Computational aspects and performance evaluation of the IFS-XIOS integration

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Abstract

The increment of the spatial resolution for computational Earth system models is nowadays one of the main concerns of the scientific community in order to solve more complex problems, and thus, achieve more accurate solutions to the reality. However, the new complexity requires more computing power that only cutting-edge supercomputers can provide. This requires to use sophisticated HPC techniques to efficiently use the computational resources. In addition, such high resolutions lead to generate an enormous amount of data to meticulously represent accurate solutions.

Current Earth system models usually have inefficient sequential I/O schemes that used to run low grid resolutions, where the generated amount of data for the simulation results was not particularly big. However, sequential I/O schemes do not scale with current models where a lot of parallel resources are used. In order to address this issue, the most adopted approach is to use scalable parallel I/O solutions that offer both computational performance and efficiency.

This document analyzes the I/O process of IFS, one of the most important atmospheric models used in Europe. IFS has two different output schemes, the MF I/O server and a sequential I/O scheme, being the latter the only available in OpenIFS. This sequential scheme is characterized by being inefficient, since all the output is made through the master process. Therefore, here it is presented an easy-to-use development that integrates an asynchronous parallel I/O server called XIOS into IFS. Moreover, different optimization techniques, such as computation and communication overlap, are applied in the integration development to minimize the I/O overhead in the resulting IFS execution.

The results show that the use of XIOS in IFS to output data is certainly good. This new parallel scheme has reduced significantly the execution time of the original sequential scheme. Using the proper configuration, XIOS proves to be a scalable I/O server that keeps a low overhead regardless the amount of IFS processes and the output size. Furthermore, XIOS offers a series of benefits that shorten the critical path of IFS experiments by concurrently running the post-processing task along the IFS execution: data format conversion, online post-processing and CMIP-compliant output. In this scenario, the total execution time is greatly reduced.
1 Introduction

Over the years, computing power of supercomputers has grown exponentially [1]. Scientific applications from all disciplines have benefited of it by increasing the complexity of the computational models used. This new complexity turns into extra computational cost added using different methods, from the increase in the horizontal or vertical resolution of spatial grids to the interaction in parallel of new components simulating additional features (biochemistry, ice, etc.). When this kind of applications have available more computational resources, they can afford to solve more complex problems leading to more accurate solutions. For example, the increase of the horizontal spatial resolution, in ocean models for areas near to the coast, allow to correctly simulate some small-scale processes such as eddies [2]. However, a higher resolution implies to generate much more data because the representation of these small-scale processes requires more points. One of the main problems is to efficiently write this such amount of data along the execution of applications. The second main problem is to post-process this data afterwards. Post-processing is the phase where data is transformed using defined operations, such as data format conversion, computation of new derived variables known as diagnostics, etc.

A good example where this extra computational power has been used is in the field of Earth System Modelling (ESM). Numerical weather and climate prediction has considerably improved the accuracy of forecasts, predictions and projections due to the increase of the grid resolution [3]. Obviously, taking advantage of this extra computational power requires to properly use efficient High Performance Computing (HPC) techniques. Traditionally, the focus of improvement from a computational point of view in ESM has been the calculation and communication of algorithms [4]: the increase of the Instruction Level Parallelism (ILP); the massively parallelization of code using heterogeneous platforms such as Graphic Processing Units (GPUs) or Intel Xeon Phis; the memory access; the compiler tuning; the optimization of Message Passing Interface (MPI) patterns; and many more.

One of the most important issues studied to improve the computational efficiency are workload imbalances between processes [5][6]. Workload imbalances happen when the amount of work to be solved for each one of the processes is not evenly distributed, so this causes that some processes have more work to solve than others. Then, processes that finish earlier have to wait for the slowest ones. This problem gets worse when more processors are used, since the workload distribution is more complex. Another issue to be taken into account are the possible workload imbalances due to the type of grid used in Earth system models. Grids are used as spatial representation to discretise the Earth’s surface to solve equations on evenly distributed grid-points. However, depending on the type of grid, the domain decomposition could be more or less complex. As an example of the type of grid used for this work, using regular reduced Gaussian grids has some intrinsic problems on the Earth poles [7][8], where the computation and communication among neighbors (to fulfill spatial dependencies) is much more expensive compared to other regions of the Earth, such as the Equator. This is because it is necessary to use a more complex domain decomposition.

Although the ESM community has done considerable efforts to improve models from an algorithmic point of view, there is a very important aspect that has almost been forgotten during many years because it was not significant enough in the past: the Input/Output (I/O). Due to the new complexity of models, it will become really difficult to exploit computational resources to achieve more accurate results without performing an efficient I/O, because Earth system models are significantly increasing the number of variables to be output, as well as the output frequency.
of variables. The output process is usually performed at the end of specific time steps, during the execution of the model. This process has increased the execution time during the last years, since more and more data has to be stored from higher grid resolutions. In the end, an inefficient I/O process could lead to a serialization where all resources are waiting to complete this critical task. Moreover, since we are rapidly approaching the exascale era, the I/O part will become a truly bottleneck [9], mainly because of the produced huge amount of data. Exascale computers are future machines that will have at least one exaFLOP, or a billion billion calculations per second. On these machines, models will be potentially able to simulate ultra-high resolutions, but if the I/O process is not optimized at the same time, everything will slow down and will not be possible to achieve the future ambitious goal of having more complex models.

One of the models that could be in this situation is the Integrated Forecasting System (IFS). IFS [10] is a global data assimilation and forecasting system developed by the European Centre for Medium-Range Weather Forecasts (ECMWF) and used by several institutions in Europe. IFS has two different output schemes: a sequential output scheme which gathers all data in the master process, and the Météo-France (MF) I/O server which is an efficient I/O scheme that uses dedicated resources to perform the I/O. While ECMWF uses the MF I/O server for its operational forecasts, external institutions have to use the sequential output scheme. This is the case of one of the global climate models most used in Europe, the EC-Earth model, which uses a limited version of IFS as its atmospheric component.

EC-Earth [11] is a global coupled climate model, which integrates a number of component models in order to simulate the Earth system. It is used for problems encompassing from seasonal-to-decadal climate prediction to climate change projections and paleoclimate simulations. In Figure 1 there is a scheme showing the components of EC-Earth.

![EC-EARTH components](image.png)

Figure 1: Components used in the EC-Earth model

The two main components are IFS as the atmospheric model and the Nucleus for European Modelling of the Ocean (NEMO) as the ocean model, both coupled using OASIS3-MCT. Climate models such as EC-Earth are a very good example to prove that an efficient I/O will be
needed for the future. They are run to simulate really extensive periods of time which turns into an enormous amount of data which has to be saved for prediction and projection analyses. For example, in a recent EC-Earth experiment consisting in a 100 years projection, it was generated a total amount of 244 TB of useful data. Only for IFS were used 4416 processors, consuming about 2.47 million computing hours.

Additionally, there are other tasks that are included typically in the workflow of an Earth system model, apart from the task used to solve the governing equations along the time. These tasks are known as pre-processing and post-processing and they are used to process the input and output data. For example, in the pre-processing task there could be the preparation of the initial conditions which are needed to run any model, whereas in the post-processing task there could be the computation of derived variables, known as diagnostics. Diagnostics are a type of variables computed from other variables, typically prognostic variables which are directly predicted by the model.

Figure 2 illustrates a really simple experiment which contains three tasks: pre-processing, simulation and post-processing. The time needed to complete the three tasks is known as critical path, because there are dependencies between them and must be sequentially executed.

![Critical path of a basic experiment with three tasks sequentially executed](image)

In particular, the post-processing task in EC-Earth is characterized by being quite expensive, because it needs to transform General Regularly-distributed Information in Binary form (GRIB) files output by IFS to Network Common Data Format (netCDF) files. This is necessary because IFS was originally developed for Numerical Weather Prediction (NWP), where the data format standard is GRIB. GRIB was designed to offer high performance for I/O operations, since in operational weather forecast, the time-to-solution is a critical process. However, the data format used in climate modelling as an accepted standard is netCDF. In addition, EC-Earth needs to compute expensive diagnostics using variables from both IFS and NEMO.

1.1 Motivation

As it has been explained, the sequential output of IFS is the scheme used in the IFS version of EC-Earth. This was not a problem using the resolution required for operational experiments, because the I/O did not represent a significant part in the total execution time.

However, we have recently started to perform very complex experiments in collaboration with different institutions from Europe, under the Horizon 2020 Process-based climate siMulation: AdVances in high-resolution modelling and European climate Risk Assessment (PRI4 Technical Memorandum No. 825).
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MAVERA project [12]. This project aims to simulate using higher resolution and it requires to produce a large number of variables. As a consequence, the community has experienced a considerably slowdown in the execution time, not only due to the higher resolution of the computational part, but also especially due to the I/O part, because it represents about 30% of the total execution time.

This is one of the critical issues to be solved for the community, because it will be present in new future experiments which will require similar PRIMAVERA configurations or even more complex.

For this reason, it is necessary to solve this problem by identifying which are the present and future community needs to use an efficient and functional I/O approach. The first need is to write in netCDF data format instead of GRIB like IFS currently does, because climate modelling works with netCDF format. The second need is to produce netCDF files as requested by the CMIP data request. The Coupled Model Intercomparison Project (CMIP) [13] is a standard experimental protocol for studying the output of coupled Atmosphere-Ocean Global Circulation Models (AOGCMs). The third need is the ability to perform online post-processing, which means that the data to be output is processed along the simulation. Those three needs are very important to avoid the costly current post-process that we have to perform. The fourth need, from a computational point of view, is to find an efficient and scalable I/O approach able to exploit parallel resources.

Additionally, the new complexity of Earth system models where different components interact among them implies new needs where data produced by each component will not be independently processed anymore. Since we currently compute some diagnostics using variables from both IFS and NEMO, we will need a mechanism to concurrently compute them online for both components along the simulation, avoiding to do it in the post-processing task.

In order to fulfill the aforementioned community needs, different I/O tools were studied, selecting for this work the most suitable one. The XML Input/Output Server (XIOS) [14] is an asynchronous MPI parallel I/O server that we chose to be integrated with IFS and as a consequence will be used in the future OpenIFS version that will substitute IFS in EC-Earth. OpenIFS is a free licensed and simplified version of IFS.

In addition, since NEMO is already using XIOS for outputting data, the future version of EC-Earth, which will use OpenIFS and NEMO, will be able to compute online diagnostics through XIOS with variables from both components at the same time.

Therefore, the XIOS integration will fulfill all the community needs and will increase the computational efficiency of IFS and will reduce the critical path by avoiding the post-processing task. Figure 3 shows how the critical path will be improved by using XIOS for IFS OpenIFS and EC-Earth.
1.2 Objectives

The main objective of this project is to improve the I/O performance of IFS to reduce the total execution time and achieve a better computational efficiency. This is to get more throughput, which means to write more data in less time.

We also set the objective of reducing the critical path by removing the post-processing devoted to perform costly operations such as GRIB to netCDF data format conversion or diagnostics computation.

In addition, we also have the objective to increase the usability of IFS by using an easier output configuration file compared to the current approach.

In order to achieve the objectives, we identify the following tasks:

- Perform the state-of-the-art of I/O techniques used in HPC and ESM to identify the most appropriate approach to be used by IFS.
- Develop an integration between IFS and XIOS.
- Do a performance analysis of the development to detect potential bottlenecks and use proper optimization techniques to finally obtain an efficient integration.

1.3 Document organization

This document is organized as follows. In Section 2 we research about the current available I/O schemes and discuss why we chose XIOS. We give an overview of IFS and XIOS in Section 3. We continue the document explaining the development done for the integration in Section 4. The next Section 5 is devoted to explain the performance analysis and optimization of the development. The computational performance of the integration is evaluated in Section 6. Finally, we conclude the work in Section 7.
2 State-of-the-art

We have presented the motivation and objectives of this project, explaining why IFS needs a new I/O scheme that fulfills the community requirements. In this section, we will review the state-of-the-art of the I/O solutions used on HPC especially in the area of ESM focusing on parallel I/O.

In Section 2.1 we give a general overview about what is I/O and how it works the sequential I/O. Then, in Section 2.2 we explain what is parallel I/O and the different approaches that we can use. We also explain some well-established I/O libraries in HPC including MPI-IO, Hierarchical Data Format (HDF) and netCDF. After that, we explain that there is a particular type of parallel I/O that uses dedicated resources, which are called I/O servers. Finally, in Section 2.4 we compare different I/O solutions and justify why we chose XIOS.

2.1 I/O overview

The action of reading and writing data, commonly known as I/O, is a basic and essential process of almost all HPC applications to communicate with the outside world. Earth system models have traditionally performed the I/O using sequential writing [9]. It is typically done using the Portable Operating System Interface (POSIX) I/O Application Programming Interface (API). Figure 4 shows the basic layers of the I/O stack involved for reading and writing using POSIX I/O. Functions such as open, write or close, directly work with files.

In parallel applications, the sequential I/O implies to send all data to the master process, which performs the sequential write. In the meantime, the rest of the processes wait until this process is completed.

![Figure 4: Sequential I/O stack traditionally used by HPC applications (Reproduced from [15])](image)

Sequential I/O was fine several years ago because the amount of data was not too high and storage systems were able to deal with it. However, as we explained in the introduction Section 1, the increase of computing power enables the capacity to execute more accurate simulations, which leads to generate more output data. In addition, the path to exascale will accentuate this problem. This means that current sequential I/O schemes are not useful anymore, even if they use optimized techniques, because it will not scale. Therefore, applications need to use parallel I/O.

A good example of an efficient I/O optimization changing the sequential scheme to a parallel one is in the Community Atmosphere Model (CAM) [16]. They published this work in 2008, 10 years ago, which indicates that they were already aware of the inherent problems of the sequential I/O for parallel applications. This work proved that the transition from sequential
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to parallel I/O gives really good improvements in terms of computational performance.

However, the need of more computational power led to use much more processors in CAM and for this reason, the parallel I/O scheme introduced in 2008, was replaced some years ago by the Parallel I/O library (PIO).

