C++ Design Patterns for Managing Parameters in Scientific Computing

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Abstract

A scientific computing module usually has many parameters to control its behaviors. Instantiate a pair of get/set functions for each parameter clutters the interface, incurs code-bloating, overburdens maintenance, and invites subtle bugs. To solve these problems, two patterns are proposed for parameter management. The “enumeration index” pattern is a simple solution for parameters of a single type. When the parameters belong to multiple types, it is difficult for a statically-typed language to access/modify these parameters using a single pair of get/set functions. The “parameter manager” pattern achieves this by templates and multiple inheritance. It also handles additional semantics such as constancy, dependency, and default values in an easy-to-use manner: a client only has to specify the set of parameters and their associated properties. Both patterns are illustrated by a common task of solving the diffusion equation in the C++ language.

Key words: Design patterns, Enumeration-Indexing, Parameter-Manager

1 Motivation

Scientific programs crunch numbers, a huge amount of numbers. Sometimes the number of parameters that controls the behaviors of a scientific program might also be overwhelming. How to handle these parameters is the central topic of this paper.

A scientific problem can often be thought of as a set of smaller problems coupled together. The solution of each smaller problem can be captured either by a function or a class; the latter differs from the former in that it has the

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notion of internal states afforded by its data members, or parameters. In the object-oriented programming (OOP) paradigm, one often faces two kinds of by-and-large choices: (i) how to break up the big problem into smaller ones, and, (ii) how to organize the individual modules to solve the big problem. A design pattern [3] contains a particular set of these structural choices, aiming to encapsulate possible changes of the big problem elegantly and efficiently.

The most common change to a scientific computation problem is probably varying its input parameters, the number of which is usually large in most practical applications. As an example, consider the diffusion equation in a rectangular $D$-dimensional domain,

$$\frac{\partial \phi}{\partial t} = \nu \nabla^2 \phi,$$

where $\mathbf{x} \in \mathbb{R}^D$ is the location vector, $t$ the time variable, $\phi = \phi(\mathbf{x}, t)$ a continuous scalar function, and $\nu$ the dynamic viscosity. (1) is often solved by integrating the system of ordinary differential equations, obtained by replacing the Laplacian term $\nabla^2 \phi$ with a spatial discretization $L\Phi$, such as the widely-used central differencing $\frac{1}{h^2}(\phi_{i+1} + \phi_{i-1} - 2\phi_{i})$. Within each time step, the following linear system is solved repeatedly as an approximation to the solution of (1):

$$(\alpha - \beta L)\Phi = \rho,$$

where the scalars $\alpha, \beta$ and the right-hand-side vector $\rho$ usually depends on the spatial discretization and the time-integrator. See [5] and [6] for two examples using implicit Runge-Kutta methods. A program implementing this algorithm with an iterative linear solver has many input parameters, some of which are listed in Table 1.

One straightforward way to handle these parameters are the traditional get/set approach, which declares these parameters as private or protected data members of class DiffusionSolver and access/modify them using public member functions such as `getTimeOrder()`, `setTimeOrder()`, `getSpaceOrder()`, `setSpaceOrder()`, etc. Although the designers of the JavaBeans specification refer to this as the get/set pattern, the author agrees with Eckel [2] that this is just a commonplace naming convention rather than a design pattern. Furthermore, there are a number of problems with this get/set approach. Firstly, iterations on these parameters will have to repeat all these functions in boilerplate, e.g., consider printing all the parameters:

```cpp
ostream& DiffusionSolver::print(ostream& os) const
{
    os << "timeOrder = " << timeOrder << endl;
    os << "spaceOrder = " << spaceOrder << endl;
    os << "t0 = " << t0 << endl;
    os << "te = " << te << endl;
```
Table 1
Example parameters for solving the diffusion equation (1). IntVect and RealVect are vectors of integers and real numbers. Their sizes equal the dimensionality of the computational domain.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Type</th>
<th>Identifier</th>
</tr>
</thead>
<tbody>
<tr>
<td>temporal order-of-accuracy</td>
<td>int</td>
<td>timeOrder</td>
</tr>
<tr>
<td>spatial order-of-accuracy</td>
<td>int</td>
<td>spaceOrder</td>
</tr>
<tr>
<td>beginning time of simulation</td>
<td>double</td>
<td>t0</td>
</tr>
<tr>
<td>ending time of simulation</td>
<td>double</td>
<td>te</td>
</tr>
<tr>
<td>number of time steps</td>
<td>int</td>
<td>numSteps</td>
</tr>
<tr>
<td>number of ghost cells</td>
<td>int</td>
<td>numGhosts</td>
</tr>
<tr>
<td>converge criteria of iteration</td>
<td>double</td>
<td>convergeRatio</td>
</tr>
<tr>
<td>max number of iterations per time step</td>
<td>int</td>
<td>maxNumCycles</td>
</tr>
<tr>
<td>lower end of the problem domain</td>
<td>RealVect</td>
<td>xLo</td>
</tr>
<tr>
<td>higher end of the problem domain</td>
<td>RealVect</td>
<td>xHi</td>
</tr>
<tr>
<td>number of cells</td>
<td>IntVect</td>
<td>numCells</td>
</tr>
<tr>
<td>grid size</td>
<td>RealVect</td>
<td>dx</td>
</tr>
<tr>
<td>write solution to a HDF5 database?</td>
<td>bool</td>
<td>writeHDF5Sol</td>
</tr>
<tr>
<td>write error to a HDF5 database?</td>
<td>bool</td>
<td>writeHDF5Err</td>
</tr>
<tr>
<td>the directory to output results</td>
<td>string</td>
<td>writeHDF5Dir</td>
</tr>
</tbody>
</table>

