Object-oriented Language Variables 
and New Continuous Media Simulation Approach

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Abstract: - Numerical computer algorithms to model physical processes in continuous media are generally based on one of the schemes of discretization of PDEs, which are considered mathematical models of phenomena. PDEs, on their own, are relations between values of given functions, unknown functions and their derivatives calculated through a limit process for small media elements interactions when elements’ sizes and evolution time steps both approach zero. The limits can typically only be calculated using (many) simplifying assumptions not based on the principles of physics. It follows that traditional numerical schemes simulate oversimplified mathematical models and not real processes. In the Direct Computer Modeling approach, an algorithm exactly reproduces interactions between small volume elements. That paradigm can be mapped onto the object-oriented computer language structure (or class) type variables.

Keywords: - Computational Physics, Continuous Media Simulation.

1 Introduction
A recently published article [1] state “… the new discipline [computational simulation] …is still troublingly immature”. One of the reasons for that may be the legacy of mathematical approaches. For roughly the last three centuries, partial differential equations (PDE) have comprised the basis for model phenomena in continuous media. Even the newly introduced computers are used simply as numerical solvers of PDEs. That means that PDEs are still considered basic models of real physical phenomena in continuous media and computers play the role of a powerful resource to numerically solve equations.

Fig.1 shows the logic of the traditional approach. What may look strange is the necessity to model, through a finite numerical scheme, a symbolic model, PDE, as opposed to the physical process itself. PDEs are derived through a discretization procedure of real physical media, with a subsequent mathematical limit process accompanied generally by many assumptions not based on the principles of physics. Any PDE is just an expression symbolically encoding some unnecessary oversimplifications of a real physical process. A computer, being a finite machine, can run a discrete time/spatial scheme evaluating functions satisfying PDE, so would it not be more practical to run discrete algorithms of interactions/states evolution of small media elements? In other words, it is more consistent to exclude blocks 3 and 4 from the traditional scheme and remove the statement “oversimplified assumptions necessary to derive PDE from” from the block 2.

In terms of the described approach, it is difficult to explain the sense of nonlinearity. When somebody says “nonlinear process”, it really means “nonlinear equations” corresponding to the process.

The approach is applicable not just to modeling continuous media but to arbitrary networks (for example, electronic schemes). The first model of soliton like behavior, was
a nonlinear elastic string simulation. Wide spectrum of potential applications expands as far as simulating relativistic charges beams.

An option existed to use a huge array of microprocessors, corresponding to elements in a physical medium [2]. Each microprocessor would save all numerical parameters defining the state of an element, implementation of all functions necessary to transform the element state parameters, and sufficient number of ports to communicate with processors within the network. In that case, the array should behave exactly like a real physical medium. To some extent, it would resemble an analog computer but have much greater capabilities. Something similar but much more simple was implemented in the “cellular automata” approach [3].
It was not obvious then which operating system could work with such multiprocessor systems, though some good projects existed. Fortunately, common computer languages such as C++ (and object-oriented PASCAL) had special types of variables that could support interacting media elements on a software (program) level.

Let me remind the reader the definition of the structure type variable in the C++ language.

In plain “C” language, the struct type variable can be thought of as a list of “standard” type variables (integers, doubles, strings, etc.), for example:

```c
struct sample {
    int i;
    double d;
    char c[30];
} st_one, st_two;
```

In C++, structure fields may be functions:

```c
struct cl {
    int i;
    int get_i(void);
    void put_i(int j);
};
```

Functions declared in the structure must have implementations:

```c
int cl::get_i(void)
{
    return i;
}

void cl::put_i(int j)
{
    i = j;
}
```

The class type variables are more flexible in the sense that you can keep class members, for example, private, accessible just to functions from the class. In struct all members are by default public. Any function from a program can get access to all of them.

It is evident that such types of variables ideally match the requirements for modeling small media elements interactions. Each single media element can be uniquely identified, at a particular instant of time, by some number of its state parameters that are “regular” type variables of the class. The class member functions derived from laws of physics then define the element behavior (its evolution in time).

All that exactly resembles what is going on in real physical media when we want to simulate it through partitioning and small element interactions. In that sense, we have analog simulation scheme implemented on software level.

3 Elements Behaviors and C++ Classes

It is more convenient to use C++ class type variables to represent elements’ states and transformations, mainly because in C++ (or any other object-oriented language) it is definitely stressed that a class is an object type variable comprising of both elements’ state variables and their transformation functions. Inheritance also makes the class approach more flexible.

Base abstract class representing a medium element can be defined as:

```c
class MediumElement {
    double current_t, delta_t;
    double* state_parameters;
    int nmb_parameters = 0;

public:
    virtual void transform_parameters(void (func*)(double* param, double* m_param, MediumElement** neighbors))=0;

    // some interface functions
    // etc.
};
```
The pointer to doubles, state_parameters, is a dynamically allocated array of numerical parameters, uniquely identifying (physical) element state at a particular instant in time.