2.2 Parallel I/O

In order to increase the scalability of Earth system models, the current feasible approach is to make the I/O scalable as well, that is, use parallel I/O.

Parallel I/O is the ability to perform multiple input/output operations at the same time, such as simultaneously writing several files or concurrently writing into different regions of the same file from different processes.

Applying this concept to Earth system models [17], the main idea is to involve all the processes of a model so that they balance or re-distribute the data to be output from subdomains in a way that writing is efficiently performed and as fast as possible.

To this aim, there are two strategies commonly adopted [18]:

- Writing multiple files: data is output among several files (as many as MPI tasks). This means that each MPI task is responsible of its subdomain. The main advantage is that is very scalable, although in some file systems the creation of a lot of files can be a problem. In addition, post-processing is needed to joint files. On the left side of Figure 5 there is a scheme showing how each process writes its own file. As said, this can be done using the POSIX I/O API or using MPI-IO point-to-point operations (explained later on in this section).

- Writing one file: data is output into one single file. In this case, no post-processing for joining files is needed, but the scalability is more difficult to achieve, which depends on the implementation and the number of MPI processes used, mainly due to overheads caused by conflicted I/O operations from all processes. On the right side of Figure 5 there is a scheme showing how all processes write into a shared file using MPI-IO collective operations.

Figure 5: Common writing strategies: on the left side the strategy for writing multiple files and on the right side the strategy for writing one file (Reproduced from [18])

Nevertheless, the two previous strategies could not be suitable to scale models using a huge
number of processes. Therefore, a feasible solution could be to use an intermediate solution. That is, writing a file for a subset of processes. Figure 6 shows a scheme of this intermediate strategy.

![Intermediate I/O strategy: processes are grouped in subsets to share a file](image)

Since POSIX I/O cannot offer the possibility to implement parallel I/O, it is necessary to use a library able to offer this feature. In this case, the most popular one is MPI-IO. It is not easy to use and as a consequence has not been adopted by many applications. However, it is indirectly the most used parallel I/O library, because many other user-friendly high-level libraries are built on top of MPI-IO, such as netCDF or HDF. Since these high-level libraries are easy to use, they are commonly used for Earth system models.

This new approach to perform I/O changes the I/O stack by adding two new layers as can be seen in Figure 7. In this new stack, applications use high-level libraries that usually offer a powerful API for efficiently organizing complex data objects with the corresponding metadata. At the same time, high-level libraries use I/O middlewares to efficiently store data into storage systems through parallel file systems.

![Parallel I/O stack adopted by HPC applications](image)

In the following points we review some well-established I/O libraries in HPC applications, i.e., MPI-IO for the I/O middleware layer, and HDF and netCDF for the high-level I/O library layer:

- MPI-IO: the MPI 2.0 standard was extended by adding specific parallel I/O functionality. Since POSIX was designed for serial I/O, MPI-IO aims to offer a high-level
interface to split the reading and writing of data across several processes taking advantage of MPI messages. It is possible to use both individual or collective I/O operations. The interface syntax is based on MPI subroutines.

Currently, many different I/O libraries are built upon MPI-IO, because it offers a good performance and deals with low-level implementation details. The performance can be tuned by setting a lot of different parameters because each HPC machine has its own hardware and software configuration.

- **HDF5**: the Hierarchical Data Format (HDF) [24] is a set of tools, libraries and file formats that are used to manage and store large amounts of data. It can also represent very complex data objects and a wide variety of metadata. In addition, it is able to store multi-dimensional arrays. HDF5 is the latest version.

- **NetCDF**: the Network Common Data Format (netCDF) [25][26] is a set of software libraries and self-describing, machine-independent data formats that support the creation, access, and sharing of array-oriented scientific data. The latest version is netCDF4 which can use HDF5 file format for storing data, but it is also compatible with the previous (and first) version of netCDF, which is netCDF3. Regarding the parallel use of netCDF, which by default uses serial writing, there are two mechanisms:
  - For writing in netCDF3 file format, it is necessary to use Parallel-netCDF (PnetCDF), built upon MPI-IO.
  - For writing in netCDF4 file format, it is necessary to use the parallel functionality of HDF5, built upon MPI-IO.

### 2.3 I/O servers

In order to keep improving the performance and the scalability of Earth System models, it is possible to go one step further and use some nodes exclusively dedicated to I/O. The model processes do not need to deal with the I/O, they only have to send the data to I/O nodes. This has the advantage that they can continue with the simulation without spending time outputting data.

This approach is known as I/O servers [27], which are the responsible of writing data into the storage system in order to theoretically hide the disks latency from the model processes and use as efficiently as possible the network bandwidth by using techniques such as aggregation.

There are different strategies to send data from model processes to I/O servers. Communication is typically done through MPI by using synchronous or asynchronous operations. Furthermore, I/O servers can have different communication patterns. One I/O server could collect a subset of subdomains, i.e., each I/O server aggregates local data from a subset of subdomains for all variables. Or one I/O server could collect a subset of global variables, i.e., each I/O server aggregates local data from all subdomains (global domain) for a subset of variables.

I/O servers need to be configured by specifying many different parameters: how to write data, the data format, which I/O method internally use, the number of fields to be written and their dimensions, the data decomposition used in the model, etc. This can be hardcoded, done using...
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an external configuration file which is parsed at runtime, and/or using an API because some variables are dynamically set up on the model during runtime.

In addition, some servers can have extra functionality such as online post-processing, data conversion or data compression. If users want to use it, they simply have to set up the proper parameters in the configuration file (or through the API).

Figure 8 shows a scheme about I/O servers, where there are the model processes communicated with I/O servers. I/O servers, before writing data into the storage system, can perform online post-processing while model processes continue with the simulation.

![Figure 8: Overview of an I/O server. On the left there are the model processes which send data to the I/O servers (orange). Then, while models processes keep running the simulation, I/O servers write data into the storage system.](image)

There are several I/O servers available in the literature. They can be generic to any kind of HPC application, such as ADaptable I/O System (ADIOS) [28] and Damaris [29], but also they can be designed specifically for ESM, such as CDI with parallel I/O (CDI-pio) [30], Climate Fast Input/Output (CFIO) [31] and XIOS [32].

### 2.4 Discussion: comparison and choice

In the motivation Section 1.1 of this project, we have clearly explained the specific needs that we have, so all of them should be fulfilled by one I/O tool. We have reviewed several options in the literature that could potentially be integrated with IFS to solve the I/O bottleneck.

If we focus on the execution time of the simulation, as well as only outputting data, we think that it is not clear which could be the fastest I/O server, because the respective authors report good performance, and I/O is something very machine-dependent and even context-dependent. For example, you could be running your scalability tests in a moment where there are other users that are executing I/O-intense applications. This would speed down your tests.

Nevertheless, there are other aspects that influence our choice. In both IFS standalone and IFS
within EC-Earth the post-processing phase is needed, so if we can move this work to the I/O servers we can save a lot of resources and time. In the current critical path of an experiment, after the simulation we have to write temporary files that will be read in the post-processing phase to be then written again once they are processed. With online post-processing, after the simulation we would be able to directly write the definitive files. I/O servers such as CDI-pio and CFIO do not offer online post-processing. In ADIOS and Damaris, there is the possibility to add a plugin to transform data, but in case of being possible, it would be needed a lot of extra work to add those post-processing functionality that in XIOS is already available.

Furthermore, EC-Earth contributes with climate simulations to the CMIP project, where data organization must follow an strict standard that XIOS is aware. Despite the fact that the other I/O servers can write in netCDF, they do not take into account the CMIP standard.

Last but not least, as we exposed in the introduction, in EC-Earth we have to calculate diagnostics that are derived using fields from both IFS and NEMO. Since NEMO is already using XIOS, if IFS was using XIOS as well, we would be able to calculate these diagnostics online instead of doing it at the post-processing phase. ADIOS may not support online diagnostics using variables from more than one component. However, if it was possible, we would have much more work, because it would be needed to integrate ADIOS with IFS and NEMO.

Therefore, according to what we need and what I/O servers offer, it is quite irrefutable that we have to choose XIOS.
3 Context

In this section we will explain the two main components used in this project. In Section 3.1 we give an overview of the scientific part of IFS as well as an explanation about the computational part, including the decomposition and parallelization of local domains, the data structures and the two available output schemes. Then, in Section 3.2 we explain the main features of XIOS and how it is used. We give an overview of all the XIOS elements: axis, domain, grid, field, file, filters and performance variables.

3.1 IFS description

We will mainly focus on the technical and computational part of IFS and only briefly describing the scientific part.

3.1.1 Brief overview

IFS is an operational global meteorological forecasting model and data assimilation system developed and maintained by ECMWF. It is a spectral model that discretises the Euler equations of motion, resolving flow features to approximately 4-6 grid-cells at the nominal resolution. The subgrid-scale features and unresolved processes are described by atmospheric physics parametrizations. There are many different unresolved physical processes in the atmosphere, such as radiation, clouds and subgrid turbulent motions. In Figure 9 there is an overview of the different physical processes.

Figure 9: Overview of the unresolved physical processes in the atmosphere (Reproduced from [34])

The dynamical core of IFS is hydrostatic, two-time-level, semi-implicit, semi-Lagrangian and applies spectral transforms between grid-point space (where the physical parametrizations and
advection are calculated) and spectral space. In the vertical the model is discretised using a finite-element scheme. A reduced Gaussian grid is used in the horizontal.

3.1.2 Decomposition and parallelization of local domains

IFS is an [MPI]+Open Multi-Processing (OpenMP) hybrid model that uses a sophisticated data structure to be efficiently parallelized. Code snippet 1 shows the current data structure used in IFS. The idea of the parallelization applied is that fields are processed using a blocking technique, where the first NPROMA dimension can be adjusted by the user at run-time to fit the memory cache. Depending on the size of NPROMA, there will be more or less NGPBLKS blocks. The 2D i-j horizontal dimension is transformed using these two new dimensions.

```
1 REAL Model_Data(1:NPROMA, 1:NFLEVG, 1:NFIELDS, 1:NGPBLKS)
```

*Code snippet 1: Data structure used in IFS*

The other two dimensions remain the same, which are NFLEVG (or vertical dimension) and NFIELDS (the number of fields). However, note that these two dimensions are not in the outer dimensions of the array. This implies that for a given NGPBLKS block, all fields are processed. Code snippet 2 shows the OpenMP parallelization applied in IFS for its data arrays. In addition, Figure 10 shows the NPROMA blocking used in IFS.

```
1 !$OMP DO SCHEDULE(STATIC)
2 DO iblock = 1, NGPBLKS
3 DO ifld = 1, NFIELDS
4 DO ilvl = 1, NFLEVG
5 DO i = 1, NPROMA
6     Model_Data(i, ilvl, ifld, iblock)
7 END DO
8 END DO
9 END DO
10 !$OMP END DO
```

*Code snippet 2: OpenMP parallelization used in IFS*
3.1.3 Data structures

In IFS there are some major data structures to store all data for spectral and grid-point fields. For spectral data, IFS uses the YOMSP module, in which the arrays SPA3 and SPA2 hold the 3D and 2D state variable spectral fields respectively.

Nevertheless, we will focus on grid-point fields. There are two core data-structures: GMV and GFL.

The GMV structure contains prognostic variables involved in the semi-implicit: wind components, temperature, surface pressure, vorticity, etc. All GMV fields have a spectral representation.

This is quite a fixed structure which is supposed to have almost no modification. Data arrays are accessed using pointers, as many as GMV fields. In addition, there can be 3D and 2D fields: the 3D ones are stored in the GMV array and the 2D ones in the GMVS array. GMV fields do not have attributes.

On the other hand, the GFL structure contains the rest of variables: specific humidity, snow, rain, ozone, etc. Unlike GMV, GFL structure is much more flexible and can be easily extended with additional fields. It only contains 3D grid-points fields that may have a spectral representation. Data is stored in array GFL.

One of the main characteristics are attributes. They are used to govern the behaviour of the individual fields of the GFL structure. The idea is to loop over all fields in GFL and perform the action defined by the setting of the appropriate attribute. However, it is still possible to treat fields separately through individual pointers.
3.1.4 IFS output schemes

IFS has two different output schemes, one better than the other, since one was replaced by the other to improve the throughput:

- Sequential output: this is the slowest one, since the output is sequentially performed by the master process. It performs a gather of all subdomains from the rest of processes, to build the global domain and write it into the storage system. This type of output is not scalable, so it has an important negative impact on the performance of the model. Although this output scheme is contained in all versions of IFS, it is never used except in OpenIFS and EC-Earth. As a consequence, in huge simulations of EC-Earth we are suffering a considerable slowdown.

- MF I/O server: the Météo-France I/O server was introduced in IFS to avoid the low performance of the sequential output scheme. It uses the concept of dedicated processes as servers to perform the I/O, such as XIOS. Although it offers really good throughput, there are some issues: it does not perform post-processing (done by FullPos), it writes data using GRIB format and it is not available in OpenIFS (future version for EC-Earth).

3.2 XIOS description

3.2.1 Overview

The XML Input/Output Server (XIOS) \cite{14,27,37,38,39,40} is an asynchronous MPI parallel I/O server that is used by Earth system models to avoid contention in the I/O. It focuses on offering high performance to achieve very high scalability with support for high-resolution output. XIOS is developed by the IPSL with an Open Source CEA CNRS INRIA Logiciel Libre (CeCILL) License. It has the following features:

- Usability in the definition and management of the I/O with a user-friendly XML configuration file.
- Avoid the I/O performance issue with dedicated parallel and asynchronous servers.
- Post-processing of fields can be performed online using an internal parallel workflow and dataflow.

XIOS is especially targeted to Earth system models with these characteristics:

- Coupled models
- Long simulations
- A lot of data is generated
- Contribute to the CMIP project
Interestingly, all the previous points (except the last one that depends on the purpose of the model) are inherent in climate models, such as EC-Earth. They are made of several coupled components that perform really long simulations. This implies to generate a lot of data that in some cases is used to contribute to the CMIP project.

In Figure 11 there is an overview of the schematic architecture used in XIOS. Each one of the model processes run its own XIOS client using the XIOS API. This is part of the client side, i.e., it is run on the model processes. Then, XIOS clients communicate data to XIOS servers using asynchronous MPI messages. They are run on independent nodes with regard to the model nodes. This is the server side, which uses its own MPI communicator to perform online post-processing over the received data. After that, XIOS servers can write post-processed data into the storage system using two different strategies: one single file or multiple files (one per XIOS server). The whole configuration is described in the iodef.xml file. In Figure 11, is used the one single file strategy.

![Figure 11: Overview of the XIOS architecture. Model processes are communicated with the XIOS servers using asynchronous MPI messages. All the framework is configured using an XML file (Reproduced from [14])](image)

Furthermore, although Figure 11 shows an XIOS configuration using the server mode (dedicated I/O processes), it is also possible to use the client mode. In client mode, XIOS servers are not used and as a consequence, XIOS clients are the responsible of doing online post-processing and writing data into the storage system, either using one single file or multiple files. This implies that model processes cannot continue with the simulation until they have finished post-processing and data writing.

Regarding the aggregation strategy used to send data from clients to servers, XIOS re-distributes clients’ data between servers as evenly as possible to use an optimal balance. Figure 12 illustrates an small example of 6 clients and 3 servers, where data is re-distributed so that servers have a proportional amount of data.
3.2.2 Concept

The philosophy of XIOS is really simple: at each time step the model can expose its data to XIOS using just one subroutine. But first, it is needed to update the calendar of XIOS to inform which is the current time step of the model. This is done using the subroutine of the Code snippet 3.

```
CALL xios_update_timestep(ts)
```

*Code snippet 3: XIOS subroutine to update the calendar*

Then it is possible to use the respective subroutines to read (Code snippet 4) or write (Code snippet 5) data. They must be called for each one of the model variables or fields that we want to read or write.

```
CALL xios_recv_field("field_id", field)
```

*Code snippet 4: XIOS subroutine to read a field*

```
CALL xios_send_field("field_id", field_out)
```

*Code snippet 5: XIOS subroutine to write a field*

Both subroutines have two arguments, the first one is the identifier of the variable or field and the second one is the data array, which can have different dimensions depending on how was defined.

Since the approach of XIOS is to call these two subroutines at each time step, there could be some situations where this is not necessary. For example, if a field is disabled in the XML configuration file so that it will not be output, we could avoid to send data that will not be used.
The Code snippet 6 shows how to use the function that queries if a field is used in the simulation.

```
1 IF (xios_field_is_active("field_id")) THEN
2 CALL xios_send_field("field_id", field_out)
3 ENDIF
```

*Code snippet 6: XIOS subroutine to query if a field is active*

There is another similar situation: if a field is not written, but it is used to calculate a diagnostic. In this case if the sampling frequency to post-process the diagnostic is lower than the time step frequency, in some time steps the data of the field is not needed. Therefore, we could avoid to send data that will not be used. We can use the optional second argument of the previous function setting it up to true. The function of the Code snippet 7 queries if a field is active or not in the current time step.

```
1 xios_field_is_active("field_id", .true.)
```

*Code snippet 7: XIOS subroutine to query if a field is active in the current time step*

Alternatively, we can set the field attribute `check_if_active` to true so that XIOS internally will check if a field is active before sending any data. Code snippet 8 shows how to do it. It is equivalent to use the function of the Code snippet 7.

```
1 <field_definition>
2  <field id="field_id" grid_ref="grid_id" operation="instant" check_if_active="true" />
3 </field_definition>
```

*Code snippet 8: XML definition to output a field if active in the current time step*

Finally, one important aspect of the XIOS philosophy is to make the initial configuration as easy as possible and posterior re-configurations to be as quick as possible by avoiding re-compilations. This is simply achieved by changing parameters in the XML configuration file. Therefore, there are two types of setup that are complemented:

- Static setup through the XML file. It is parsed at runtime at the beginning of the execution to modify the XIOS internal workflow and the user output definition. It follows a hierarchical approach using the inheritance concept.
- Dynamic setup through the Fortran API. It is used to modify the XML definition, or extend it. For example, the grid setup should be done dynamically since subdomain decomposition is different according to the number of MPI tasks and is done at model runtime.

The key point is that if the integration of XIOS is properly done, users only need to make use of the static setup. The dynamic setup is only needed in particular cases, probably for model...
developers. For example, the model internally computes a new diagnostic that should be output through [XIOS]. It would be needed to make some easy changes to send the new diagnostic data array to [XIOS].

3.2.3 Initialization

There are two steps to initialize [XIOS]:

- **XIOS** initialization: in this step [XIOS] is initialized, the [XML] is parsed and [MPI] may be initialized (if it has not already been done). The `MPI_COMM_WORLD` communicator is shared between the model and [XIOS], so it is returned a local communicator to be potentially used by the model. This is done with the subroutine of the Code snippet 9.

- Context initialization: it performs the configuration of the context associated with "context_id" defined in the [XML] file. In addition, it opens the scope where to put setup subroutines: geometry, calendar, fields, etc. This is done with the subroutine of the Code snippet 10.

```
1 CALL xios_initialize("code_id", return_comm=communicator)

Code snippet 9: Subroutine to initialize XIOS
```

```
1 CALL xios_context_initialize("context_id",communicator)

Code snippet 10: Subroutine to initialize the XIOS context
```

3.2.4 Finalization

The two steps of the initialization are inversely undone:

- Context finalization: close contexts once they are processed. This is done with the subroutine of the Code snippet 11.

- **XIOS** finalization: close servers, opened files and generate the performance report to know if we are using the optimal number of servers. It also finalizes [MPI] if it was initialized by [XIOS]. This is done with the subroutine of the Code snippet 12.

```
1 CALL xios_context_finalize

Code snippet 11: Subroutine to finalize the XIOS context
```
3.2.5 Grid definition

It is necessary to set up on XIOS the grid that is exactly used by the model and the data decomposition for each MPI process as well.

XIOS is able to use grids of any dimension: 0D, 1D, 2D, 3D, etc. The most commonly used is a 3D grid which is made of a 1D-vertical axis and a 2D-horizontal domain. Axis and domain are the two elements used in XIOS to define any kind of grid.

Axes are generally used to describe the vertical direction of a grid. There are several variables to describe how data is stored in memory and mapped into the grid. Figure 13 shows the basic variables of the axis element.

Domains describe the type of horizontal layer that maps the Earth’s surface. In XIOS there are four different domains available:

- Regular Cartesian
- Curvilinear
- Reduced Gaussian
- Unstructured

There is an example of a reduced Gaussian domain in Figure 14. As mentioned, this type of domain is the new one used in IFS.
Figure 14: Example of a reduced Gaussian domain (Reproduced from [14])

As in the axes, domains also have some basic variables used to describe the geometry of the global domain and local subdomains. Figure 15 shows the basic variables, which for describing a regular Cartesian domain would be enough, but for a reduced Gaussian one, we would need to set up some different and a bit more difficult variables. This will be explained in the development Section 4.

Figure 15: Basic variables to describe the global domain and a subdomain (Reproduced from [14])
3.2.6 Fields

Fields are typically declared through the XML file as it shows Code snippet 13.

```
<field_definition>
  <field id="temp" grid_ref="grid_3d" />
  <field id="precip" grid_ref="grid_3d" />
  <field id="pressure" domain_ref="domain_2d" />
</field_definition>
```

*Code snippet 13: Example of an XML field definition*

Data of fields is distributed according to the underlying grid description. They have several attributes that will be associated in the output file: name, standard name, unit, etc. In addition, there are other configurable parameters such as the sampling frequency, the compression level, etc.

3.2.7 Files

The declaration of files is similar to the fields one, also using the XML file. In Code snippet 14 there is one file defined with the name "daily_output", containing several fields with different online post-processing options.

```
<file_definition>
  <file name="daily_output" freq_output="1d">
    <field field_group_ref="fields_3d" operation="average" />
    <field field_group_operation="instant">
      <field field_ref="temp" name="temp_inst" />
      <field field_ref="pressure" name="pressure_inst" />
    </field_group>
    <field field_ref="pressure" operation="average" />
  </file>
</file_definition>
```

*Code snippet 14: Example of an XML file definition*

There are some configurable parameters: write one or multiple files, compression, netCDF version, etc.

3.2.8 Filters

An essential part of XIOS for performing online post-processing are filters. Depending on the type of post-processing operations, there are several different filters that are applied to fields during runtime. Internally, filters are applied to data fluxes with a timestamp, which represent fields. Figure 16 shows a workflow with its respective fluxes and filters to generate diagnostics to be written into files. These workflows are internally used by XIOS, users do not have to deal
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with them.

![Diagram of workflow applying filters to fields]

*Figure 16: Example of a workflow applying filters to fields*

There are three types of filters:

- **Arithmetic filters**: they combine fluxes of the same timestamp together applying arithmetic operations. Fluxes must be represented with the same grid. Code snippet 15 shows how to represent the two following arithmetic operations using the XML:

  \[ C = \frac{A + B}{A \cdot B} \]

  \[ D = \frac{e^{-C \cdot D}}{3} \]

- **Time integration filters**: they integrate a flux over a period of time. There are different types of operations: once, instant, maximum, minimum, average and accumulate. It is possible to chain time filters with "@". Code snippet 16 shows an example.

- **Spatial filters**: they are used to change the geometry of the fields. There are different parallel operations:
  - Data extraction: zooming, slicing, etc
  - Global or spatial reduction: mean, max, min, etc
  - Horizontal interpolation
  - Polynomial vertical interpolation
  - Pressure levels interpolation
  - Connectivity discovery
  - ...

Code snippet 17 shows a code to interpolate a regular domain to an unstructured domain.

```xml
1 <field id="A" />
2 <field id="B" />
3 <field id="C">(A + B)/(A*B)</field>
4 <field id="D">exp(-C*this)/3</field>
```

*Code snippet 15: Example of an arithmetic filter*
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3.2.9 Performance tuning

At the end of the execution, XIOS generates individual performance reports for all clients and servers to know details about memory consumption and different execution times (total, waiting, ratio, etc).

There are different options to tune the performance of XIOS:

- Test the two possible run modes: attached mode or server mode. Theoretically, server mode should give more performance, but additional computational resources are needed.

- Test the two possible writing modes: single file or multiple files. Theoretically, multiple files should give more performance due to higher scalability, but post-processing to merge all files is needed.

- Test different values for some memory buffer parameters.
3.2.10 CMIP

Some climate models are tuned to perform simulations that will contribute to the CMIP project. The Coupled Model Intercomparison Project (CMIP) is a standard experimental protocol for studying the output of coupled AOGCMs. The ongoing version is the sixth, known as CMIP6. One of the requisites is that data must follow a very strict standard, so almost all models need to post-process data to fulfill the standard. This process is known as Climate Model Output Rewriter (CMOR) post-processing or CMORization.

As we mentioned, XIOS is able to ideally produce netCDF files according to the CMIP data request, being ready to be distributed and published. However, this is not automatically done, so a minimum user effort is needed to write the proper XML configuration file.

This XIOS feature avoids (or at least minimize) to perform the costly and slow CMORization.
4 Development of the IFS-XIOS integration

4.1 Overview

In previous Section 3 we have given an explanation of both IFS and XIOS to understand the details of their integration.

Figure 17 shows the IFS-XIOS integration scheme implemented and illustrates how the different parts of both components are interconnected. In green, it is shown the IFS processes which execute the client side of XIOS through its API. They send data using asynchronous MPI communications to XIOS servers. XIOS servers are run on server side, which are represented in orange. Finally, servers send data to the storage system (in purple) through system calls. Both XIOS clients and servers are configured with the iodef.xml file.

Furthermore, this figure also shows that post-processing is performed in both clients and servers. This is because depending on the type of post-processing, some operations are performed on client side, such as horizontal interpolations, and some other on server side, such as netCDF compression.

There is a pair of scripts, model and run_parallel, that are in charge of executing the whole integrated system. In particular, run_parallel executes both binaries in Multiple-Program Multiple-Data (MPMD) mode using aprun.
If we focus on the structure of our code, we have followed the coding standards of IFS. They have different prefixes for file names, where in general "yom" is used for variables, "su" is used for setup routines and "c" is used for control routines. We have implemented three different files:

- **yomxios.F90**: it contains variables related to XIOS. For example the context handle, time step, data arrays, etc.
- **suxios.F90**: it contains all the routines needed to initialize XIOS from IFS. There are three public subroutines: **suxios_init**, **suxios_fin** and **suxios_ctxt**.
- **cxios.F90**: it contains one public routine called **ifs_xios_send_fields** to send data.

The development has been designed so that it is really simple to use and maintain. It is possible by calling the previous mentioned four public subroutines:

- **suxios_init**: it is called at the beginning of IFS because MPI is initialized by XIOS
- **suxios_fin**: it is called at the end of IFS when MPI is no needed anymore, because XIOS finalizes MPI
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- suxios_ctx: it is called during the IFS setup. It performs the configuration of XIOS with the parameters of IFS.

- ifs_xios_send_fields: it is called at the end of each time step to send fields to XIOS.

The ifs_xios_send_fields subroutine is internally designed so that it has three different parts or steps. This three steps are: update calendar, NPROMA blocks gather and send fields. If IFS is running an output time step, all three steps will be sequentially executed; otherwise if IFS is running a non-output time step, only the update calendar step will be executed.

Figure 18 shows how these three steps are executed at the end of an output time step. For illustrative purposes, in this example all time steps perform output.

\[\text{IFS time steps}\]

\[\text{C}_x\text{- Update calendar}\]

\[\text{G}_x\text{- Gather}\]

\[\text{S}_x\text{- Send}\]

Figure 18: Output scheme developed for IFS: all three steps, update calendar, NPROMA blocks gather and send fields are sequentially executed at the end of an output time step.

In the following sections we will explain more in detail how the previous files are implemented and the changes in the scripts. In Section 4.2 we explain the necessary steps to perform the XIOS setup from IFS, i.e., pass the IFS parameters that XIOS requests. It includes the design of both yomxios.F90 and suxios.F90 files. Then, in Section 4.3 we explain how we implemented the grid-point fields transfer. This is basically the design of the cxios.F90 file. Finally, in Section 4.4 we explain the necessary changes in the IFS environment, including model and run_parallel scripts, to be able to run the whole integrated system.

4.2 XIOS setup

In order to set up XIOS to be fully operational, we have to follow a series of steps. First of all, XIOS must be initialized, which internally initializes MPI. Then, we have to setup the whole context. This part includes the calendar, which is very important to have both IFS and XIOS fully coordinated when the simulation progresses along the time. It also includes the geometry of IFS. This is the definition of the IFS grid, including the horizontal domain and the vertical axis. Thus, XIOS knows how received data from different independent processes has to be placed over
the world. Finally, once the simulation finishes, we have to finalize XIOS, which internally also finalizes MPI.

### 4.2.1 Data and variables module

According to our development structure, we have a module in yomxios.F90 that contains variables related to the XIOS context, calendar variables, identifiers for the axes and domains, and three data arrays that are used to send fields to XIOS. Code snippet 18 shows the yomxios module.

```fortran
MODULE yomxios
USE PARKIND1, ONLY : JPIM, JPRB
USE xios
IMPLICIT NONE
SAVE

! XIOS context
TYPE(xios_context) :: context_handle
CHARACTER(len=3) :: model_name = "ifs"
CHARACTER(len=3) :: ifs_context = "ifs"

! Calendar management
TYPE(xios_date) :: time_origin
TYPE(xios_date) :: start_date
TYPE(xios_duration) :: time_step

! Axes definition
CHARACTER(len=12) :: model_axis_name = "model_levels"

! Domains definition
CHARACTER(len=16) :: gaussian_domain_name="reduced_Gaussian"

! Arrays to gather NPROMA blocks and send fields to XIOS
REAL(KIND=JPRB), ALLOCATABLE :: xios_gmv(:,,:), xios_gmvs(:,), xios_gfl(:,)

END MODULE yomxios
```

Code snippet 18: Overview of the data and variables module

### 4.2.2 Initialization

The first step is to initialize XIOS and MPI. It is possible to initialize MPI through IFS or XIOS. According to the XIOS documentation, they recommend to initialize it through XIOS. This means that we have to ensure that IFS does not call MPI_Init.

As mentioned, XIOS will use the MPI_COMM_WORLD communicator for both components, it will use its own communicator, and it will return a local communicator to be used by IFS. IFS is already prepared to work with MPI_COMM_WORLD or with a provided local communi-
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cator. This avoided us to adapt IFS to support provided communicators.

The mechanism to change the IFS communicator is through two variables: LMPLUSERCOMM and MPLUSERCOMM. Code snippet 19 shows how to do it. Setting up LMPLUSERCOMM to true, we are specifying to use a provided communicator, which will be provided with the MPLUSERCOMM. When we initialize XIOS the subroutine xios_initialize returns a local communicator.

```
! Enabling the usage of MPLUSERCOMM communicator, rather than MPI_COMM_WORLD
LMPLUSERCOMM = .TRUE.

! Initialization of XIOS and definition of the MPLUSERCOMM communicator to be used by IFS
CALL xios_initialize(model_name,return_comm=MPLUSERCOMM)
```

Code snippet 19: Mechanism to change IFS communicators

This code is part of the public suxios_ini subroutine.

4.2.3 Finalization

After the simulation, it is needed to finalize XIOS and MPI. Code snippet 20 shows how to do it. First of all, we deallocate some arrays that are used for sending fields to XIOS (it is explained in Section 4.3).

Then, we have to call xios_context_finalize and xios_finalize subroutines to finalize XIOS and MPI. Since in the initialization we specified to IFS to use a provided communicator, it will not call MPI_Finalize, which is called by XIOS.

```
! Deallocating XIOS arrays
DEALLOCATE(xios_gmv)
DEALLOCATE(xios_gmvs)
DEALLOCATE(xios_gfl)

! Finalization of XIOS context
CALL xios_context_finalize()

! Finalization of XIOS and MPI
CALL xios_finalize()
```

Code snippet 20: MPI and XIOS finalization

This code is part of the public suxios_fin subroutine.

4.2.4 Context

One important step is the XIOS context setup. Code snippet 21 shows how to perform the setup. There are three first subroutines to start the IFS context definition scope, which is closed using the xios_close_context_definition. It contains three main private subroutines:
• **ifs\_xios\_set\_calendar:** it is used to inform XIOS which type of calendar uses IFS, the start date of the simulation and the duration of the time step. Section 4.2.5 explains how it is implemented.

• **ifs\_xios\_set\_axis** and **ifs\_xios\_set\_domain:** both subroutines are used to set up the geometry of the model. It is essential to inform XIOS about the distribution of each one of the grid-points over the Earth, as well as how they are distributed between all IFS processes. Thus, XIOS can output valid data, where each grid-point has a longitude and a latitude over the Earth’s surface. Section 4.2.6 explains how they are implemented.

After the context definition close, we allocate some arrays used to send fields to XIOS (the same arrays that we deallocated in Section 4.2.3, i.e., in finalization).

```fortran
! Context initialization
CALL xios\_context\_initialize(ifs\_context, MPLUSERCOMM)
CALL xios\_get\_handle(ifs\_context, context\_handle)
CALL xios\_set\_current\_context(context\_handle)

! Date setting
CALL ifs\_xios\_set\_calendar

! Definition of axes
CALL ifs\_xios\_set\_axis(YDGEOMETRY)

! Definition of domains
CALL ifs\_xios\_set\_domain(YDGEOMETRY)

! Close context definition
CALL xios\_close\_context\_definition()

! Allocating XIOS arrays
ALLOCATE(xios\_gmv(YDGEOMETRY\%YRGEM\%NGPTOT,YDGEOMETRY\%YRDIMV\%NFLEVG) )
ALLOCATE(xios\_gmvs(YDGEOMETRY\%YRGEM\%NGPTOT))
ALLOCATE(xios\_gfl(YDGEOMETRY\%YRGEM\%NGPTOT,YDGEOMETRY\%YRDIMV\%NFLEVG))
```

This code is part of the public **suxios\_ctxt** subroutine.

### 4.2.5 Calendar

XIOS needs to know the type of calendar that IFS is using, in this case the Gregorian one. It is set up calling xios\_define\_calendar. Then, we set up the time origin and the start date using xios\_set\_time\_origin and xios\_set\_start\_date subroutines respectively. In this case, we use the same date and time for both. IFS stores the initial date using format AAAAMMDD in variable NINDAT, and the initial time in seconds in variable NSSSSS.

Finally, the last parameter to be set up is the time step that IFS is using. IFS stores the time step in a variable called TSTEP, and it is passed to XIOS using the xios\_set\_timestep subroutine.
Code snippet 22 shows how the calendar setup is done.

```fortran
INTEGER(KIND=JPIM) :: year, month, day, hours, minutes, seconds
year = NINDAT/10000
month = MOD(NINDAT/100, 100)
day = MOD(NINDAT, 100)
hours = NSSSS/3600
minutes = (NSSSSS - hours*3600)/60
seconds = NSSSSS - hours*3600 - minutes*60
CALL xios_define_calendar(type="Gregorian")
!
CALL xios_set_time_origin(time_origin=xios_date(year, month, day, hours, minutes, seconds))
!
CALL xios_set_start_date(start_date=xios_date(year, month, day, hours, minutes, seconds))
!
! Updated date = start_date + NSTEP*YRRIP*TSTEP
time_step%second = YRRIP*TSTEP
CALL xios_set_timestep(time_step)
```

*Code snippet 22: XIOS setup to use IFS calendar*

This code is part of the private *ifs_xios_set_calendar* subroutine.

### 4.2.6 Geometry

One of the most difficult parts in setting up XIOS is the IFS geometry transfer to XIOS. Since we are working with a 3D grid, it consists of two parts: the 1D axis definition and the 2D domain definition.

Code snippet 23 shows how to perform the vertical axis definition. Using the loop, it basically builds and array with as many positions as the number of vertical levels. Each position of the array contains the number of the vertical level. IFS stores the number of vertical levels in variable NFLEVG.

Then, using the *xios_set_axis_attr* subroutine, we set up different attributes for the vertical axis: total number of vertical levels, the array containing the vertical levels, the units (we are working on sigma levels, so there are no units), and the direction of the axis, which is positive.
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```
INTEGER(KIND=JPIM) :: i
REAL(KIND=JPRB) :: j
REAL(KIND=JPRB), ALLOCATABLE :: zML(:)

! Definition of model levels axis
ALLOCATE(zML(YDGEOMETRY%YRDIMV%NFLEVG))

j = 1.0
DO i = 1, YDGEOMETRY%YRDIMV%NFLEVG
  zML(i) = j
  j = j + 1.0
END DO

CALL xios_set_axis_attr(model_axis_name, n_glo=YDGEOMETRY%YRDIMV%NFLEVG, value=zML, unit="-", positive="up")

DEALLOCATE(zML)
```

**Code snippet 23: Vertical axis definition**

Code snippet 23 is part of the private *ifs_xios_set_axis* subroutine.

The second part is the domain definition. First of all, we need to build the local domain data (Code snippet 24) by iterating over all grid-points in the global domain to find and store in the `i_index` array which are the grid-points of the local domain. Thus, each IFS process will communicate to its XIOS client which are the local grid-points.

```
! Local domain data
!
j = 0
DO i = 1, n_glo
  IF (YDGEOMETRY%YRMP%NGLOBALPROC(i) == MYPROC) THEN
    j = j + 1
    sasha(i) = i - 1
  END IF
END DO
```

**Code snippet 24: Local domain data setup**

After that, there is a bigger loop (Code snippet 25) which is in charge of setting up the longitudes and latitudes of each one of the local `i_index` grid-points. They are stored in arrays `lonvalue_1d` and `latvalue_1d`. Furthermore, we also have to set up the boundaries for each grid-point. Boundaries are used to delimit the area that represents each grid-point. There are 4 corners for the area of each grid-point. The easiest way to determine the boundaries of each grid-point is using the middle point between two latitudes and the middle point between two longitudes. We have to be careful in the first and last latitudes, since there are not grid-points on the poles. Boundaries are stored in arrays `bounds_lon_1d` and `bounds_lat_1d`. 
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Code snippet 25: Longitudes, latitudes and corners setup

Finally, all these arrays that we have build, are passed to XIOS using the \texttt{xios\_set\_domain\_attr} subroutine. In addition, we send other parameters, such as the type of domain (Gaussian), the number of global grid-points and the number of local grid-points. We also specify that the local domain and the data domain have the same size, i.e., there will be no halos in the data arrays that we will send for each one of the fields. Code snippet 26 shows how it is done.
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1. Define global domain and local domain index
   
   ! Define global domain and local domain index
   CALL xios_set_domain_attr(gaussian_domain_name, type='gaussian', ni_glo=ni_glo, ibegin=0, ni=ni,
   \( i_{\text{index}}=i_{\text{index}} \))

2. Define local domain data
   
   ! Define local domain data
   CALL xios_set_domain_attr(gaussian_domain_name, data_dim=1, data_ibegin=0, data_ni=ni)

3. Define longitudes and latitudes for grid-point cells
   
   ! Define longitudes and latitudes for grid-point cells
   CALL xios_set_domain_attr(gaussian_domain_name, lonvalue_1d=lonvalue_1d, latvalue_1d=latvalue_1d)

4. Define cell’s boundaries
   
   ! Define cell’s boundaries
   CALL xios_set_domain_attr(gaussian_domain_name, nvertex=nvertex, bounds_lon_1d=bounds_lon_1d,
   \( \rightarrow \text{bounds_lat}_1d=\text{bounds_lat}_1d \))

5. Deallocate memory
   
   DEALLOCATE(i_index)
   DEALLOCATE(lonvalue_1d, latvalue_1d)
   DEALLOCATE(bounds_lon_1d, bounds_lat_1d)
   DEALLOCATE(zrgauslat)

---

**Code snippet 26: Horizontal domain definition**

Code snippets 24, 25 and 26 are part of the private *ifs_xios_set_domain* subroutine.

4.2.7 *iodef.xml* file

In order to complement the dynamic setup done through the [XIOSAPI](https://example.com), we also had to implement the *iodef.xml* file to perform the static setup. It contains several sections that we will briefly mention.

The first one is the field definition (Code snippet 27). There are declared 12 3D fields and one 2D field: temperature, ozone, surface pressure, etc. They have a reference to the corresponding grid.
Then, there is the file definition (Code snippet 27). It contains two different files, file_3D_bench and file_2D_bench, which are used to output 3D and 2D fields respectively. Both files have an output frequency of 3 hours, they output instant values (no post-processing applied), they are output in multiple file mode, the frequency sampling of each field is 3 hours and files are forced to be flushed into the storage system at the end of 1 simulated day.
After that, there are the definitions of axes, domains and grids (Code snippet 29). We simply have one vertical axis, one reduced Gaussian domain, and two different grids: `model_levels` to represent 3D fields and `surface_fields` to represent 2D fields.

Finally, there are some variables which are used to set up from a computational point of view: size of the buffers, server or attached mode, info level, print performance reports, etc. Code snippet 30 shows the variables used.
4.3 Grid-point fields transfer

Once the setup is done, at the end of each time step, we have to inform XIOS about the current time step and query if we have to send any field. If so, we will send the requested data.

Code snippet 31 shows how the time step is updated and how to process just one field, in this case a 3D GMV field which is temperature. First of all, we update the time step calling the xios_update_calendar subroutine. IFS stores the current time step in variable NSTEP. After that, we query if the field that we want to send is active in the current time step. In the example, the subroutine xios_field_is_active is used to query if we have to send the temperature field.

If the field is active, we have to build an XIOS-style array before sending data. If we remember what we explained in Section 3.1.2, IFS uses a data structure that is not usual, having the following dimensions: NPROMA, NFLEVG, NFIELDS and NGPBLKS. This does not match with the XIOS data arrays, where we need to send an array with the following dimensions: longitude, latitude and vertical levels. When it uses unstructured or reduced Gaussian grids, XIOS merges the first two dimensions, so it needs and array with these two dimensions: unidimensional 2D domain and vertical levels. Therefore, we have to re-organize data of each field before sending it, which means to perform the NPROMA blocks gather.

To perform the gather, in Code snippet 31 we iterate over blocks of size NPROMA, until all the NGPTOT grid-points of the subdomain are processed. Then, for each block we have to iterate all the block grid-points for all NFLEVG vertical levels. We do not need to iterate over the total number of fields because we are gathering data of just one field.

Note that the gather is at the subdomain level, i.e., intra-node shared-memory, so MPI communications are not needed at all.

Once data is re-organized following the XIOS-style array, we can send it calling the xios_send_field subroutine.
Computational aspects and performance evaluation of the IFS-XIOS integration