... // repeat for all other parameters

This leads to “code bloating”: the number of lines grows as that of the parameters. It also reminds us of the dubious “copy-and-paste” code-duplication in a subtle form. Secondly, this leads to “interface bloating”: the number of member functions is twice as that of parameters. For classes with a large number of parameters, this might be an unacceptable violation of the fundamental OOP principle that the interface of a class be minimized [4]. Thirdly, hard-coding the names of these parameters make the maintenance of class DiffusionSolver tedious and error-prone; if one were to add or remove some parameters, she would have to go through all functions like print and add additional pairs of get/set functions to the interface of class DiffusionSolver.

Lastly, DiffusionSolver might have a constructor that takes all the parameters:

```cpp
class DiffusionSolver {
```
public:
    DiffusionSolver(int a_timeOrder, int a_spaceOrder,
        double a_t0, double a_te,
        int a_numSteps, int a_numGhosts,
        double a_convergeRatio, int maxNumCycles,
        ...);

...}
}

The task of checking the correct order of the sequence is tedious and error-prone. Worse, a bug caused by switching the order of parameters of the same type is very hard to catch since the compiler won’t complain at all! In fact, if those two parameters have different orders in the header file and the implementation file, the compiler won’t notice either! On top of these, just imagine adding or removing some parameters from the list!!

Often, a programmer frustrated by the get/set approach will just declare the parameters as public data members so that client programs can get/set them freely. Worse than the get/set approach, public members violate ‘encapsulation’, the most important principle of OOP. Also, if the value of a parameter has to be within a certain range to guarantee the correct behaviors of the module, it is easy to enforce and hide this semantics in a set function while it is complicated, if not impossible, to do so for the public member approach.

Ideally, we want to have a single pair of accessing/modifying functions as follows

    /// pseudo signature for accessing function
    ParameterType get(ParameterName) const;

    /// pseudo signature for modifying function
    void set(ParameterName, ParameterType);

    /// pseudo code of looping on parameters
    for (Parameter p=ParameterIterator.begin(); p!=ParameterIterator.end(); p++)
    {
        ... // do something with p
    }

In a dynamically typed language such as python and lisp, implementing the above pseudo-code is easy; however, in a statically typed language like C++, the specific types of Parameter, ParameterName, and so on must be resolved at compile time. To comply to this, we either require all parameters have the same type or employ a compile-time branching mechanism based on parameter type. In Section 2 and Section 3, the author proposes a pattern in each of these
two categories. The targeted use cases are scientific-computing modules with
a large number of parameters and a small number of parameter types.

2 The Enumeration Index Pattern

The enumeration index pattern exploits the C++ enumeration construct by
mapping a list of user-defined names to a set of consecutive and non-negative
integers, which can be used as indices to parameter arrays. The following C++
code illustrates this pattern for the integer parameters in Table 1.

class IntegerManager {
public:
    enum ParmInt {
        timeOrder = 0, spaceOrder, numSteps,
        numGhosts, maxNumCycles, nParmInt
    };
    typedef std::map<ParmInt, int> MAP;

    static char const*const toStr(const ParmInt& p) {
        switch(p) {
            case timeOrder  : return "timeOrder ";
            case spaceOrder : return "spaceOrder ";
            case numSteps   : return "numSteps  ";
            case numGhosts  : return "numGhosts ";
            case maxNumCycles: return "maxNumCycles ";
            default         : return ";
        }
    }

    int getInt(const ParmInt& p) const {
        return m_int[p];
    }

    void setInt(const ParmInt& p, int v) {
        m_int[p] = v;
    }

    void setInts(const MAP& map) {
        for (MAP::const_iterator i=map.begin(); i!=map.end(); i++)
            setInt(i->first, i->second);
    }

    ostream& print(ostream& os) const {
        for (int i=0; i<nParmInt; i++)
            os<<
        return os;
    }
}
ParmInt pInt = static_cast<ParmInt>(i);
    os << toStr(pInt) << " = " << getInt(pInt);
}

private:
    int m_int[nParmInt];
};

After setting the value of the first enumerate to zero, the values of the rest enumerates are automatically assigned to consecutive integers. Consequently, the last enumerate in the list, nParmInt, can be used as the size of the integer array for storing the values of the integer parameters. Although an enumerate list can be mapped to string literals by preprocessing macros, a toStr function is shown here for clarity.

Compare to declaring two member functions for each parameter, this enumeration index pattern has a number of appealing advantages. Firstly, the number of get/set functions remains the same so long as the number of parameter types is constant. Printing the parameters reduce to a loop on the enumeration list. Secondly, adding and removing a parameter can be achieved by simply changing the enumeration list and the corresponding toStr function; the changes automatically propagates to an iterative function such as print due to the enumeration mechanism. Thirdly, the function setInts can be used as a constructor to set all the int parameters in a much more semantically clear way. For example, the client program might read

typedef IntegerManager IM;
IM::MAP map;
map[IM::timeOrder] = 4;
map[IM::spaceOrder] = 4;
... // other int parameters
IM im;
im.set(map);

The association between the keys and values is much easier and clearer than that of the get/set approach: it eliminates the tedious task of checking the correct order of parameters passed to the constructor.

Sometimes the parameters of a class are simple in that (i) they only have a single type (ii) they do not have additional semantics such as dependency on each other. In this case, the enumeration-indexing pattern is an excellent solution. However, as the number of types grows, the interface quickly becomes large since an enumeration list, a toStr function, and a pair of get/set functions are needed for each type; this is analogous to the bloated interface in
Fig. 1. The main idea of the parameter manager pattern. For each parameter type, `ParameterManagerN` has a corresponding associative container that takes the name of a parameter and returns its value in the particular type. The type of the parameter is resolved at compile time while the name of the parameter is passed at run time to the container of the specific type. The dashed box belongs to the interior of a `ParameterManagerN` object.

the simple get/set approach. In addition, it is difficult to enforce constancy or dependence of parameters. This is where the parameter manager pattern comes to help.

3 The Parameter Manager Pattern

The main idea of the parameter manager is illustrated in Fig. 1, where the get function is templated on the type of the parameter. Inside a `ParameterManagerN` object, an associative data structure, e.g. `std::map`, is instantiated for each possible type at compile time. At run time, the container with the specific type of the parameter is queried for the value of the parameter.

3.1 Capture the notion of ‘parameter’

The first step to implement the above idea is to capture the notion of parameter in a templated class. This is also conducive to binding more semantics to a parameter such as constancy, dependency, and default values.

```cpp
template<class T>
```
class Parameter {
public:
    Parameter(const std::string& key,
              const T& defaultValue = T(),
              bool constant = false,
              bool dependent = false) :
        key_(key),
        defaultValue_(defaultValue),
        constant_(constant),
        dependent_(dependent){}

    bool isDependent(void) const { return dependent_; }
    bool isConstant(void) const { return constant_; }
    char const*const key(void) const { return key_.c_str(); }
    const T defaultVal(void) const { return defaultValue_; }

    friend bool operator<(const Parameter<T>& p1,
                          const Parameter<T>& p2) {
        return (p1.key() < p2.key());
    }

private:
    std::string key_;  
    T defaultValue_; 
    bool constant_;   
    bool dependent_; 
};

Now that Parameter has its own abstraction layer, one can easily append more semantics to it.

The next step is to map parameters to their values. For this purpose, std::map is an excellent choice and we wrap around it a class ParameterManager<T>, with the template parameter class T the same as that of Parameter<T>. Although the key of std::map in the following code is Parameter<T> instead of string in Fig. 1, they are essentially the same since the compare operator of the Parameter class is based on its name as a string. Hence, two different parameters must not have the same key.

template<class T>
class ParameterManager {
public:
typedef std::map<Parameter<T>, T> MAP;

virtual ~ParameterManager(){}

const T get(const Parameter<T> &p) const {
    typename MAP::const_iterator mit = map_.find(p);
    if (mit==map_.end()){
        std::cerr << "Warning: Parameter " << p.key() << " not set yet! Return default value...\n";
        return p.defaultVal();
    }
    return mit->second;
}

void set(const Parameter<T> &p, const T &v) {
    if (!p.isConstant())
        map_[p] = v;
    // initialize a constant p, check value
    else if (map_.find(p) == map_.end() && v==p.defaultVal())
        map_[p] = p.defaultVal();
    else
        std::cerr << "Warning: " << p.key() << " = " << p.defaultVal() << " is already set!\n ";
}

void print(std::ostream& os) const {
    for (typename MAP::const_iterator mit = map_.begin(); mit != map_.end(); mit++)
        os << mit->first.key() << " = " << mit->second << std::endl;
}

protected :
    MAP map_;}

If a parameter is not initialized yet, ParameterManager::get will issue a warning and return the default value of the parameter; if a constant parameter is already initialized to its default value, ParameterManager::set will also issue a warning for another attempt to set the value. Compare to getInt/setInt of SolverManager in Section 2, the additional semantics makes the contract more precise and helps to enforce the correct behaviors of the class.
3.2 From one to many

Within one type $T$, we have achieved using a single pair of get/set functions to access/modify the parameters. Also, this provides us with enough ammunition to address the problem of multiple types. Consider the parameters listed in Table 1 in Section 1. Conceptually we are trying to gather a number of ParameterManager<T> instances, each corresponding to a certain type, into one entity. The goal of still having a single pair of get/set functions can be achieved by function overloading and multiple inheritance, thanks to Parameter being a templated and strongly-typed class. Here, the ambiguity is removed by branching different parameter types to different parameter managers.

```cpp
template<class T1, class T2>
class ParameterManager2 : private ParameterManager<T1>,
    private ParameterManager<T2>
{
    public:
    virtual ~ParameterManager2(){}

    const T1 get(const Parameter<T1>& p) const {
        return ParameterManager<T1>::get(p);
    }
    const T2 get(const Parameter<T2>& p) const {
        return ParameterManager<T2>::get(p);
    }

    void set(const Parameter<T1>& p, const T1& v) {
        ParameterManager<T1>::set(p,v);
    }
    void set(const Parameter<T2>& p, const T2& v) {
        ParameterManager<T2>::set(p,v);
    }

    friend std::ostream& operator<<
        (std::ostream& os, const ParameterManager2<T1, T2>& pmm)
    {
        pmm.ParameterManager<T1>::print(os);
        pmm.ParameterManager<T2>::print(os);
        return os;
    }
};
```
Note that the private inheritance closes all the interfaces of individual parameter managers and the ambiguity of multiple inheritance is resolved by function overloading. Without this explicit branching, calling a get/set function of ParameterManager2 will result in a compilation error.

For $N$ bigger than 2, ParameterManager$N$ can be defined similarly. The code-bloating caused by the repetition can be alleviated by defining the following preprocessing macros.

```cpp
/// define a non-constant, independent parameter with default value.
#define PM_DEF(TYPE, NAME, VALUE)
    const Parameter<TYPE> PM_CLASSNAME::NAME(#NAME, VALUE)

/// define a constant parameter
#define PM_DEFC(TYPE, NAME, VALUE)
    const Parameter<TYPE> PM_CLASSNAME::NAME(#NAME, VALUE, true)

/// define a dependent parameter
#define PM_DEFD(TYPE, NAME, VALUE, C)
    const Parameter<TYPE> PM_CLASSNAME::NAME(#NAME, VALUE, C, true)

/// initialize a parameter to its default value.
#define PM_INIT(P)
    set(P, P.defaultVal())

#define PMM_GET(N)
    const T##N get(const Parameter<T##N>& p) const {
        return ParameterManager<T##N>::get(p); }

#define PMM_SET(N)
    void set(const Parameter<T##N>& p, const T##N& v) {
        return ParameterManager<T##N::set(p,v);} 
```

The stringizing operator `#` encloses the argument in double quotation marks, thus `#NAME` will expand to a string literal at compile time. The token-pasting operator `##` concatenates its argument to tokens immediately before it to form a new token. For example, `T##N` will expand to `T1`, `T2`, and so on depending on the value of the argument $N$. The above macros reduce repetition and clarify the code. Using them, ParameterManager3 is defined as follows.

```cpp
template<class T1, class T2, class T3>
class ParameterManager3 : private ParameterManager<T1>,
    private ParameterManager<T2>,
    private ParameterManager<T3>
{
```
In ParameterManager3, the insertion operator ‘<<’ is serial in that the corresponding function ParameterManager<T>::print is called for each type within one scope; a get or set function is parallel in that the macros generate three functions for it.

Alternatively, when \( N \) is large, ParameterManager\( N \) can be automatically generated by meta-programming, the main enabling mechanisms being a type list, multiple inheritance, and template-based recursion. See, e.g., Chapter 3.13 in the wonderful book by Alexandrescu [1]. However, the approach in [1] can not be applied here without modification because of serial methods such as the insertion operator mentioned in the previous paragraph. In other words, one must clearly differentiate serial methods in ParameterManager\( N \) from parallel ones in order to define them correctly. In addition, there are other reasons a simple repetition is preferred here: (1) For the targeted use cases of scientific computing, the number of types is not large; (2) The interface of ParameterManager is thin and very static; (3) It is self-contained; (4) It is much easier to understand.

### 3.3 Which parameters to manage?

By now the reader might have raised a few questions. For examples, a constant parameter can be declared as a static constant data member, so why bother with the parameter manager? Should parameter manager manage all data members of a class? If not all, which parameters should be managed? In order
to answer these questions, let us examine the relationship between a module and its parameters.

A module takes some input and produces some output, it represents a contract between the client who uses the module and the programmer who writes the module. To realize the contract, the two parties have to agree on some conditions: those limit the possible values of the input and the scope of the contract are called preconditions; those applies to the output are called postconditions. Together they unambiguously define the problem to be solved by the module. In summary, the behaviors of a module depend on the preconditions and postconditions, which are represented by the parameters of the module.

A parameter is said to be fundamental if it quantifies either a precondition or a postcondition. The set of fundamental parameters spans a use-case space, of which each consisting point, represented by a combination of the values of the fundamental parameters, uniquely determines the behaviors of the module. Usually there is more than one way to choose the set of fundamental parameters for a certain module, in this case we require the set of fundamental parameters to have the minimum number of parameters.

The fundamental parameters should be well organized — clearly represented, easily accessed and modified. Indeed, this is the main motivation of the parameter manager pattern. At this point the questions posed in the beginning of this section are easy to answer. Internal data members irrelevant to pre- and post-conditions should not be managed. Whenever a data member, constant or non-constant, belongs to the set of fundamental parameters, it should be managed by ParameterManager. As for a data member whose value depends on others, it obviously does not belong to the set of fundamental parameters. Whether it should be managed by ParameterManager is a design choice. For example, the grid size $dx$ depends on number of the cells, the starting and ending position of the computational domain. However, it should be managed since it will be frequently accessed by other modules. To sum up, the fundamental parameters and frequently used dependent data members should be managed by ParameterManager.

To enforce the additional semantics, two pure virtual functions are added to ParameterManagerN:

```cpp
template<class T1, class T2, ..., class TN>
class ParameterManagerN : private ParameterManager<T1>,
    ...,
    private ParameterManager<TN>
{
public:
    virtual void updateDependentParameters(void) = 0;
```
A derived class is responsible for defining the default values and dependency of its parameters. Making the virtual functions pure is a friendly reminder through the compiler if this is forgotten.

3.4 The parameter manager pattern in action

Finally we can apply the parameter manager pattern to the example of solving the diffusion equation.

```cpp
#include "ParameterManagerN.H"

class DiffusionSolver :
  public ParameterManager6<bool, int, double, string, IntVect, RealVect>
{
  public:
    // Independent Parameters : preconditions and postconditions
    static const Parameter<int> timeOrder;
    static const Parameter<int> spaceOrder;
    static const Parameter<double> t0;
    static const Parameter<double> te;
    static const Parameter<int> numSteps;
    static const Parameter<int> numGhost;
    static const Parameter<double> convergeRatio;
    static const Parameter<int> maxNumCycles;
    static const Parameter<RealVect> xLo; // constant
    static const Parameter<RealVect> xHi;
    static const Parameter<IntVect> numCells;
    static const Parameter<bool> writeHDF5Sol;
    static const Parameter<bool> writeHDF5Err;
    static const Parameter<string> writeHDF5Dir;
    // Dependent Parameters
    static const Parameter<RealVect> dx;

    DiffusionSolver()
    {
      initialize();
      updateDependentParameters();
    }
```
The handlers of the parameters in Table 1 are declared as static constant members. As self-documenting code, the declarations clearly capture the preconditions and postconditions of the diffusion solver module.

Regardless of different types, the value of all declared parameters can be accessed or modified using the single pair of get/set functions inherited from ParameterManager6, as shown in the following example.

```plaintext
typedef DiffusionSolver DS;
DiffusionSolver ds;
ds.set(DS::timeOrder, 4);
ds.set(DS::convergeRatio, 1.e-10);
ds.set(DS::writeHDF5Err, false);
const double t0 = ds.get(DS::t0);
const int spaceOrder = ds.get(DS::spaceOrder);
const bool writeHDF5Sol = ds.get(DS::writeHDF5Sol);
...
```

Inside the constructor, initialize() set the properties of the independent parameters for the first time, then the value of the dependent parameters are updated by updateDependentParameters(). The definition file shows more details.

```plaintext
#include "DiffusionSolver.H"
#include "ParameterManagerMacros.H"

#define PM_CLASSNAME DiffusionSolver

// Table of default values

PM_DEF ( int, timeOrder, 4);
PM_DEF ( int, spaceOrder, 4);
PM_DEF ( double, t0, 0.0);
PM_DEF ( double, te, 1.0);
PM_DEF ( int, numSteps, 10);
PM_DEF ( int, numGhost, 2);
```
PM_DEF ( double, convergeRatio, 1.e-12);
PM_DEF ( int, maxNumCycles, 20);
PM_DEFC( RealVect, xLo, RealVect::Zero);
PM_DEF ( RealVect, xHi, RealVect::Unit);
PM_DEF ( IntVect, numCells, IntVect::Unit);
PM_DEFD( RealVect, dx, RealVect::Unit, false);
PM_DEF ( bool, writeHDF5Sol, true);
PM_DEF ( bool, writeHDF5Err, true);
PM_DEF ( string, writeHDF5Dir, "hdf5");

///--------------------------------------------------------------

void GridsManager::initialize(void)
{
  PM_INIT(timeOrder);
  PM_INIT(spaceOrder);
  PM_INIT(t0);
  PM_INIT(te);
  PM_INIT(numSteps);
  PM_INIT(numGhosts);
  PM_INIT(convergeRatio);
  PM_INIT(maxNumCycles);
  PM_INIT(xLo);
  PM_INIT(xHi);
  PM_INIT(numCells);
  PM_INIT(writeHDF5Sol);
  PM_INIT(writeHDF5Err);
  PM_INIT(writeHDF5Dir);
}

void GridsManager::updateDependentParameters(void)
{
  const RealVect tmp = (get(xHi)-get(xLo))/get(numCells0);
  set(dx, tmp);
}

RealVect::Zero and RealVect::Unit represent zero and unit vectors with
their size as the dimensionality of the computational domain. dx is shown as
an example of the dependent parameter. Alternatively, one could choose dx as
an independent parameter and numCells as a dependent parameter. So long
as the number of independent parameters is minimum, choosing independent
parameters is influenced by use cases or even personal preference.

Although each parameter identifier is repeated twice, the macros nicely or-
nize the definition and initialization of the parameters. Consider augmenting
Table 1 with two additional columns, one for default values and the other for
dependency or constancy. A code-generator can be written to automatically generate `DiffusionSolver.cpp` (except the function for updating dependent parameters) from the augmented table. We will not elaborate on the code-generator since it can be easily done in a scripting language such as python or even in C++ itself.

### 4 Concluding Remarks

The author has proposed two patterns for managing parameters in scientific computing. Both patterns use a single pair of get/set functions to access and modify the parameters. The enumeration index pattern is a good choice if all the parameters are of a single type and do not have additional semantics. The parameter manager pattern is a better choice for multiple types and additional parameter semantics.

Using the parameter manager pattern can be automated as much as possible. One only have to inherit from a `ParameterManagerN` class and specify for each parameter its identifier, type, default value, constancy, and dependency. It is then straightforward to convert this specification table to similar code shown in Section 3.4 by a code generator, except that updating dependent parameters cannot be automated.

Scientific computing often uses input files to record the values of the parameters. Using the parameter manager pattern, an input file can be mapped to a parameter manager object by a helper class that parses each key in the input file and then sets the corresponding parameter value for that object.

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### References


