milimetr délky z sestupné části tubulu do dřene
 f_{Na}^* .. tok sodíku ze vzestupné části tubulu do dřene na milimetr délky –
 aktivní transport – parametr
 L .. délka tubulu
 P_{H_2O} .. prostupnost cévy pro vodu (permeabilita)

$$\begin{aligned}
\frac{\partial Q}{\partial x}(x, t) + f_{H_2O}(x, t) &= 0 \\
(c_{out}(x, t) - c_{in}(x, t)) \cdot P_{H_2O} &= f_{H_2O}(x, t) \\
f_{H_2O}(x, t) &= \frac{dV}{dt}(t) \\
Q(L, t) \cdot c_{in}(L, t) &= f_{Na}^* \cdot L + Q(L, t) \cdot c^*(t) \\
\frac{\partial}{\partial x} (\bar{c}_{in}(x, t) \cdot Q(x, t)) &= f_{Na}^* \\
f_{Na}^* \cdot L &= \frac{dm_{Na}}{dt}
\end{aligned}$$

3.3.2 Oxygen diffusion in tissue around vessel

polar coordinates (r, φ)

$$\begin{aligned}
\frac{\partial \varrho}{\partial t} + q \left(\frac{1}{r} \frac{\partial \varrho}{\partial r} + \frac{\partial^2 \varrho}{\partial r^2} + \frac{1}{r^2} \frac{\partial^2 \varrho}{\partial \varphi^2} \right) + w &= 0 \\
\varrho(r_0, \varphi) &= \varrho_0 \\
\varrho(r, 0) &= \varrho(r, 2\pi) \\
\varrho_{nnn}(R, \varphi) &= 0 \quad (= \varrho_{tn}(R, \varphi))
\end{aligned}$$

ϱ .. oxygen concentration

ϱ_0 .. concentration in the vessel

q .. diffusion coefficient

w .. local oxygen consumption

R .. Ω diameter

The last equation should simulate infinite continuation of the domain.

3.3.3 Heat diffusion

domain boundary

$$\partial\Omega = (a_b \cos(v), b_b \sin(v)), \quad v \in \langle 0, 2\pi \rangle$$

equation [16]

$$\frac{\partial T}{\partial t} + \frac{\lambda}{c\varrho} \left(\frac{\partial^2 T}{\partial x^2} + \frac{\partial^2 T}{\partial y^2} \right) = W$$

λ .. thermal conductivity

$W(x, y)$.. heat power density of tissue (input)

$$W(x, y) = \begin{cases} W_0 & \text{if } x^2 + y^2 \leq r^2 \\ 0 & \text{else} \end{cases}$$

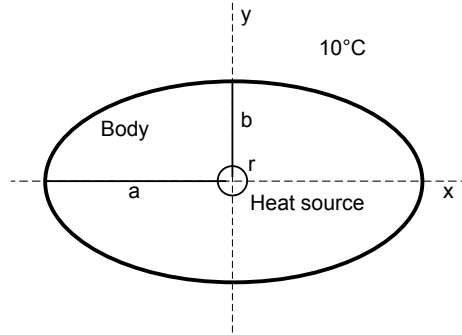


Figure 3.3: Scheme of heat diffusion in body

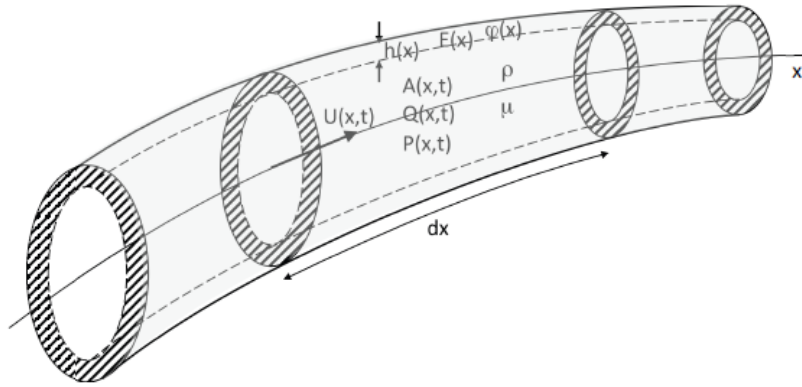


Figure 3.4: Arteria scheme

boundary condition

$$\lambda \frac{\partial T}{\partial n} = -\alpha(T - T_{out}), (x, y) \in \partial\Omega$$

α .. tissue-air thermal transfer coefficient [17]

initial condition

$$T(x, y, 0) = T_0(x, y)$$

3.3.4 Pulse waves in arteries caused by heart beats [2, 9, 11]

$A(x, t)$.. crosssection of vessel

$U(x, t)$.. average velocity of blood

$Q(x, t)$.. flux

$$Q = AU$$

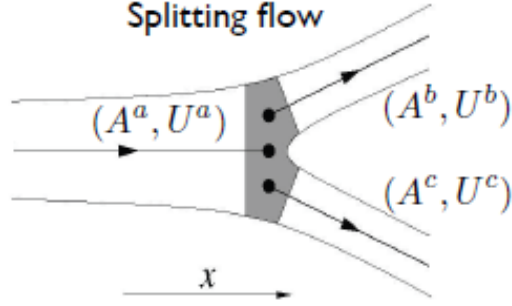


Figure 3.5: Arteria splitting

- $P(x, t)$.. pressure
 P_{ext} .. external pressure
 A_0 .. vessel crosssection at $(P = P_{ext})$ (24mm)
 $\beta = \frac{\sqrt{\pi} h_0 E}{(1-\nu^2) A_0}$
 h_0 .. vessel wall thicknes (2mm)
 E .. Young's modulus (0.24 - 6.55MPa)[5, 4, 3]
 ν .. Poisson ratin (1/2)
 $\varrho = 1050 \text{ kg m}^{-3}$.. blood density
 $\mu = 4.0 \text{ mPa s}$
 α .. other ugly coefficient, let us say its 1
 f .. frictional forces per unit length, let us assume inviscide flow $f = 0$, or
 $f = -AQ8\mu/(\pi r^4) = -8\pi\mu Q/A$ [15]
 μ .. dynamic viscosity of blood $(3.4) \cdot 10^{-3} \text{ Pa}\cdot\text{s}$ [14]

$$\begin{aligned}
 \frac{\partial A}{\partial t} + \frac{\partial Q}{\partial x} &= 0 \\
 \frac{\partial Q}{\partial t} + \frac{\partial}{\partial x} \left(\alpha \frac{Q^2}{A} \right) + \frac{A}{\varrho} \frac{\partial P}{\partial x} &= \\
 = \frac{\partial Q}{\partial t} + \alpha \left(2 \frac{Q}{A} \frac{\partial Q}{\partial x} - \frac{Q^2}{A^2} \frac{\partial A}{\partial x} \right) + \frac{A}{\varrho} \frac{\partial P}{\partial x} &= \\
 \frac{\partial Q}{\partial t} + 2\alpha \frac{Q}{A} \frac{\partial Q}{\partial x} + \left(\frac{\beta}{2\varrho} \sqrt{A} - \alpha \frac{Q^2}{A^2} \right) \frac{\partial A}{\partial x} &= \frac{f}{\varrho} \\
 P_{ext} + \beta \left(\sqrt{A} - \sqrt{A_0} \right) &= P
 \end{aligned}$$

Three segment geometry – splitting arteria

We model arteria being splited into two minor arteries. Three same equation systems (super-indexes A, B, C) are solved on three different domains. Systems are connected via BC.

Boundary conditions

input

$$\begin{cases} P^A(0, t) = P_S & t \in \langle 0, \frac{1}{3}T_c \rangle \\ Q^A(0, t) = 0 & t \in \langle \frac{1}{3}T_c, T_c \rangle \end{cases}$$

T_c .. cardiac cycle period

junction

$$\begin{aligned} Q^A(L^A, t) &= Q^B(0, t) + Q^C(0, t) \\ P^A(L^A, t) &= P^B(0, t) \\ P^A(L^A, t) &= P^C(0, t) \end{aligned}$$

terminal we simulate the continuation of segments B and C with just a resistance

$$Q(L, t) = \frac{P(L, t)}{R_{out}}, \text{ for } B \text{ and } C$$

For check: the result should be in agreement with Moens–Korteweg equation.

3.3.5 Vibrating membrane (drum) in air

Membrane [18]:

$\Omega_m = \{(x, y) | x^2 + y^2 < r^2\}$
 $u(x, y, t)$.. membrane displacement, $u : \Omega_m \times \langle 0, T \rangle \rightarrow \mathbb{R}$
 r .. membrane radius
 c_m .. membrane wave speed

$$\frac{\partial^2 u}{\partial t^2} = c_m^2 \nabla^2 u$$

Initial and boundary conditions

$$\begin{aligned} u(x, y, 0) &= u_0(x, y) \\ u(x, y, t) &= 0 \quad (x, y) \in \partial\Omega_m \end{aligned}$$

Air[12]:

$\Omega_a = \langle 0, l_x \rangle \times \langle 0, l_y \rangle \times \langle 0, l_z \rangle$
 $\mathbf{v}(x, y, z, t)$.. air speed, $\mathbf{v} : \Omega_a \times \langle 0, T \rangle \rightarrow \mathbb{R}^3$
 $p(x, y, z, t)$.. air pressure, $p : \Omega_a \times \langle 0, T \rangle \rightarrow \mathbb{R}$
 ρ_0 .. density
 c_a .. speed of sound

$$\begin{aligned}\rho_0 \frac{\partial \mathbf{v}}{\partial t} + \nabla p &= 0 \\ \frac{\partial p}{\partial t} + \rho_0 c_0^2 \nabla \cdot \mathbf{v} &= 0\end{aligned}$$

Initial and boundary conditions

$$\begin{aligned}\mathbf{v}(x, y, z, 0) &= \mathbf{0} \\ p(x, y, z, 0) &= p_0 \\ \mathbf{v}(x, y, z, t) \cdot \mathbf{n}(\partial\Omega_a) &= 0 \quad (x, y, z) \in \partial\Omega\end{aligned}$$

Position of membrane centre in room $\mathbf{a} = (a_x, a_y, a_z)$.. position vector of membrane centre in room

membrane lies in $\tilde{\Omega}_m = \{(a_x + x, a_y + y, a_z) | x^2 + y^2 < r^2\}$ in term of coordinates defined in Ω_a .

Equation connecting membrane and air

$$\mathbf{v}(x, y, z, t) \cdot \mathbf{n}_{\tilde{\Omega}_m} = \frac{\partial u}{\partial t}(x - a_x, y - a_y, t) \text{ in } \tilde{\Omega}_m$$

Appendix A

Articles and books

I want to read:

A DIFFERENTIATION INDEX FOR PARTIAL DIFFERENTIAL-ALGEBRAIC
EQUATIONS [7]

INDEX AND CHARACTERISTIC ANALYSIS OF LINEAR PDAE SYS-
TEMS [8]

Finite difference methods for ordinary and partial differential equations [6]

Appendix B

Questions & problems:

important topics are written in bold

B.1 Modelica language extension

- **Allow writing equations independent on particular domain and also coordinate system?**
 - perhaps yes
 - * using coordinate free differential operators (**grad**, **div** etc.)
 - * or using **dom** (as **this** in oop, **described later**) to address domain generally in differentiation
 - * or using **replaceable** and **redeclare** on domain class
 - but how to define fields in a domain independent way?
 - how to apply domain independently written equations on particular domain?
 - Write also boundary conditions domain independent?
 - Notation for normal vector to domain boundaries e.g. **omega.left.n**. How to write normal vector domain-independently?
- **How to deal with (name) coordinate (independent) variables**, so that it doesn't meddle with other variables (ODE)? Should it be fixed or defined within the domain definition? (Or cartesian fixed and others defined extra by user if needed?) Some approaches (possibly good ideas underlined):
 - in domain_name.region_name opens scope. Use keyword domain (or just dom), similarly as this in oop to address coordinate variables defined within this domain. Than coordinate names may be fixed.
E.g. pder(u, dom.x)=0 in omega.left

- can be used in combination with fixed or user-defined coordinate names
- name by user (use some special data type (Coordinate or Independent) to define independent variables within domain record (or class) – see 3.1
 - * in this case shall we have nevertheless some coordinates defined by default (perhaps cartesian)?
- NO. avoid coordinate variables at all
 - * allow writing equations coordinate-free, using only `pder(u,time)`, `grad`, `div`, ... operators (does it mean, we need no coordinates defined in domain?).
 - * use operators `pderx(u)`, `pdert(u)` or
- NO. `domain_name.variable_name` (e.g. `omega.x`). – This makes equation domain dependent.
- NO. Fixed names `x`, `y`, `z` used stand-alone. If they meddle with other variable, infer which one is it from the fact that we differentiate with respect to this variable and from the actual domain (indicated with `in op.`). – Makes model confusing.
- NO. fixed names and approach ODE variables from PDE in some special way.
- NO. use longer name for coordinate variables (e.g. `spaceX` ...)
- **Allow other independent variables than space variables? It would require not to define `x`, `y`, `z` coordinates (independent variables) by default but define independent variables always in the domain definition as discussed above.**
- How to map shape function return values on particular space variables (e.g. `x`, `y`, `z`) when they are not ordered? And what if there are more coordinate systems defined (e.g. cartesian and polar)? The same problem holds for functions that set start values to fields.
- How to define general differential operators (as `grad`, `div` ...) , if we use user defined coordinates?
- How to call attribute of Coordinate variable saying the type of the coordinate (now called `name`) should be the value assigned to this attribute written in quotes? It is also related with the previous question.
e.g. something like `Coordinate x (name = "cartesian");`
- **How to write boundary conditions that combine field variables from different domains?** Allow some connection of regions of different domains.
- Initialization.

- Rename *region* to *manifold*[1]?
- unify somehow concept of region and domain?
- How to call divergence operator (standard `div` is already used for integer division)
- How should the shape, geometrical structure, mesh structure, etc. be described by an external file?
- Philosophical problem: What exists first, domain or coordinates? I would say coordinates must exist first as domain shape is defined using shape-function using some coordinates.
- Is allowed passing functions as variable? It is used in setting start value to fields.

Solved problems:

- Rename ranges to intervals?
 - yes
- Domain description where some parameters are in range and others are fixed: $\{\{1,1\}, \{0.5,0.5\}\}$ or $\{\{1,1\}, 0.5\}$?
 - allow both
- How to deal with vector fields? How to access its elements – using an index or scalar product with standard base vectors?
 - both
- How to distinguish the main domain (now called `DomainLineSegment1D`, `DomainRectangle2D` ...) and its “subsets” where some equations hold (now called `Domain0D`, `Domain1D` ...). I think only one of them should be called domain.
 - “subsets” remained to regions – (`Region0D`, `Region1D`, `Region 2D`, `Region3D`)
- directional derivative
 - `der(u,v)` (u is scalar or vector field in \mathbb{R}^n , v is vector in \mathbb{R}^n)
- Should it be possible to override initial and boundary conditions given in model with some different values from external configuration file?

- yes
- How to set initial condition for field derivative in similar way as using `start` attribute (i.e. not using equation in `initial` section)? See 3.2.2
 - `start` attribute is array where index denotes the derivative `start[0]` - actual value, `start[1]` - first derivative

B.2 Generated code

- How to represent on which particular boundary an boundary condition hold in generated code (or even on which interior domain hold which PDE equation system, if we support various interiors)? – Some domain struct could hold both shapeFunction parameter ranges and pointer (or some index) to function with the corresponding equations. Or boundary condition function knows on which elements (indexes) of variable arrays should be applied.
- **Should be generated functions independent on grid? It means either**

```
functionPDEIndependent(u,u_x,t,x)
u_t = ...
return u_t
or
functionPDEDependent(data)
for (int i ...)
u_t[i] = ...
```

B.3 Numerics and solver

- Shall we support higher derivatives in solver?
- What about space derivatives? – All state and algebraics have corresponding array for its space derivative, not all of them are used. – Or all space derivatives of states and algebraics are stored as different algebraic fields. – Or there is array for space derivatives that is utilised by both states and algebraics that need it.
- What about multi step methods (RK, P-K)?
- How to generate even (or arbitrary) meshes with nonlinear shape functions?
- How to generate mesh points just on the boundary? 1D – simple – just two points. 2D – We can go through the boundary curve and detect crossings of grid lines. 3D – who knows?!

- How to plugin an already existing solver?
- How to determine causality of boundary conditions and other equations that hold on less dimensional manifolds.
- Build whole solver in some PDE framework, perhaps Overture (<http://www.overtureframework.org/>)

B.4 TODO

- Write a library for vector fields defining scalar and vector product, divergence, gradient, rotation...
- Write model in coordinates different from cartesian

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