```fortran
! Update calendar of XIOS with the current time step
CALL xios_update_calendar(NSTEP)

! GMV
!
! Temperature
IF (xios_field_is_active("t", .TRUE.,)) THEN
  ! Array accessing optimized for GMV, not for xios_gmv
  DO jstglo = 1, YDGEOMETRY%YRGEM%NGPTOT, YDGEOMETRY%YRDIM%NPROMA
    icend = MIN(YDGEOMETRY%YRDIM%NPROMA,YDGEOMETRY%YRGEM%NGPTOT−jstglo ↪ +1)
    ibl = (jstglo−1)/YDGEOMETRY%YRDIM%NPROMA + 1
    DO jlev = 1, YDGEOMETRY%YRDIMV%NFLEVG
      i = jstglo
      DO j = 1, icend
        xios_gmv(i,jlev) = YDFIELDS%YRGMV%GMV(j,jlev,YDFIELDS%YRGMV%YT0%MT,ibl)
        i = i + 1
      END DO
    END DO
  END DO
END IF
```

One important thing to highlight are the XIOS thread-safe calls. Since IFS uses OpenMP for intra-node parallelization and XIOS does not support threading, we must ensure that only one thread at each MPI process calls XIOS subroutines. Although in previous code snippets we have not used any kind of mechanism, in our development we actually used the OpenMP SINGLE construct. Code snippet 32 illustrates the mechanism that we followed to have thread-safe calls.

```fortran
!OMP SINGLE
! One single thread region
!OMP END SINGLE
```

### 4.4 Environment setup

Besides the source code development, it is really important to adapt the scripts in order to run IFS and XIOS. In the following sections we will give an overview of the main changes that were needed.

#### 4.4.1 XIOS compilation

One of the challenges was the XIOS compilation. Since we are working in a Cray machine and the IFS 43r3 version and its libraries that we are using are only compiled with Cray, we had to compile XIOS using Cray as well. The problem is that there are not many Cray machines...
in Europe, so XIOS is almost always compiled with Intel or GNU. This probably implies that
XIOS is not usually tested with Cray and as a consequence, there are some issues.

First of all, the parallel compilation does not work in Cray compilers due to different bugs,
something that has been reported to the XIOS developers to solve the problem. For example, a
parallel compilation with Intel using eight threads, it takes about five minutes, while a sequential
Cray compilation it takes about eight hours.

The bigger issue was related with the version of the Cray Developer Toolkit (CDT). By default,
it loads the version 17.03, which successfully compiled XIOS, but it always failed when passing
the XIOS tests. We tried different compilation flags, but this did not solve the issue. At the
end, we discovered that using an older version of the CDT the 16.04 version, we were able to
pass the XIOS tests.

However, we also had other minor issues, such as that by default, Cray compiler generates mod
files in capital letters, but they were not recognized when linking with XIOS. We solved this
issue by using the -ef flag, which generates .mod files in lowercase letters.

Finally, we used a conservative compilation for XIOS i.e., using the -O1 optimization flag,
because we read that for some old Cray compilers there was a bug using the -O3 flag.

4.4.2 Including and linking XIOS

XIOS needs to be included and linked like all libraries. This is not difficult, but it was needed
to find the exact place between the large amount of IFS scripts.

Code snippet 33 shows how the include can be done.

```
1 VPATH := . $(VPATH) /perm/ms/spesiccf/c3xy/xios2/inc
```

**Code snippet 33: XIOS including**

Code snippet 34 shows how the link can be done. Note that we also include the libcray-c++-rts
library, since XIOS is programmed in C++.

```
1 −L/opt/cray/cce/8.4.6/CC/x86−64/lib/x86−64 −lcray−c++−rts \
2 −L/perm/ms/spesiccf/c3xy/xios2/lib −lxios
```

**Code snippet 34: XIOS linking**

4.4.3 Parallel netCDF and HDF5 versions

Although IFS output is in GRIB format, it is linked through environment modules with netCDF
and HDF5, libraries which have been covered in Section 2.2. IFS uses sequential versions of them,
but depending on the XIOS configuration, it needs the netCDF and HDF5 parallel versions.

We changed the two lines of Code snippet 35 by the two lines of Code snippet 36.
Computational aspects and performance evaluation of the IFS-XIOS integration

1  LOAD_MODULE cray-netcdf/$\{\text{netcdf\_version}\}$
2  LOAD_MODULE cray-hdf5/$\{\text{hdf5\_version}\}$

*Code snippet 35: Old netCDF and HDF5 modules*

1  LOAD_MODULE cray-netcdf-hdf5parallel/$\{\text{netcdf\_version}\}$
2  LOAD_MODULE cray-hdf5-parallel/$\{\text{hdf5\_version}\}$

*Code snippet 36: New netCDF and HDF5 modules*

**4.4.4 Model script**

One of the tasks in the workflow of the IFS experiments is called *model*, which is in charge of running the model. The associated script is obviously called *model* and it also calls other scripts, like the *run\_parallel* one, which is in charge of executing the IFS binary. The *run\_parallel* script modification is explained in next Section 4.4.5.

However, before calling *run\_parallel* we have to prepare the environment. Code snippet 37 shows some preliminary steps. First of all, there are two variables that will be substituted by XCdp. Through XCdp we can set up at run-time variables which are between percentage symbols. Then, we have to copy the *iodef.xml* file and replace the string *XIOS\_USING\_SERVER* to indicate if we will run XIOS in attached or server mode. We also have to copy the XIOS binary if it is run in server mode.

There is another important setup regarding the Lustre striping. Lustre \[43\] is an open-source parallel file system that supports many requirements of leadership class HPC simulation environments. Lustre stores a file in one or more Object Storage Target (OST) devices. By default, it uses one OST but it is possible to divide the file into chunks that are stored in different OSTs. This is known as striping, and it is used for performance purposes, especially for really big files. ECMWF recommends to stripe a file of size more than a few hundred GBs.

According to our *iodef.xml* file, XIOS will store netCDF files into the *xios\_output* directory, but depending on the configuration, files can be really huge. To be cautious, we apply an striping of 4 OSTs for each one of the netCDF files that will be stored in *xios\_output*.

If we had not changed the number of OSTs per file, we could affect the performance of the whole system, including the rest of users that are performing I/O at that moment. This is because we could fill up the OST in the parallel file system where the big netCDF file would be in. When such OST runs out of space, it also affects other user’s jobs.
Once all setup is done, it is possible to call run_parallel to execute the [IFS] binary. Code snippet 38 shows a simplified version of how to use run_parallel. If we want to run [IFS] and [XIOS] in [MPMD] mode, we need to pass to run_parallel a file that contains in different lines each one of the binaries and their arguments. Otherwise, we call run_parallel as usual, which means that [IFS] will use [XIOS] in attached mode.

```bash
if [ $XIOS_USING_SERVER = true ]; then
cp $XIOS_PATH/bin/xios_server.exe .
fi
```

**Code snippet 38: MPMD mode preliminary setup in model script**

4.4.5 Supporting MPMD mode

The last major change that we did was in the run_parallel script to run [IFS] and [XIOS] in [MPMD] mode. This script was already prepared to run binaries in [MPMD] mode, but it seems that was never tested and it contained some errors. We fixed them and extended the script to be fully integrated with [XIOS]. We will show some code snippets that are useful to understand what we did.

At the beginning, we set up through XCdp several variables that are used by [XIOS] (Code snippet 39). They are related to the affinity of [XIOS]: number of [XIOS] servers, number of servers per node, number of server per [NUMA] socket within a node, etc. We also specify that we do not want threads, since [XIOS] does not use [OpenMP].
Another important variable is `MPICH_DMAPP_APP_IS_WORLD`, which by default is set to 0. When we were testing the [MPMD](https://en.wikipedia.org/wiki/Message Passing Interface) mode, we were having a deadlock in the execution without any kind of error. We spent a lot of time trying to understand why it was happening. We fixed the error by using the setup of Code snippet 40. Cray MPICH internally uses DMAPP, and if we set `MPICH_DMAPP_APP_IS_WORLD` to 1, it uses [MPMD](https://en.wikipedia.org/wiki/Message_Passing_Interface) for [MPI](https://en.wikipedia.org/wiki/Message_Passing_Interface), but treats each DMAPP application as if it is a distinct job. [MPI](https://en.wikipedia.org/wiki/Message_Passing_Interface) ranks are globally contiguous and global [MPI](https://en.wikipedia.org/wiki/Message_Passing_Interface) communication is possible.

```
export MPICH_DMAPP_APP_IS_WORLD=1
```

*Code snippet 40: Environment variable to run in MPMD mode*

Finally, in Code snippet 41 we show the command line that will execute both [IFS](https://www.ecmwf.int/en/ifs) and [XIOS](https://www.xiossoftware.com). This command line is the result of executing the code that we implemented in [run_parallel](https://www.ecmwf.int/en/). It has several parameters that determine the geometry of the parallel job.

Binaries are separated by a colon and their arguments are passed after the name. For example, [IFS](https://www.ecmwf.int/en/ifs) has several arguments, while [XIOS](https://www.xiossoftware.com) does not have anyone. Then, each binary has its own aprun parameters: the number of processes (-n), the number of processes per node (-N), the number of processes per [NUMA](https://en.wikipedia.org/wiki/Non-uniform_memory_access) socket (-S), the number of cores per each process (-d) and the number of logical threads, i.e., hyperthreading (-j). In addition, there are global parameters for both binaries, such as [CPU](https://en.wikipedia.org/wiki/Central_processor_unit) binding (-cc), amount of memory per each process (-m) and the [NUMA](https://en.wikipedia.org/wiki/Non-uniform_memory_access) memory affinity (-ss).

Note that for [IFS](https://www.ecmwf.int/en/ifs) we also set the number of OpenMP threads.

```
aprun −cc cpu −m8000h −n 702 −N 12 −S 6 −j 2 −d 6 −ss env OMP_NUM_THREADS=6 ./
    ifsMASTER −f h240 −t 600.0 −v ecmwf −e b08 : −cc cpu −n 10 −N 1 −j 1 −d 1 ./xios_server.
    → exe
```

*Code snippet 41: Example of an aprun command running IFS and XIOS*
5 Performance analysis and optimization

After the integration, the next step is to study the computational performance to detect potential bottlenecks and optimize them if possible.

The methodology that we used to analyze the performance and optimize the code is illustrated in Figure 19.

Firstly, we measure the execution time to use it as a reference. Then, using a profiling and/or tracing tool, we detect any possible bottleneck. Following up, we apply the proper optimization technique, and we also verify that the results are correct.

After that, we measure the execution time again, and depending on the time improvement of the optimization, we definitely integrate it or we discard it. If we discard the optimization, we try to find a better optimization, while if we integrate the optimization, we have to choose the next step: if the overall speedup including all the optimization is good enough, we end up the process; otherwise, we start a new iteration by performing a new performance analysis.
Before explaining the optimization techniques that we performed, it is better to give some execution times about the IFS-XIOS integration. It is very important to keep in mind that all the performance numbers that appear in this document are using the IFS configuration explained in later Section 6.2. In summary, we use: the T1279 octahedral reduced Gaussian grid with 137 verticals levels; the forecast has a length of 10 days with a time step of 600 seconds.
and outputs at a frequency of 3 hours. IFS uses 702 MPI processes with 6 OpenMP threads per process, which gives a total of 4212 cores. In addition, it is very important to mention that our development does not execute FullPos, which will be needed in the future to properly output data and perform vertical interpolations that are not supported by XIOS. This part of the work is currently under development, but for analyzing the IFS-XIOS integration from a computational point of view, it was not necessary to have the whole project finished. Therefore, we have to keep in mind that comparing our development against the MF I/O server and the sequential output scheme is not 100% fair, since they execute FullPos.

The total execution time of IFS and XIOS without any optimization takes 7773 seconds for a total of 1440 time steps, being this total execution time the summation of all time steps plus a period of initialization and finalization of the model. To have a reference and changing only the I/O scheme, the sequential output scheme takes 9054 seconds, the MF I/O server takes 7535 seconds, and the same execution disabling the output takes 7356 seconds.

If we focus on the execution of one time step, the execution times are of the order of less than 10 seconds. One of the IFS log files reports the execution time per each time step. Code snippet 42 illustrates a simplified output from time step 311 to 321. The 7th column indicates the individual execution time. Regular time steps last about 4.7 seconds, while one out of six time steps last about 6.7 seconds. The slower ones are in charge of also executing the radiation (it is not computed at each time step due to computational cost reasons).

Furthermore, in Code snippet 43 we provide specific execution times of our developed code and included in the IFS execution. These times are obtained using the GSTATS profiling tool, briefly explained in following Section 5.1. There are three profiled subroutines, SUXIOS_INI, SUXIOS_CTX and CXIOS, which corresponds to our public suxios_ini, suxios_ctx and ifs_xios_send_fields subroutines respectively. Note that suxios_fin does not appear, although we tried to profile it. It could be related to the fact that it finalizes MPI and as a consequence GSTATS is not able to properly finish the profiling.

The time that these three subroutines take is not considerable: initialization takes 0.5 seconds, context setup takes 9.5 seconds, and the NPROMA blocks gather and send of fields take 84.8 seconds. In relative terms, they sum 1.24% of the total execution time (last column).
In Section 5.1, we will briefly explain the tools that we used to study the computational performance.

After that, we will explain the different optimization techniques that we used (5.2, 5.3, 5.4) and the reason why we used them.

Finally, in Section 5.5 we explain three additional optimization techniques that we did not implement due to the following reasons:

- It was not possible to use it.
- It was implemented by the compiler using the proper compilation flag.
- It is enabled by default, but it was interesting to see what happens if not used.

5.1 Tools

We basically used two kind of tools: profiling and tracing. They are explained in the following two sections.

5.1.1 Profiling

Profiling is the analysis of the application’s behavior using information gathered as the program executes in order to determine which parts need to be optimized. To achieve this goal, a profiling tool takes into account aspects such as execution time per subroutine, execution time per line of code (this implies to know which instructions have more cost), which are the dependencies between functions (it builds a tree with calls), the number of times that each function has been called, etc. We can get all this data without looking at the source code or even without having to instrument it.

One popular tool is gprof, which offers simplicity. However, since we are not interested in profiling the entire code, we used the GSTATS timers. IFS has its own low overhead timers called GSTATS. They are really simple to use: declare an identifier of the region that you want to instrument, and add two calls, at the beginning and at the end of the region.

At the end of the execution, GSTATS generates a profile for each instrumented region: number of executed times, the average, the aggregated, the maximum, standard deviation, etc.
5.1.2 Tracing

Tracing is the process of recording event-based performance data along the execution of a program. Using a viewer we can see the behavior of the application in our machine, focusing on hardware counters, communication patterns or memory hierarchy. The tools used to trace the model were Extrae and Paraver, which are open-source and developed by the BSC Performance Tools group [44]:

- Extrae is a package used to instrument the code automatically and/or manually through its API. It generates Paraver trace-files with hardware counters, MPI messages and other information for a post-mortem analysis. It can be downloaded and installed in any HPC facility.
- Paraver is a trace browser that can show a global visual qualitative perspective of the application behavior for later focus on the bottlenecks with a detailed quantitative analysis. The tool allows to create views with any parameter recorded and points to a region of a code, making process of modification easier.

Figure 20 shows an example of a trace, which in this case is a regular time step from an IFS-XIOS execution. A trace has the timeline on the x axis and the MPI processes on the y axis. Along the timeline, some (or many) events happen, which can be related to MPI calls, cache misses, Million Instructions Per Second (MIPS) and many other performance metrics. These events can be chosen with Paraver, but it depends on the configuration of Extrae, i.e., we have to set up in an XML file the events that we want to generate.

The trace of the Figure 20 shows MPI call events, where each color represents an MPI function. Figure 21 illustrates in detail each one of the possible colors, i.e., possible MPI functions. Note that the first color, light blue, it is actually not an MPI function, but it represents computation (outside MPI).

Figure 20: Example of a trace with MPI call events. It shows a complete regular time step of an IFS-XIOS execution. Timeline is on the x axis and MPI processes on the y axis.
Although other metrics have been evaluated too, the figures showed in this document are traces related to MPI call events because it is enough for the performance analyses and optimization techniques that we used.

5.2 Threading with OpenMP

One of the optimization was to parallelize the NPROMA blocks gather using OpenMP threads. As we explained in Section 4.3, the gather was necessary because the requested XIOS arrays have different dimensions with regard to the IFS ones, so it was necessary to build those new arrays.

According to the profiling analysis with GSTATS timers, the gather does not represent too much time. Code snippet 44 shows in detail the gather of the ciwc field. It takes about 5.8 ms, so the optimization would not have a large impact in the execution time for these tests. However, while the OpenMP master thread is performing the gather, the rest of threads are idle, so this operation is actually inefficient. In addition, it could become a bottleneck for future experiments demanding more computational power. Thus, it is better to use all the threads.

<table>
<thead>
<tr>
<th>ROUTINE</th>
<th>CALLS</th>
<th>SUM(s)</th>
<th>AVE(ms)</th>
<th>STDDEV(ms)</th>
<th>MAX(ms)</th>
<th>SUMB(s)</th>
<th>FRAC(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ciwc_GATHER</td>
<td>80</td>
<td>0.5</td>
<td>5.8</td>
<td>0.9</td>
<td>6.5</td>
<td>0.0</td>
<td>0.01</td>
</tr>
</tbody>
</table>

*Code snippet 44: Profiling analysis with GSTATS of a field gather*

We followed the parallelization strategy of Code snippet 45. It contains the gather and send of two fields, specific humidity and temperature, in order to understand how we try to overlap the send of one field with the gather of the next one. This is extended to many other fields using the same strategy, but showing two fields is enough for illustrative purposes.

Note that now, the condition of each IF is a logical variable, since we have performed the call to the xios_field_is_active subroutine and stored the condition before the OpenMP parallel region. This is because XIOS subroutines are not thread-safe.

We start by parallelizing the gather using an OMP DO directive, where the granularity of chunks is an NGPBLKS block. This strategy is beneficial for OpenMP because IFS data arrays were designed to that end. We explained the benefits of this design in Section 3.1.2.
After that, once we have an XIOS-style array, we can send it to XIOS. Only one of the threads will perform the send, while the others will perform the gather of the next field. This overlap is possible because we use the NOWAIT clause in the SINGLE construct.