Virtual (specifically implemented for each considered physical process) function transform_parameters should model arbitrary element behavior, depending on its current state (pointer param), global media parameters (pointer m_param) and an array of pointers to other elements interacting with the current element.

A derived class corresponding to some special type of a medium and specific model then can be defined as:

```cpp
class SpecMediumElement: public MediumElement {
    double* model_parameters;
    void Model(double* param, double* m_param, SpecMediumElement** neighbors);

    // and necessary interface functions
}
```

The function Model( ...) is used to input variable for the transform_parameters. Actual implementation of it should be written in accordance with physical laws applied to a particular model. Unquestionably, additional interface functions to initialize elements’ state parameters, model parameters and external inputs should be defined.

The whole medium volume of interest may be represented as an array, SpecMediumElement*, (with one index per element as assumed in previous sections) or as SpecMediumElement***, with three indexes for each single element, which may be more convenient.

### 4 String Elements and C++ Class Type Variables

Let me demonstrate how all that looks for the well-known elastic string physical and mathematical model.

Consider an element of a string. Let’s create a class, describing the element in the same way as in deriving the string wave equation. The element state is fully defined by:
- moment in time, \( t \)
- element’s position on x-axis, \( x \)
- element’s length projection on x-axis, \( \Delta x \)
- element’s left and right ends’ tangent directions, \( \alpha_1 \) and \( \alpha_2 \), or their trigonometric functions
- element’s current vertical displacement from x-axis, \( u \)
- element’s velocity in the direction perpendicular to x-axis, \( v \)
- two global parameters, stretch tension force value, \( T \) and linear density, \( \rho \)

We have to consider string element’s state evolution in time. Let’s take the simplest rule of time parameter transformation:
\[
t 
\Delta t 
\]
\( \Delta t \) is constant.

Parameters \( x \) and \( \Delta x \), for a single element, do not change values.

For now, we can express \( u \) and \( v \) as:
\[ v(t + \Delta t, x) = V(t, x, \Delta x, v(t), u(t), \alpha_1, \alpha_2) \]
\[ u(t + \Delta t, x) = U(t, x, \Delta x, v(t), u(t), \alpha_1, \alpha_2), \]

where \( V \) and \( U \) are some (integration) functions (schemes).

An interesting issue is how we can define parameters \( \alpha_1 \) and \( \alpha_2 \). I do not want to consider unnecessary complicated schemes involving predefined \( \alpha_1 \), and have to average element’s curvature, etc. I will consider a much more relevant scheme with explicitly involved elements’ interactions (that do not take place in a traditional scheme). In that scheme, the element representing class should contain two pointers to (left and right) neighbors:

```cpp
class _element _{
    ....
    element* _left;
    element* _right;
    ....
}
```

Then, we have:

\[
\tan \alpha_1 = \frac{u - (\text{left} \to u)}{\Delta x}
\]

\[
\tan \alpha_2 = \frac{u - (\text{right} \to u)}{\Delta x}
\]

At this point, we already have:

```cpp
    class _element _{
        ....
        double_t;
        double_\Delta t;
        double_x;
        double_\Delta x;
        double_\alpha _left;
        double_\alpha _right;
        double_u;
        double_v;
        element* _left;
        element* _right;
        public :
            double_U(t, \Delta t, x, \Delta x, u*, v*, \alpha _left, \alpha _right);
            double_V(t, \Delta t, x, \Delta x, u*, v*, \alpha _left, \alpha _right);
            double_get_alpha(element* _elmnt);
        }
    }
```

Implementation of the last function is already known:

```cpp
    get_alpha(element* _elmnt)
    {
        return \frac{\tan^{-1}(u - (\text{elmnt} \to u))}{\Delta x};
    },
```

that gives both \( \alpha _\text{left} \) and \( \alpha _\text{right} \), depending on the \( \text{elmnt} \) argument.

5 Conclusion

I have tried to briefly describe an approach to computer modeling of physical phenomena in continuous media. The Direct Computer Modeling approach is an attempt to use computers and software more adequately, instead of using an outdated method of symbolically representing real processes flows. This approach (or something similar) may become a good alternative to numerical schemes, based on hieroglyphic PDE descriptions.
The approach was initially outlined [4] and described putting more stress on object-oriented implementation [5] in earlier works. A general scheme for continuous media dynamics has also been addressed [6]. As I mentioned before, the first computer program modeling an elastic string dynamics with Direct Computer Modeling demonstrated soliton type string excitation behaviors and showed that split stresses, not the stretch ones, actually defined the whole picture. I mention this example to emphasize that with Direct Computer Modeling we can make further discoveries even in well-known situations [7].

References:


