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Title of your project

In partial fulfillment of MSc in High-performance Computing
in the School of Mathematics

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Supervision:

Your primary supervisor

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ACKNOWLEDGEMENTS

bsadassaaWrite acknowledgements to your supervisor, classmates, friends, family, partner. . . anyone who supported you during the MSc.

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ABSTRACT

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2 RELATED WORK

To gain well quality solution of various types of PDEs is prohibitive and notoriously challenging. The number of methods available to determine canonical PDEs is limited as well, includes separation of variables, superposition, product solution methods, Fourier transforms, Laplace transforms and perturbation methods, among a few others. Even though these methods are exclusively well-performed on constrained conditions, such as regular shaped geometry domain, constant coefficients, well-symmetric conditions and many others. These limits strongly constrained the range of applicability of numerical techniques for solving PDEs, rendering them nearly irrelevant for solving problems practically.

General, the methods of determining numerical solutions of PDEs can be broadly classified into two types: deterministic and stochastic. The mostly widely used stochastic method for solving PDEs is Monte Carlo Method [?] which is a popular method in solving PDEs in higher dimension space with notable complexity.

2.1 FINITE DIFFERENCE METHOD

The Finite Difference Method (FDM) is based on the numerical approximation method in calculus of finite differences. The motivation is quite straightforward which is approximating solutions by finding values satisfied PDEs on a set of prescribed interconnected points within the domain of it. Those points are which referred as nodes, and the set of nodes are so called as a grid of mesh. A notable way to approximate derivatives are using Taylor Series expansions. Taking 2 dimension Poisson Equation as instance, assuming the investigated value as, φ ,

$$\frac{\partial^2 \varphi}{\partial x^2} + \frac{\partial^2 \varphi}{\partial y^2} = f(x, y) \quad (2.1.1)$$

The total amount of nodes is denoted with $N = 15$, which gives the numerical equation which governing equation 2.1.1 shown in equation 2.1.2 and nodes layout as shown in the figure 1

$$\frac{\partial^2 \varphi_i}{\partial x_i^2} + \frac{\partial^2 \varphi_i}{\partial y_i^2} = f(x_i, y_i) = f_i, \quad i = 1, 2, \dots, 15 \quad (2.1.2)$$

In this case, we only need to find the value of internal nodes which i is ranging from 1 to 10. Next is aiming to solve this linear system 2.1.2.

2.2 PHYSICS INFORMED NEURAL NETWORKS

With the explosive growth of available data and computing resources, recent advances in machine learning and data analytics have yielded good results across science discipline, including Convolutional Neural Networks (CNNs) [?] for image recognition, Generative Pre-trained Transformer (GPT) [?] for natural language processing and Physics Informed Neural Networks (PINNs) [?] for handling science problems with high complexity. PINNs is a type of machine learning model

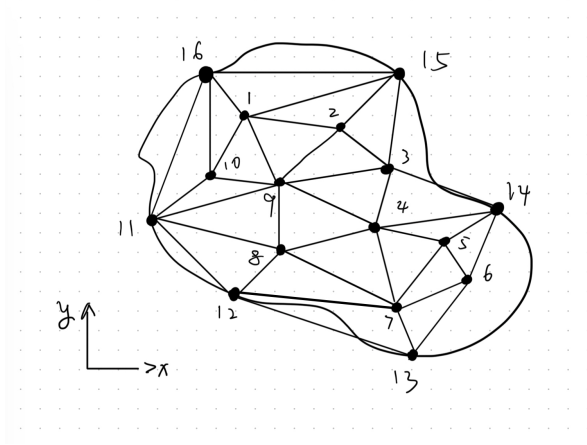


Figure 1: The Schematic Representa of of a 2D Computiationl Domain and Grid. The nodes are used for the FDM by solid circles. Nodes 11 – 15 denote boundary nodes, while nodes 1 – 10 denote internal nodes.

makes full use of the benefits from Auto-differentiation (AD) [?] which led to the emergence of a subject called Matrix Calculus [?]. Considering the parametrized and nonlinear PDEs of the general form [E.q. 2.2.1] of function $u(t, x)$

$$u_t + \mathcal{N}[u; \lambda] = 0 \quad (2.2.1)$$

The $\mathcal{N}[\cdot; \lambda]$ is a nonlinear operator which parametrized by λ . This setup includes common PDEs problems like heat equation, and black-stokz equation and others. In this case, we setup a neural network $NN[t, x; \theta]$ which has trainable weights θ and takes t and x as inputs, outputs with the predicting value $\hat{u}(t, x)$. In the training process, the next step is calculating the necessary derivatives of u with the respect to t and x . The value of loss function is a combination of the metrics of how well does these predictions fit the given conditions and fit the natural law [Fig. 2].

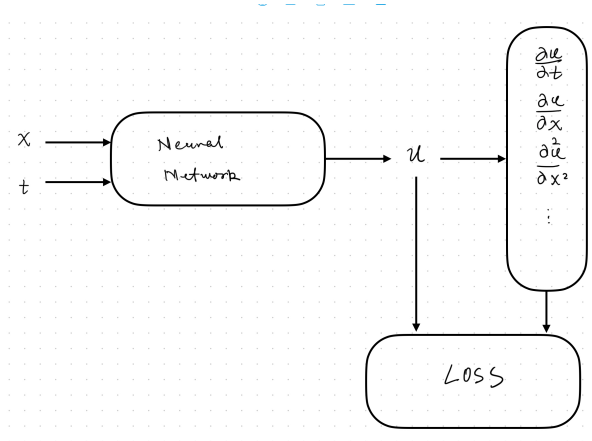


Figure 2: The Schematic Representa of a structure of PINN.

2.3 FINITE DIFFERENCE TIME DOMAIN METHOD

As described previously in the section 2.1, FDM could solve the PDEs in its original form where Finite Element and Finite Volume Methods gained results by solving modified form such as an integral form of the governing equation. Though the latter methods are commonly get better results or less computational hungry, the FDM has many descendants, for instance the Finite Difference Time Domain Method (FDTD) where it still finds prolific usage are computational heat equation and computational electromagnetics (Maxwell's equations [?]). Assuming the operator $\mathcal{N}[\cdot; \lambda]$ is set to ∇ where it makes E.q. 2.2.1 become to heat equation 2.3.1.

$$\frac{\partial u}{\partial t} - \lambda \left(\frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} \right) = 0 \quad (2.3.1)$$

Using the key idea of FDM, assuming the step size in spatio-time space are Δx , Δy and Δt , we could have a series equations which have form [E.q. 2.3.2].

$$\frac{u_{i,j}^{n+1} - u_{i,j}^n}{\Delta t} = \lambda \left(\frac{u_{i+1,j}^n - 2u_{i,j}^n + u_{i-1,j}^n}{\Delta x^2} + \frac{u_{i,j+1}^n - 2u_{i,j}^n + u_{i,j-1}^n}{\Delta y^2} \right) \quad (2.3.2)$$

when the time step size satisfies the Couran, Friedrichs, and Lewy condition (CFL[?]). We could get the strong results by iterating the equation 2.3.2, or more specifically using equation 2.3.3 to get the value u of next time stamp $n + 1$ on nodes i, j .

$$u_{i,j}^{n+1} = u_{i,j}^n + \frac{\lambda \Delta t}{\Delta x^2} (u_{i+1,j}^n - 2u_{i,j}^n + u_{i-1,j}^n) + \frac{\lambda \Delta t}{\Delta y^2} (u_{i,j+1}^n - 2u_{i,j}^n + u_{i,j-1}^n) \quad (2.3.3)$$

also shown in the figure 3.

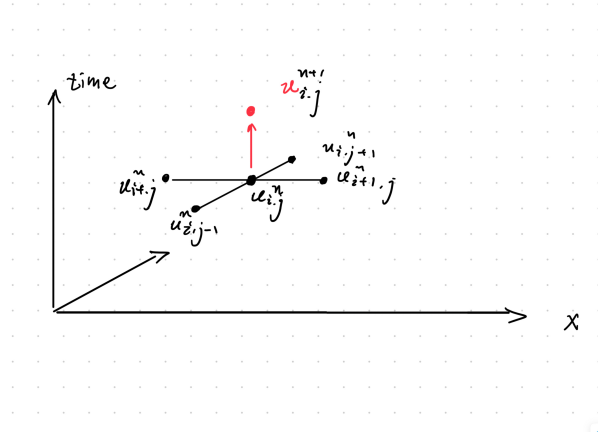


Figure 3: The Schematic Representa the computational spatio-time domain of FDTD methods.

3 PROBLEM SETUPS

Due to the inherent limitations of current computing systems, obtaining sufficiently precise solutions is both computationally expensive and time-consuming. These challenges arise from the constraints imposed by the clock speed of computing units, as described by Moore's Law [?], as well as the relatively low communication speeds between these units. While modern numerical methods have advanced to a level where they can produce satisfactory results within acceptable time frames across many research domains, the increasing scale of problems we aim to solve has driven the search for more cost-effective approaches. This has led to a growing interest in neural networks as a promising alternative.

In this project, I aim to evaluate the performance of the Finite-Difference Time-Domain (FDTD) method and the Physics-Informed Neural Network (PINN) model within parallelized computing environments by find the steady-state solution of PDEs. These two methodologies broadly represent the current approaches to handling PDEs, specifically CPU-based parallelization and GPU-based parallelization.

3.1 GENERAL FORM

Starting with the general form of the PDEs, rather than the specific euqations, is because different equations give perform differently on the same compute system. To this end, consider the previously discussed form of PDEs shown in equation 2.2.1 which parametrized by number λ and an operator $\mathcal{N}[\cdot; \lambda]$. Moreover, we assume the variable x is a 2D or 3D spatio vector which is written in $\vec{x} \in \mathbb{R}^d$, $d = 2, 3$.

$$\begin{aligned} \frac{\partial u}{\partial t}(t, \vec{x}) + \mathcal{N}[u(t, \vec{x}); \lambda] &= 0, & \vec{x} \in \Omega, t \in [0, +\infty) \\ u(0, \vec{x}) &= \varphi(\vec{x}), & \vec{x} \in \Omega \\ u(t, \vec{x}) &= g(t, \vec{x}), & \vec{x} \in \overline{\Omega}, t \in [0, +\infty) \end{aligned} \quad (3.1.1)$$

The domain of this PDE system is considered between 0 and 1, where is denoted with $\Omega = [0, 1]^d$, $d = 2, 3$. To these setups, we have the general form of the PDEs we are going to investigated, shown in equations 3.1.1 The boundary condition shown in E.q. 3.1.1 is Dirichlet Condition known as first type boundary condition, where as the second type boundary condition (Von Neuman) [E.q. 3.1.2] gives the other form of $u(t, \vec{x})$ at the boundary $\overline{\Omega}$.

$$\frac{\partial u}{\partial \vec{x}} = g(t, \vec{x}), \quad \vec{x} \in \overline{\Omega}, \quad t \in [0, +\infty) \quad (3.1.2)$$

3.2 SPECIFIC FORM

With general form proposed in section 3.1[E.q. 3.1.1], I specify a particular form of this heat problem to help us to have better understand the quality of our solutions and programs. In

2 dimension space, the domain $\Omega = [0, 1]^2 \in \mathbb{R}^2$ and its boundary denoted with $\overline{\Omega}$, the initial condition $\varphi(x, y) = 0$. such problem has the certain form below

$$\begin{aligned} \frac{\partial u}{\partial t} &= \alpha \left(\frac{\partial u^2}{\partial^2 x} + \frac{\partial u^2}{\partial^2 y} \right) & (x, y) \in \Omega, t \in [0, +\infty) \\ u(0, x, y) &= \varphi(x, y) = 0 & (x, y) \in \Omega \\ u(t, x, y) &= g(x, y) = \begin{cases} y, & x = 0, y \in (0, 1) \\ 1, & x = 1, y \in (0, 1) \\ x, & y = 0, x \in (0, 1) \\ 1, & y = 1, x \in (0, 1) \end{cases} & t \in [0, +\infty) \end{aligned} \quad (3.2.1)$$

With given format, and $\alpha = 1$, we have the analytical solution of this equations, where is

$$u(t, x, y) = x + y - xy, \quad (x, y) \in \Omega, t \in [0, +\infty) \quad (3.2.2)$$

In 3 dimension space, similarly, with identical initial condition set up to 0, coefficient $\alpha = 1$, the boundaries are

$$u(t, x, y, z) = g(x, y, z) = \begin{cases} y + z - 2yz, & x = 0, \\ 1 - y - z + 2yz, & x = 1, \\ x + z - 2xz, & y = 0, \\ 1 - x - z + 2xz, & y = 1, \\ x + y - 2xy, & z = 0, \\ 1 - x - y + 2xy, & z = 1 \end{cases}, t \in [0, +\infty) \quad (3.2.3)$$

In such case, the analytical solution has for form below

$$u(t, x, y, z) = x + y + z - 2xy - 2xz - 2yz + 4xyz, \quad (x, y, z) \in \Omega, t \in [0, +\infty) \quad (3.2.4)$$

3.3 DISCRETIZATION

To begin with discretizing the objects or regions we intend to evaluate via matrices, we consider a straightforward approach: using the coordinates in $d = 2, 3$ dimensional spaces and the function values at those points to simplify the objects. This naive approach works well for investigating objects with regular shapes, such as a cube.