```fortran
!*OMP PARALLEL PRIVATE(jstglo,icend,ibl,jlev)
! GFL – Specific humidity
IF (q) THEN
!*OMP DO SCHEDULE(DYNAMIC)
DO jstglo = 1, YDGEOMETRY%YRGEM%NGPTOT, YDGEOMETRY%YRDIM%NPROMA
   icend = MIN(YDGEOMETRY%YRDIM%NPROMA,YDGEOMETRY%YRGEM%NGPTOT–jstglo ↦+1)
   ibl = (jstglo–1)/YDGEOMETRY%YRDIM%NPROMA + 1
   DO jlev = 1, YDGEOMETRY%YRDIMV%NFLEVG
      xios_gfl(jstglo:jstglo+icend–1,jlev) = YDFIELDS%YRGFL%GFL(1:icend,jlev,YGFL%YQ%MP,ibl)
   END DO
END DO
!*OMP END DO
!*OMP SINGLE
CALL xios_send_field("q",xios_gfl)
!*OMP END SINGLE NOWAIT
END IF
!
! GMV – Temperature
!*OMP DO SCHEDULE(DYNAMIC)
DO jstglo = 1, YDGEOMETRY%YRGEM%NGPTOT, YDGEOMETRY%YRDIM%NPROMA
   icend = MIN(YDGEOMETRY%YRDIM%NPROMA,YDGEOMETRY%YRGEM%NGPTOT–jstglo ↦+1)
   ibl = (jstglo–1)/YDGEOMETRY%YRDIM%NPROMA + 1
   DO jlev = 1, YDGEOMETRY%YRDIMV%NFLEVG
      xios_gmv(jstglo:jstglo+icend–1,jlev) = YDFIELDS%YRGMV%GMV(1:icend,jlev,YDFIELDS%YRGMV%YT0%MT,ibl)
   END DO
END DO
!*OMP END DO
!*OMP SINGLE
CALL xios_send_field("t",xios_gmv)
!*OMP END SINGLE NOWAIT
END IF
!
! . Gather and send many other fields
!*OMP END PARALLEL
```

This optimization slightly improves the performance as it can be seen in results Section 6.4 (see Figures 27 and 28). The execution time is reduced from 7773 seconds to 7705 seconds. It is important to highlight that depending on the distribution of cores between MPI and OpenMP, we could exploit much more this optimization. For example, if we have a configuration where we use more cores for OpenMP threads and less cores for MPI processes, we would have less subdomains, but much bigger. Thus, the gather with just one thread would be much more significant, because subdomains would have much more data to be gathered.
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Code snippet 46 shows the profiling analysis of the field ciwc with GSTATS timers using the optimization. The average time is reduced from 5.8 ms to 1.2 ms. Note that this is also extensible to all other fields that we need to output.

<table>
<thead>
<tr>
<th>ROUTINE CALLS</th>
<th>SUM(s)</th>
<th>AVE(ms)</th>
<th>STDDEV(ms)</th>
<th>MAX(ms)</th>
<th>SUMB(s)</th>
<th>FRAC(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ciwc_GATHER</td>
<td>80</td>
<td>0.1</td>
<td>1.2</td>
<td>0.2</td>
<td>2.3</td>
<td>0.0</td>
</tr>
</tbody>
</table>

*Code snippet 46: Profiling analysis with GSTATS of an optimized gather*

5.3 Optimized compilation of XIOS

The following optimization was the compilation of XIOS. The compilation of XIOS using the -O1 flag should be improved in order to take advantage of the optimization introduced by the compiler automatically, where the use of -O2 or -O3 are the options recommended by default. However, as we explained in Section 4.4.1 we had a lot of issues to use them.

We were analyzing the performance reports of XIOS and we realized that it was spending a considerably amount of time for just writing data, especially on client side. Code snippet 47 shows that IFS spent about 132 seconds in executing XIOS client code, while Code snippet 48 shows that XIOS servers spent about 3196 seconds in processing events.

We know that enabling post-processing leads to an increase of the XIOS execution time. Depending on the post-processing filter used, this post-processing is performed on client side or on server side. For example, horizontal domain interpolations are performed on client side, while data compression is performed on server side. However, since in our tests there is not post-processing at all, the execution time for XIOS seems to be too much high. For this reason, we thought that it would be interesting to try to improve the XIOS compilation.

As we mentioned in Section 4.4.1 we read that for some old Cray compilers there was a bug using the -O3 flag. However, this bug may have been solved in new versions, so once updated, we tried to use -O3. Interestingly, XIOS compiled and tests successfully passed.
When we tried this new optimized version with IFS, we experienced an improvement in the execution time as it can be seen in results Section 6.4 (see Figures 27 and 28). The execution time is reduced from 7705 seconds to 7629 seconds. Code snippet 49 shows that the execution time on client side is reduced from 132 seconds to 40 seconds. On server side (Code snippet 50), same effect occurs and execution time is reduced from 3196 seconds to 1382 seconds.

Code snippet 49: XIOS report on client side once optimized

Code snippet 50: XIOS report on server side once optimized

This optimization will be especially useful when post-processing is used, since it typically requires a lot of computation.

5.4 Overlapping computation and communication

The last optimization that we implemented was an overlapping between the computation of IFS and the MPI communications to XIOS.

We mentioned that non-output time steps have execution times of about 4.7 seconds (regular time step) and about 6.7 seconds (time step with radiation) as showed in Code snippet 42. However, we realized that in an output time step, there is a slight increase in the execution time of the three following time steps. Code snippet 51 illustrates this effect: time step 324 performs output through XIOS however, the time steps which experience an increase are 325, 326 and 327. This increase actually has many variability among similar output time steps, which means that sometimes is negligible and sometimes is really noticeable.

Nevertheless, in general we can observe this pattern in the following three time steps of an output time step.
In order to understand what was happening, we traced the execution with Extrae. Figure 22 illustrates three time steps, where the first one is performing output at the end. The major part of processes of the trace are the IFS ones, while some processes on the bottom are the XIOS ones.

In this trace, IFS is using the output scheme that we explained in Section 4 and showed in Figure 18. This scheme sequentially executes at the end of an output time step these three steps: update calendar, NPROMA blocks gather and send fields. If it is not an output time step, it only performs the update calendar step.

The trace reveals an interesting point: at the beginning of the next time step after an output time step (second time step in Figure 22), there is an increase in the communications among IFS processes. They are basically performing an MPI_Waitany and an MPI_Alltoallv, but they seem to be delayed according to the time step where this should be done.

The problem is related to the moment where some fields are sent to XIOS. Although they are sent using asynchronous communications to overlap with computation, at the beginning of the time step IFS is not performing computation, but synchronous communication. This leads IFS to wait for XIOS asynchronous communications and its own synchronous communications. IFS communications have to wait because nodes and network are already occupied by XIOS communications.

In addition, at the beginning of the third time step there is also a delay in the MPI_Waitany and MPI_Alltoallv execution, though at the end of the second time step no field is sent at all. So we concluded that when executing the update calendar after an output time step, XIOS performs costly communications that occupy the network resources which cannot be used by IFS at that moment.
Then, knowing that update calendar and send fields steps can potentially collide with IFS communications, we considered to re-order them along the execution time to truly overlap XIOS communication with IFS computation. According to Figure 22, at each time step there is a huge area (light blue) where IFS is only performing computation. This region is in charge of executing the physics of IFS. Ideally, it is a perfect place to perform the XIOS communications.

Our new scheme with re-ordered steps is illustrated in Figure 23: the update calendar step ($C_x$) is prior called at the beginning of the physics in the same time step; the NPROMA blocks gather ($G_x$) is the only step that remains at the end of the same time step; and the send fields step ($S_x$) is delayed to the next time step at the beginning of the physics, called before the update calendar of the next time step ($C_{x+1}$).

Once all time steps are performed, we may need to execute the send fields step ($S_n$) of the last time step, something that it is negligible for the final execution time.
The approach to implement the first scheme was to use just one subroutine called `ifs_xios_send_fields`, which contained the three essential steps: update calendar, NPROMA blocks gather and send fields.

The implementation of the new scheme is not difficult: we split the three steps into three different subroutines:

- **ifs_xios_calendar**: it contains the call to the `xios_update_calendar` subroutine and all the calls to `xios_field_is_active` for each one of the fields to individually store in logical variables if they need to be sent or not.

- **ifs_xios_gather_fields**: depending on the values of the logical variables, this subroutine will perform the NPROMA blocks gather of the fields that need to be sent.

- **ifs_xios_send_fields**: depending on the values of the logical variables, this subroutine will send the fields which are active.

These three public subroutines are placed in the proper places of the IFS code so that the new scheme is fully implemented.

According to the results that are shown in Section 6.4 (see Figures 27 and 28), it is the optimization that gives more improvement in the total execution time: it is reduced from 7629 seconds of the previous optimization to 7507 seconds.

This new scheme improves the execution time of the three time steps that follow an output time step. Code snippet 52 shows this behaviour: time step 324 performs output, but it almost does not affect the execution time of time steps 325, 326 and 327.
If we trace this new scheme, Figure 24 indicates that there is no delay in the \textit{MPI} \texttt{Waitany} and \textit{MPI} \texttt{Alltoallv} operations at the beginning of an IFS time step. It is visible how IFS executes the NPROMA blocks gather at the end of the first time step and the call to send fields at the beginning of the physics in the second time step.

However, it is also important to remark that now there is a delay at the end of time steps two and three, but it is less significant than the delay of the first scheme.

![Code snippet 52: IFS log file with one output time step once optimized](image)

Figure 24: Trace with three IFS time steps once optimized

Figure 25 shows a comparison of the two output schemes. Traces of Figures 22 and 24 are compared using the same elapsed time. It is visible how the three time steps of the trace that uses the optimized output scheme (below) finish earlier than the other three time steps, the ones that use the non-optimized scheme.

![Figure 25: Trace with two IFS time steps once optimized](image)
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5.5 Explored options

We also considered three additional optimization techniques, but as we explained at the beginning of this section, we did not implement them.

5.5.1 Vectorization with SIMD instructions

One of the key points in current x86 machines is the use of vector instructions, also known as Single Instruction, Multiple Data (SIMD) instructions. We thought that they would be really important for performing the NPROMA blocks gather, in order to speed up the whole process.

However, before any kind of implementation, we checked the assembly code to know if compiler was using them. Effectively, Cray compiler was vectorizing the gather, as it is shown in Code snippet 53. At the beginning, it uses a series of _vmovaps_ instructions to move data from memory to vector registers (_xmm_). Then, this data is moved from vector registers to the stack (_rsp_). Finally, once all data is in the stack, Cray compiler calls the _xios_send_field_r8_2d_ subroutine using the _callq_ instruction.
5.5.2 Memory affinity

The majority of [HPC] machines use [Non-Uniform Memory Access (NUMA)] nodes. This type of computers are characterized by having more than one processor, which share memory without uniform access time. This means that the whole memory is distributed across the processors, where each processor locally owns a part of the memory. However, processors are able to access other processors’ local memory, but more slowly.

Cray XC40 nodes have two [NUMA] sockets (or processors), each one with 18 cores and its own
local memory. When running a parallel job with aprun, it is possible to specify if sockets are allowed to allocate memory from each other or not. The -ss flag indicates that a core or a thread can allocate only the memory local to its assigned NUMA socket.

In theory, using this flag it should give more computational performance, since slow crossed memory allocations between sockets are not allowed. This type of tests are known as memory affinity. Our idea was to test and potentially use the -ss optimal option. However, by default IFS uses the -ss flag to run, so this optimization was already in use.

Nevertheless, we thought that it would be useful to see if -ss was certainly optimizing the run of IFS so we repeated the experiment removing this flag. We observed that the total execution time dramatically increased by 20%. Thus, we proved that the -ss flag to achieve NUMA affinity is mandatory for this kind of applications.

5.5.3 Derived MPI Datatypes

We have seen that IFS-style arrays and XIOS-style arrays are different, so we needed to perform an NPROMA blocks gather before sending fields to XIOS. We occurred that using derived MPI Datatypes could be possible to avoid the gather.

The idea of using derived MPI Datatypes [45] is to describe the layout of data structures in memory. This means that they are used to describe how data is stored: stride between elements, how many contiguous elements per block and the block count. It is possible to describe vectors, structs, etc. Thus, the NPROMA blocks gather would not be needed.

Nevertheless, we were not able to use this technique because we do not directly work with MPI subroutines when we send data to XIOS. The only way is by calling the xios_send_field subroutine, which is from the XIOS API. Although this cannot be done, it could be suggested to the XIOS developers to add it as a new feature in the future.
6 Evaluation

In this section we will evaluate the results of the development and the optimization techniques. In Section 6.1 we overview the HPC platform. In Section 6.2 we explain the IFS configuration that we used to perform the benchmark tests. Then, in Section 6.3 we show the two metrics that are used to measure the computational performance. Finally, in Section 6.4 we explain all the different tests that we performed.

6.1 HPC platform and compilation environment

Performance evaluation was conducted on the ECMWF’s High Performance Computing Facility (HPCF) [46], which is a Cray system that has two identical Cray XC40 clusters. It has a peak performance of 8499 teraflops.

Each Cray XC40 cluster has 3610 compute nodes running the Cray CLE 5.2 UP04 operating system. Nodes are made of 128 GB of memory and two Intel E5-2695v4 ”Broadwell” processors, each with 18 cores. It is possible to activate hyper-threading, offering a total of 72 threads.

Cores have available three levels of cache, L1, L2 and L3, having 64 KiB, 256 KiB and 45 MiB (shared) of memory respectively. They operate at a clock frequency of 2.1 GHz.

The cluster uses the Aries™ Interconnect network technology, which implements a ”dragonfly” topology. In addition, Lustre is used as the parallel file system.

The Cray Developer Toolkit (CDT) version 16.04 and the Cray Compiling Environment (CCE) version 8.4.6 were used to compile both IFS and XIOS. They were built with Cray MPICH 7.3.3.

6.2 IFS configuration

It is really important to choose a proper IFS configuration to benchmark our development, since we want to be as close as possible to real experiments. Thus, we could have an exact idea of the benefits of using XIOS.

We will compare and discuss in Section 6.4 the different output schemes: sequential output, MF I/O server, XIOS server (different optimized versions) and also no output, to know which is the maximum achievable speedup.

Table 1 shows a summary of the basic IFS parameters that are used in the configuration to compare all the output schemes. We use the octahedral reduced Gaussian grid in high resolution, which is the same that ECMWF uses for its operational forecast. We also use 702 MPI processes with 6 OpenMP threads for each one, which gives a total of 4212 processing elements. This distribution of processing elements is recommended by ECMWF because it offers a good balance between performance and efficiency. Moreover, we do not change neither the number of MPI processes nor the number of OpenMP threads because it would be a scalability study, but it is not part of this work. In order to evaluate the computational performance of our development, it is preferable to focus on the number of processes of XIOS and the output size.

Furthermore, as we have previously mentioned in Section 5, unlike the MPI I/O server and the sequential output scheme, our IFS XIOS integration does not execute FullPos. This part of the project is under development, but for analyzing the benefits of using XIOS from a computational
point of view is enough using the development presented in this document. Therefore, it is important to keep in mind that the comparison is not fully fair.

We almost used all the default values that prepIFS sets up when a new experiment is created. We just changed a few parameters, especially the ones related to performance and output.

<table>
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<tr>
<th>Basic IFS parameters</th>
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<tr>
<td>Grid type</td>
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<td>Vertical resolution</td>
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<td>Time step</td>
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</tr>
<tr>
<td>3D prognostic fields</td>
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</tr>
<tr>
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<td>No</td>
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<tr>
<td>OpenMP threads per forecast task</td>
<td>6</td>
</tr>
<tr>
<td>Hyperthreading</td>
<td>Yes</td>
</tr>
</tbody>
</table>

*Table 1: Summary of the IFS configuration*

The parameters that change between output schemes are the following:

- Sequential output: we do not need additional processes for I/O servers. We do not want to write data using the Fields Data Base (FDB).

- MF I/O server: we have to specify the number of dedicated processes for I/O, which the recommended amount by ECMWF is 18. We do want to write data using FDB.

- XIOS: we have to specify the number of dedicated processes for I/O, which the recommended amount is analyzed in later Section 6.4.1. We cannot use FDB, so we disable it.

The output is characterized by being quite large in size: we are outputting data at a frequency of 3 hours, while in the operational forecast is at 6 hours. Thus, we are able to stress the schemes and prove that XIOS works well. In any case, there are real configurations where a frequency of 3 hours could be used.

We output the following 12 3D prognostic fields:

- U component of wind
- V component of wind
- Temperature
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- Divergence
- Vorticity
- Specific humidity
- Cloud fraction
- Ozone
- Specific rain water content (rain)
- Specific snow water content (snow)
- Specific cloud liquid water content (liquid water)
- Specific cloud ice water content (ice water)

In addition, we also output this 2D prognostic field:

- Surface pressure

It is important to be aware of the output size of this particular configuration because it is considerably large and it is directly related to the time used for the output process. However, it depends on the data format:

- **GRIB** format: files’ size is about 770 GB.
- **NetCDF** format: files’ size is about 3.2 TB.

This difference in size between both formats is due to compression. While **GRIB** files are compressed, **NetCDF** files do not have any kind of compression. However, we could enable the **NetCDF** files compression in the **XIOS** configuration, which would not affect the overall performance because it is done on server side. But, we do not do that because we are only evaluating **I/O** aspects.

Finally, in order to evaluate the results, for each case we make the average of three executions. It is true that when measuring **I/O** the sampling should be bigger, about 10 runs, because the variability considerably grows. Nevertheless, we do not do it because of two reasons. The first one is because the **I/O** variability is not reflected in the total execution time, since **I/O** servers have enough time to "buffer" it. The second reason is because we are running huge experiments, so we cannot afford wasting such amount of computing hours.

### 6.3 Metrics

In order to compare and evaluate the two **IFS** output schemes against the different optimization versions of our development, we will use two basic metrics in computer sciences: execution time and speedup.
We measured the execution time as the elapsed time from the beginning of the IFS-XIOS execution to completion.

The speedup measures the relative performance between two systems solving the same problem. Systems can be of any type: hardware systems, software systems, etc. In our case, we compare different output schemes for IFS which always uses the configuration explained in previous Section 6.2.

We define the speedup as:

\[ S = \frac{T_{\text{baseline}}}{T_{\text{optimized}}} \]

Where:

- \( S \) is the speedup.
- \( T_{\text{baseline}} \) is the execution time of the case that we take as a reference.
- \( T_{\text{optimized}} \) is the execution time of cases that we want to know the relative improvement against the reference case.

6.4 Results

In this section we will explain all the different tests that we performed to measure our development. First of all, in Section 6.4.1 we will show how the optimal number of XIOS servers change in function of the output size. After that, in Section 6.4.2 we perform a comparison between all the output schemes that we previously mentioned. In Section 6.4.3 we perform the same type of comparison, but adding the time needed to convert GRIB to netCDF files. Finally, in Section 6.4.4 we perform a comparison test by adding to non-XIOS schemes the same number of cores and nodes that XIOS uses.

Figures use some abbreviations for XIOS: v1, v2, v3 and v4. They include the different optimization versions which are in increasing order. For example, v3 includes v1 and v2. The list is the following:

- **XIOS** v1: non-optimized version. It is only the IFS-XIOS integration development. It corresponds to Section 4.
- **XIOS** v2: it includes the optimization of the NPROMA blocks gather parallelization with OpenMP threads. It corresponds to Section 5.2.
- **XIOS** v3: it includes the optimized compilation of XIOS. It corresponds to Section 5.3.
- **XIOS** v4: it includes the overlapping computation and communication optimization. It corresponds to Section 5.4.
6.4.1 Optimal number of XIOS servers

In this test we execute [IFS] with [XIOS] under the same conditions, only changing the number of fields to be output. The purpose is to find the minimum number of [XIOS] servers needed that do not increase the total execution time depending on the output size. This means that we are finding the optimal number of [XIOS] servers for different number of fields to be output.

[XIOS] servers consume a considerable amount of memory to maintain the value of variables needed in some post-processing operations. This means that depending on the volume of the set of fields, more or less memory is allocated by [XIOS] servers. This is solved using the memory of more or less nodes of the supercomputer. Additionally, the more servers are working at the same time, the more fields can be processed in parallel, avoiding a bottleneck during the output process. In Figure 26 we use three different output sizes to find the corresponding optimal numbers: 12 3D fields, 6 3D fields and 1 3D field. When we output 12 fields, we need 10 servers, with 6 fields we need 5 servers and with 1 field we just need 1 server. It is visible that the number of needed servers scales in function of the output size.

![Optimal number of XIOS servers](image)

There is an important point to mention. For this test we are placing one server per node, because if we try to fit those number of cores in less nodes, the execution crashes due to lack of memory. Thus, we can also say that Figure 26 shows the number of nodes needed in function of the output size. However, note that if we increase the number of [XIOS] servers per node, but keeping the needed amount of nodes, the execution successfully finishes spending the same elapsed time.

Therefore, this figure suggests that [XIOS] is memory sensitive. As we have explained, [XIOS]...
computes a lot of memory because it accumulates data on server side just in case it is needed to apply time integration filters, i.e., post-processing over a period of time.

6.4.2 Comparison test between different output schemes

In this test we compare the different output schemes that we have available: IFS sequential output, MF I/O server, different IFS-XIOS optimized versions and IFS with no output. The fact that I/O schemes are compared with the no output case is to know the maximum achievable speedup because it does not have I/O overhead at all. It is important to remind that: all executions are done using 702 MPI processes with 6 OpenMP threads each one; and our integrated IFS-XIOS framework does not use FullPos, which is used in the MF I/O server and in the sequential output, so comparison is not fully fair. In addition, we are going to show results that take into account the whole execution of IFS-XIOS, this is, from the beginning to the end of the execution.

Figure 27 shows the execution time of the different schemes. The first column is the sequential scheme, which takes 9054 seconds. It is considerably slower than the rest of schemes. The second one, the MF I/O server, takes 7519 seconds. It is a good time compared to no output, which takes 7356 seconds.

The next four columns are the different XIOS versions. It is visible how each optimization improves the previous one. The XIOS v1 is only the IFS-XIOS integration, which takes 7773 seconds. It is faster than the sequential output, but slower than the MF I/O server. The next one, XIOS v2, parallelizes the NPROMA blocks gather using OpenMP threads. This optimization avoids that only the OpenMP master thread works while the rest of threads are idle. It takes 7705 seconds, which is a reduction of 68 seconds regarding the XIOS v1.

After that, the XIOS v3 uses the optimized compilation of XIOS. It is very important to reduce both client and server execution time, especially if it is necessary to run post-processing, which typically implies a lot of computation. The execution time is improved by 76 seconds with regard to XIOS v2, taking 7629 seconds.

Finally, the most optimized version, XIOS v4, re-orders the three necessary steps to perform output: update calendar, NPROMA blocks gather and send fields. The send fields step is performed in the IFS physics computation, taking advantage that there are not communications at all. Thus, it is possible to achieve better overlapping between XIOS communication and IFS computation. The execution time takes 7507 seconds, 122 seconds faster than the previous XIOS v3. It is also slightly faster than the MF I/O server (12 seconds), and much more than the sequential output (1547 seconds). In addition, it is only 151 seconds slower than no output.
Figure 27: Comparison of the execution time of different output schemes

Figure 28 shows the speedup of the same previous comparison. The maximum achievable speedup is 1.231x, i.e., a 23.1% faster than the sequential output. Both MF I/O server and XIOS v4 execution time improvements are 20.4% and 20.6% respectively.

Comparing the sequential output (9054 seconds) and the no output tests (7356 seconds), it is easy to check that the output process requires an average of 1698 seconds, which represents a 23% more of the execution time of IFS. However, thanks to the most optimized version of the IFS XIOS integration, this time is reduced from 1698 to 151 seconds, which represents only a 2% of the execution time of IFS achieving that almost all the output process is hidden with the IFS computation.
6.4.3 Comparison test adding the GRIB to netCDF post-processing

This test is the same as the previous one (Section 6.4.2), but with the additional cost of converting GRIB files to netCDF files. This cost is only added in sequential output and MF I/O server schemes, since they are the ones that write in GRIB format. The format conversion used is the same that EC-Earth uses for its experiments. In particular, GRIB files are serially converted to netCDF files, this is, one file at a time. Note that we do not include the no output case, which does not have neither GRIB format nor netCDF format, since it does not generate data.

This test is useful to know the potential benefit of using XIOS to avoid the costly GRIB to netCDF post-processing.

Figure 29 shows a comparison between the different output schemes, where the time needed to convert GRIB files to netCDF files is taken into account. The post-processing time needed for the conversion is really huge, about 13680 seconds (3.8 hours). It is much more slower than the simulation itself.

The execution time of the sequential output scheme is increased from 9054 seconds to 22734 seconds, and for the MF I/O server from 7519 seconds to 21199 seconds. It represents an increase of 151.1% and 181.9% respectively.
Figure 29: Comparison of the execution time of different output schemes, where both sequential output and MF I/O server have the additional cost of converting GRIB to netCDF files.

Figure 30 shows the speedup of the same previous comparison. It is considerable the speedups that XIOS achieves in all its optimization versions: 2.92x in v1, 2.95x in v2, 2.97x in v3 and 3.02x in v4.

It is really evident that if we take into account both model and post-processing tasks, the benefit of using XIOS compared to the current I/O schemes is huge.
6.4.4 Comparison test with additional computational resources

Finally, we perform a comparison with additional computational resources between the sequential output scheme and the most optimized version of IFS-XIOS, i.e., XIOS v4. The idea is to use in both cases, firstly, the same amount of cores, and secondly, the same amount of nodes. Thus, we guarantee that we compare cases using the same amount of computational resources. Furthermore, at the end there is the same comparison test, but also taking into account the GRIB to netCDF conversion cost for the sequential output scheme.

It is important to keep in mind that each Cray XC40 node has 36 physical cores, so using hyperthreading 72 threads. In our case, we are running 12 MPI processes per node, where each one has 6 OpenMP threads. Thus, we are completely fulfilling nodes.

We always take as a reference the optimal amount of XIOS servers (and nodes) that will be added to the other cases. Since we are outputting 12 3D fields, we use 10 XIOS servers spread along 10 nodes. Therefore, those additional resources used by XIOS are added to the sequential output scheme as follows: if using the same amount of cores, we use 10 additional cores to run IFS (702 + 10 = 712). On the contrary, if using the same amount of nodes, we use 120 additional cores to run IFS (702 + 120 = 822).

Note that we do not include the MF I/O server. This is because of two reasons. Firstly, if we want to use 10 cores as dedicated I/O processes, the execution crashes since there are not enough resources. The minimum amount is 18, the default value. And secondly, if we want to use 10 nodes to place I/O processes, we cannot set it up because I/O processes are internally distributed by IFS. It is not possible to choose the geometry of the parallel job through aprun.
Figure 31 shows the execution time comparison of the two evaluated schemes. The first case is the sequential output. It has a considerable improvement using the same amount of nodes, where execution time is reduced from 9054 seconds to 8282 seconds. The second case is the XIOS v4, where the execution time is always kept to 7507 seconds, because it is the reference case regarding the amount of resources.

![Figure 31: Comparison test with additional computational resources of the execution time of two different output schemes. In blue they are using the original amount of resources, in orange they are using the same amount of cores, and in yellow they are using the same amount of nodes.](image)

From previous Figure 31 we compute the speedup of Figure 32. It illustrates how the XIOS v4 scheme achieves a lower speedup when the sequential output scheme uses more computational resources.
Figure 32: Speedup of XIOS v4 against the sequential output scheme with additional computational resources. In blue the original amount of resources, in orange the same amount of cores, and in yellow the same amount of nodes.

Figure 33 illustrates the same execution time comparison of previous Figure 31 but also taking into account the execution time of converting GRIB to netCDF files. In this scenario, the benefit of using more computational resources for the sequential output scheme is less significant, because the increase added by the format conversion is really big.
Figure 33: Execution time comparison test with additional computational resources and GRIB to netCDF conversion cost of two different output schemes. In blue they are using the original amount of resources, in orange they are using the same amount of cores, and in yellow they are using the same amount of nodes.

Figure 34 shows the speedup computed from the previous figure, where it is evident that XIOS offers a considerable improvement with regard to the sequential output scheme. The speedup of XIOS v4 is 3.00x using the same amount of cores and 2.92x using the same amount of nodes.
Figure 34: Speedup of XIOS v4 against the sequential output scheme with additional computational resources and GRIB to netCDF conversion cost. In blue the original amount of resources, in orange the same amount of cores, and in yellow the same amount of nodes.
7 Conclusions and future work

7.1 Conclusions

In this project we have integrated XIOS into IFS with a twofold objective: reduce the total execution time of the model and reduce the workflow’s critical path by using online post-processing. In addition, we had a secondary objective which was to increase the usability of IFS by using an easier output configuration file compared to the current approach.

In order to achieve the objectives, we have presented a development which is characterized by being very easy to use. IFS initializes, sets up and finalizes XIOS through three subroutines. Then, we have designed a scheme to write data at the end of each output time step. This output scheme is made of three steps: update calendar, NPROMA blocks gather and send fields. This first development already improved the execution time with regard to the sequential output scheme, despite of not being optimized. The execution time takes 7773 seconds, while the sequential output takes 9054 seconds, which adds a 23% of time to the execution of IFS. However, note that unlike the MF I/O server and the sequential output scheme, the integrated IFS-XIOS version does not execute FullPos, which is needed to output and post-process correct data. This is currently an ongoing task, but for analyzing the benefits of using XIOS from a computational point of view, it is enough with the work presented in this document.

After that, we have applied several optimization techniques to improve the bottlenecks that we detected in the performance analysis.

In the first optimization we have used OpenMP threads to parallelize the NPROMA blocks gather. Although it does not give a large improvement, it is important to not have threads in idle, while only works the master thread. This optimization would be especially beneficial when having larger subdomains with less MPI processes and more OpenMP threads per MPI process to ensure a good scalability for any configuration of future developments. The execution time is reduced 68 seconds with regard to the first development.

The second optimization is an improvement in the XIOS compilation. After sorting out all the issues that we were having, we achieved to compile XIOS using the -O3 optimization flag. Not surprisingly, there is an improvement in the execution time, which is reduced by 76 seconds regarding the previous optimization. This proves that it is important to compile external libraries using the best optimization flags of the compiler. Although this could seem trivial, a non-optimized compilation of external libraries could be a bottleneck for the scalability of a model.

The last optimization is the most beneficial in terms of computational performance. We have used more sophisticated tools such as Extrae and Paraver to find what could be improved. We have realized that there was not a real overlap in the computation and communication of IFS and XIOS respectively, so we have designed a new output scheme to re-arrange the steps and achieve an improved overlap. In this new scheme, the three steps of the first output scheme are split and placed along the IFS time step. This optimization takes 7507 seconds, improving the execution time by 122 seconds. Thanks to this last optimization, it is even slightly faster than the MF I/O server, which takes 7519 seconds. Furthermore, it is important to mention that this optimization, which also includes the previous ones, is only 151 seconds slower than IFS without outputting data at all, which represents only a 2% of the execution time of IFS. Within 151 seconds, IFS outputs through XIOS 3.2 TB of data. This optimization proves that the use of asynchronous communications to overlap with computation is sometimes not
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enough. In some cases, an additional study is needed to detect in which areas computation and communication can be effectively overlapped, taking into account other communication stages along the complete execution.

We have also considered other optimization techniques such as vectorization with \textbf{SIMD} instructions, memory affinity and derived \textbf{MPI} Datatypes. However, we have not implemented them due to different reasons: the vectorization of the code that we would have implemented, it is already done by the compiler; the memory affinity improvement is already in use in \textbf{IFS} (however, we have proved that its use is very important for the performance of the execution); and the derived \textbf{MPI} Datatypes are not possible to be used, since data is sent to \textbf{XIOS} through Fortran subroutines.

When we also take into account the post-processing task to compare performance results, the benefit of using \textbf{XIOS} becomes enormous. In both sequential output and \textbf{MPI} I/O server schemes is necessary to convert \textbf{GRIB} files to \textbf{netCDF} files, which are serially converted. It takes 13680 seconds (3.8 hours), which increases the execution time of the sequential scheme up to 22734 seconds and 21199 seconds in the \textbf{MPI} I/O server. Our most optimized version is a 202\% faster than the sequential output scheme and a 182\% faster than the \textbf{MPI} I/O server. Thus, we have implemented an scalable development that will address the I/O issue that is limiting the computational performance of \textbf{IFS} within EC-Earth. It represents a first step in EC-Earth that will considerably reduce the total execution time of large experiments, as well as it will increase the efficiency. This does not only imply to save thousands of computing hours, but also storage space because we will only store processed data ready to be used, instead of a huge amount of temporary raw data.

We have actively worked, discussed and collaborated with two European institutions, \textbf{ECMWF} and \textbf{NLeSC} to succeed in the completion of this project. \textbf{ECMWF} is interested in this work because they will distribute OpenIFS with \textbf{XIOS} as an optional I/O scheme.

Furthermore, the EC-Earth community is very interested in this work, because it is one of the key developments that need to be implemented for a future major release of EC-Earth. Some of the institutions involved in this community are \textbf{BSC} [\textbf{KNMI}] [\textbf{SMHI}] etc.

7.2 Future work

In order to keep improving this project, there are several tasks that will be gradually and sequentially implemented in the future by different institutions:

- As we have previously mentioned, it is needed to use FullPos to output correct data and apply vertical interpolations that are not currently supported by \textbf{XIOS}. However, we have to ensure a user-friendly output configuration file because choosing an inappropriate mechanism to set up both FullPos and \textbf{XIOS} can lead to a usability loss. This task is currently ongoing.

- All the work done for \textbf{IFS} will be ported to OpenIFS.

- At that time, \textbf{XIOS} will be already integrated with OpenIFS and \textbf{NEMO}, so it will be necessary to adapt EC-Earth components to generate on-line diagnostics through \textbf{XIOS}.
Acronyms

**ADIOS** ADaptable I/O System.
**AOGCM** Atmosphere-Ocean Global Circulation Model.
**API** Application Programming Interface.
**BSC** Barcelona Supercomputing Center.
**CAM** Community Atmosphere Model.
**CDI-pio** CDI with parallel I/O.
**CDT** Cray Developer Toolkit.
**CeCILL** CEA CNRS INRIA Logiciel Libre.
**CFIO** Climate Fast Input/Output.
**CMIP** Coupled Model Intercomparison Project.
**CMOR** Climate Model Output Rewriter.
**CPU** Central Processing Unit.
**ECMWF** European Centre for Medium-Range Weather Forecasts.
**ESM** Earth System Modelling.
**FDB** Fields Data Base.
**GPU** Graphic Processing Unit.
**GRIB** General Regularly-distributed Information in Binary form.
**H2020** Horizon 2020.
**HDF** Hierarchical Data Format.
**HPC** High Performance Computing.
**I/O** Input/Output.
**IFS** Integrated Forecasting System.
**ILP** Instruction Level Parallelism.
**IPSL** Institute Pierre Simon Laplace.
**KNMI** Koninklijk Nederlands Meteorologisch Instituut.
**MF** Météo-France.
MIPS  Million Instructions Per Second.
MPI  Message Passing Interface.
MPMD  Multiple-Program Multiple-Data.
NEMO  Nucleus for European Modelling of the Ocean.
netCDF  Network Common Data Format.
NLeSC  Netherlands eScience Center.
NUMA  Non-Uniform Memory Access.
NWP  Numerical Weather Prediction.
OpenMP  Open Multi-Processing.
OST  Object Storage Target.
PIO  Parallel I/O library.
POSIX  Portable Operating System Interface.
PRIMAVERA  PRocess-based climate sIMulation: AdVances in high-resolution modelling and European climate Risk Assessment.
SIMD  Single Instruction, Multiple Data.
SMHI  Swedish Meteorological and Hydrological Institute.
XIOS  XML Input/Output Server.
XML  Extensible Markup Language.
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