For the FDTD (Finite-Difference Time-Domain) method, we use a finely generated d -cube with shape $\{n_i\}_i^d$. Including the boundary conditions, the cube has $\prod_i (n_i + 2)$ nodes. It requires $4 \prod_i (n_i + 2)$ bytes for float32 or $8 \prod_i (n_i + 2)$ bytes for float64 to store in memory. With this setup, for equally spaced nodes, we have:

$$\Delta x_i = \frac{1}{n_i - 1} \quad (3.3.1)$$

Unlike the previously generated regular grid of points with dimensions $n_x n_y n_z$, another strategy is to randomly generate the same number of points based on the same known conditions,

covering both the central part and the boundary. In this scenario, shown in figure, there are $n_x n_y n_z$ central points, with function values set according to the boundary conditions, and $2 \times (n_x n_y + n_y n_z + n_z n_x)$ boundary points to be solved. This set of points can be used for training a PINN model.

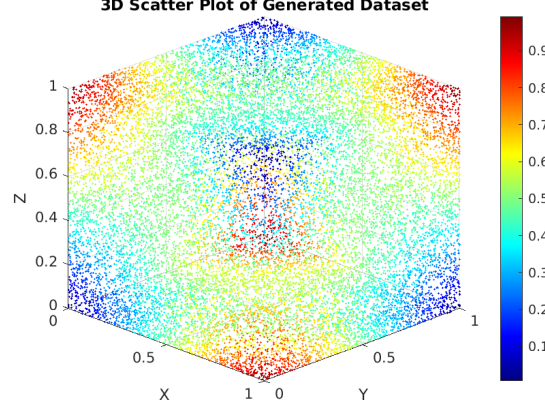


Figure 4: Randomly general dataset for 3D PINN model training.

3.4 ACCURACY

When we tried to represent numbers using arithmetic in binary, decimal or hexadecimal, truncation always affects the precision of every number, or so called as round-off-error.

ROUND-OFF ERROR

In IEEE-754 [?] standards, a 32-bit floating pointer number, single precision, obligatorily represented with 23-bit mantissa, 8-bit exponent and 1-bit for sign. Where as 64-bit floating number, double precision, also ubiquitous used, which has 11-bit exponent and 52-bit mantissa. After almost three decades development, not only single and double precisions (float32, float64) are ubiquitously in use, also more formats such as fp4, fp8, and fp16 etc. Both of them follows the simple form of exponent k , sign n and mantissa N . [?]

$$2^{k+1-N} n$$

Round-off errors are a manifestation of the fact that on a digital computer, which is unavoidable in numerical computations. In such case, the precision of the number depends on how many bytes are used to store single number. For instance, a float32 number provides $2^{-23} \approx 1.2 \times 10^{-7}$, and a double precision number gives $2^{-53} \approx 2.2 \times 10^{-16}$, such number is called machine ϵ which is the smallest number the machine can represent with given format.

In numerical methods I investigated, the FDTDs are conventionally using double precision number so that the programs can treat extremely large and small numbers simultaneously in

the same computation, without worrying about the round-off errors. However, as mentioned, fp32 and fp16 are also popular use in scientific computing, especially in machine learning training process. While the lastest training GPUs are integrated compute accelerate unit for low precision floating numbers. [?].

FLOATING-POINT ARITHMETIC

The other type loss comes from the arithmetic operations on two numbers x, y . The standard model holds that

$$fl(x \text{ op } y) = (x \text{ op } y)(1 + \delta), \quad |\delta| < \epsilon \quad (3.4.1)$$

where the op stands for the four elementary operations: $+, -, \times, /$. [?, ?, ?, ?].

3.5 COMPUTIATIONAL TOPOLOGY

The computational topology is critically important when we are programming parallel PDEs solver softwares. Put the strongly speed-dependent data into the slow memory could make entire program slower.

CALLAN The cluster we are using for this project is **Callan** [?] which has 2 CPUs per compute node, and each CPU has 32 cores with single thread. The Non-Uniform Memory Access (NUMA) nodes are layout as following Accessing the other NUMA node's memory reduces the

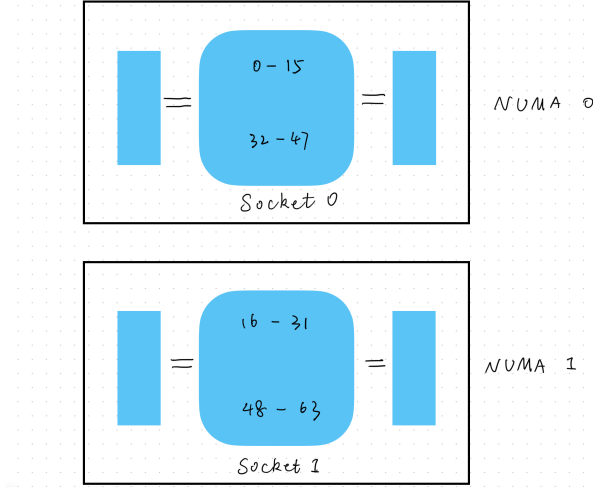


Figure 5: NUMA topology of single node on Callan

bandwidth and also the latency, though the bandwidth is commonly high enough, the latency can increase by 30% to 400% [?]. This latency becomes dangerous when writing shared memory parallel programs.

4 METHODOLOGY

4.1 GENERAL SETUPS

Initially, we need to determine the data types to be used and define macros for assertions and helper functions to ensure that the program can detect common bugs and report their locations. These features can also be disabled in the release version for performance optimization. Such details are defined in the `assert.hpp`, `helper.hpp`, and other related header files located in the subfolder.

4.2 MULTI-DIMENSION MATRIX DESIGN DETAIL

The FDTD method is a type of FDM. The main idea behind FDM is briefly outlined in Section 2.1. The challenge lies in implementing it in a computer system to ensure it runs both correctly and efficiently. Also, the data types I am using for storing sizes are `unit32_t` and `unit64_t` while I used them as `Dworld` and `Qworld` respectively, also are defined as `size_type` and `super_size_type` in `__detail` namespace.

PERFORMANCE BALANCING

To achieve this, it is crucial to develop an efficient Matrix object. Rather than using the standard library's (STL) vector module, which can be slower due to the overhead of row pointers, I opted to build the Matrix object using a unique pointer (`std::unique_ptr`), which includes only basic features such as reset, swap, and most importantly, a destructor that automatically deletes pointers. This approach enhances the safety of memory management in our programs. Additionally, from a safety perspective, given that I implemented many features within the matrix object, I followed standard library conventions for naming. This includes using the `__detail` namespace within namespace `multi_array` to hide objects and features that are not intended for direct use by the end user.

TEMPLATE OBJECT DESIGN OF MATRIX SHAPE

STRIDES Besides that, in the multidimensional cases, the size in each dimension is not enough for accessing variables, this is where we need the `strides` member variable, which stores the number of element the operator needs to skip in each dimension. The `__multi_array_shape` object is encapsulated within the `__detail` namespace and serves as a member variable of the later template object for the multi-dimensional matrix. This object includes a member variable defined using the STL vector, as the shape object primarily stores the sizes for each dimension, which typically requires only a small amount of space. Additionally, this object provides member functions to access the size of a given dimension.

Algorithm 1 Stride implementation

```
1: dims                                # STL vector, stores the matrix's size in each dimension.
2: strides                             # STL vector, has the same size with dims.
3: n = dims.size()                   # Store the dimension of matrix.
4: strides[d-1] = 1;                   # Stride is 1 in the first dimension.
5: for d = n - 1; d > 0; -d do
6:   strides[d-1] = strides[d] * dims[d]    # Determine stride in the latter dimension.
7: end for
8: return strides
```

PERFORMANCE BALANCING In certain scenarios, we only require the shape information of a matrix without needing to access the entire matrix object. Accessing the shape information through well-defined operators is a more efficient way to handle multidimensional matrices. This is particularly crucial in parallel programming, where understanding the shape of a matrix is of critical importance. Sometimes, a process may need to know the shape of matrices stored on other processes. In such cases, using this matrix object as a local variable within functions increases the likelihood that the compiler will store it in a register, which is generally faster than using heap or stack memory. In addition, it includes check and cast functions that allow the user to verify if the template data type `__T` is signed using `constexpr`. The `constexpr` keyword ensures that this check occurs at compile time, and if the data type is not legal, the program will assert and provide a message indicating that the indexing value must be a non-negative number.

TEMPLATE OF MULTI-DIMENSIONAL MATRIX IMPLEMENTATION

The Matrix in this project is designed to support various data types in C++. Consequently, the matrix is implemented as a template class with several essential features, template variable `__T` and `__NumD`, for the value data type and number of dimension, also including iterators, swap functionality, fill operations, and support for the IO stream operator `<<`. To facilitate this, the `__array_shape` object is used to explicitly manage and access the array's shape information.

OPERATOR () The hard part of this object designed is the support template number of dimension, whereas the dimension is integer not less than 1, the operator of access element is designed by following algorithm

OVERLOAD OPERATOR << In order to print the multi-dimension array with operator `<<`, I designed a recursive helper function to print the matrix on given dimension. Thus we could call the function on the first dimension, and it will recursively print all dimensions.

Algorithm 2 Operator (Ext ... exts) of template matrices object `__detail::__array`

```
1: __NumD, __T;                                # Template variables: dimension, data type.
2: FINAL_PROJECT_ASSERT_MSE                    # Number of Arguments must Match the dimension.
3: index = 0, i = 1                             # Initialize variables in advance.
4: indices[] = __shape.check_and_cast(ext) # The indexes number must none-negative
    number.
5: for i < __NumD; ++i do
6:   index += indices[i] * __shape.strides[i]
7: end for
8: FINAL_PROJECT_ASSERT_MSE                      # Boundary checking.
9: return __data[index]
```

Algorithm 3 Recursive Function to Print Multi-Dimensional Array

```
1: current_dim, offset;                        # Parameters: current dimension, offset.
2: Dims __Dims;                               # Template variable: number of dimensions.
3: if current_dim == __Dims - 1 then
4:   os << "|"                                # Start printing last dimension.
5:   for i from 0 to arr.__shape[current_dim] - 1 do
6:     os << std::fixed << std::setprecision(5) << std::setw(9) <<
        arr.__data[offset + i];                # Print array elements with formatting.
7:   end for
8:   os << " |\n";                            # End of current row in the last dimension.
9: else
10:  for i from 0 to arr.__shape[current_dim] - 1 do
11:    next_offset = offset;                    # Initialize next offset.
12:    for j from current_dim + 1 to __Dims - 1 do
13:      next_offset *= arr.__shape[j];        # Update next offset based on shape.
14:    end for
15:    next_offset += i * arr.__shape[current_dim + 1]; # Finalize next offset for
    recursion.
16:    self(self, arr, current_dim + 1, next_offset); # Recursive call to print next
    dimension.
17:  end for
18:  os << "\n";                               # Print a newline after each dimension.
19: end if
```

4.3 TEMPLATE MULTI-DIMENSION MATRIX DESIGN OF USER

With contiguity of safety, this object of multi-dimension array is accessible to users without direct visit to the memory space where store values of matrix.

RESOURCE ACQUISITION IS INITIALIZATION (RAII)

This private object has only a member variable, a unique pointer to the template `__array`, and other member function provide necessary features to operating on it. Smart pointers acquire resources in their constructor and automatically release them in their destructor, which is the essence of RAII. By releasing resources in the destructor, smart pointers help prevent resource leaks. When an exception occurs, smart pointers automatically release resources, preventing resource leaks, thus it enhanced the safety level of using resources, reduce the potential memory leak problems.

TEMPLATE MULTI-DIMENSION IO FOR WRITING TO/READING FROM FILE

Initially, the multi-dimension matrix has variables shape, and values which given dimension and size in each dimension. This template design end up with these variable can be stored in given data types also leads with lower portability. To avoid such problems and from other point of views, I chose to store the matrices in binary format, rather than other type files. There are couple benefits of doing so,

1. Compatibility and Portability: The format of binary files is relatively stable and can be easily used in different programming environments or applications. Unlike `.txt` files, `.mat` files those has less compatibility across different platforms.
2. I/O Performance: Binary files can perform block-level I/O operations directly without needing to parse text formats or convert data types. This usually makes reading and writing binary files much faster than `.txt` files, especially when dealing with large-scale multidimensional matrix data.
3. Support MPI IO: Binary files support the MPI IO, which provides a significant reduction in the cost of communication, when storing and reading the large scale matrices.

However, the IO does not play a critical role in effects performance of FDTD algorithms, if and only if we need to store or load the data during evolving the arrays.

PURE MESSAGE PASSING PARALLEL

HYBRID PARALLEL

4.4 PHYSICS INFORMED NEURAL NETWORKS

CUDA PARALLEL

HYBRID PARALLEL

5 IMPLEMENTATION

5.1 FINITE DIFFERENCE METHODS

PURE MESSAGE PASSING PARALLEL

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5.2 PHYSICS INFORMED NEURAL NETWORKS

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5.3