NEMO 4.0 performance: how to identify and reduce unnecessary communications

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Abstract

A non-intrusive instrumentation of the NEMO code and the development of a simplified configuration (called BENCH) brought information about MPI communications cost and structure. It helped us to identify the most appropriate incremental developments that model needs to enhance its scalability. We prioritised the reduction of extra calculations and communications required at the North Polar folding, the grouping of boundary exchanges and the replacement of global communications by alternative algorithms. Appreciable speed up (x2 in some cases) is measured. Scalability limit is pushed below a size of 7x7 grid points per sub-domain, showing that the limitation of the North Polar folding solution can be compared with the supposed icosahedral grid one. We consider that scalability is not the major well of future performance gain, neither horizontal resolution increase, whereas potentiality of extra developments accelerating cache access (horizontal domain tiling and single precision computations) is favourably evaluated. Taking note of the limited technological gain between the two 6-month old and 4-year-6-month old machines we operated, in order to avoid future net decrease in computing performance, we recommend from now to limit new expensive code developments to what technology and engineering are able to sustain.
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The cost of MPI communications is clearly identified as the most important factor of scalability limitation of the NEMO model (e.g. Tintó Prims et al. 2018 [1]). Any new scalability improvement in this model requires to carefully identify the place of the most costly MPI exchanges in the code, to characterise their utility, before being able to propose a strategy of algorithm modification, a communication reduction or a simple removal.

1. Light code instrumentation dedicated to calculation / communication ratio measurement

In NEMO, MPI communications are wrapped in a small number of high level routines co-located in a single FORTRAN file (lib_mpp.F90). This code structure strongly facilitates a quick and non intrusive instrumentation for a study dedicated to the MPI communication structure. Basically, the NEMO communication library provides functionalities for horizontal halo exchange and global averages or min/max operation between all horizontal subdomains. Every halo exchange of any variable of the code necessarily needs to be done through the generic "ROUTINE_LNK". The global operations (sum, min/max) are also located in a few number of routines, that can be also gathered following the same generic strategy (one routine fits all argument kinds – integer/float/double, 2D/3D arrays - With very little code changes focused at these positions, it is possible to identify and characterise, for further modification, the whole MPI communication pattern. This instrumentation does not replace existing external solutions like “extrae-paraver” described in Tintó Prims et al. 2018. The amount of information collected by our solution is much smaller, and the possibilities of analysis reduced, but we are able to deliver without any external library, without additional computing cost or additional post-processing, a synthetic information for any kind of machine.

Because of some initial coding assumptions of the existing internal timing library routine (timing.F90), we prefer not to modify and re-use its functionalities, but to slightly instrument the ROUTINE_LNK generic interface with MPI_Wtime calls. Following the strategy already deployed to estimate the coupled system load imbalance [2], a counter of the time spent in MPI routines, also called “waiting time”, is incremented at any call of a MPI send or receive operation in one hand, and at any MPI collective operations on the other hand. Symmetrically, a second counter is filled outside these two kind of operation. Only two informations per core are collected in two real scalar variables: the cumulated time spent to send/receive/gather or (mostly) wait MPI messages and the complementary period of other operations (named “calculations”).

The strategic position of our instrumentation allows to not only estimate the time spent in the communication routines but also to identify their calling routines and the size and kind (2D/3D or pointers of 3D arrays) of exchanged halo arrays. The calling of any lbc_lnk or MPI-collective subroutine is modified to communicate, via sub-routine argument, the name of the calling routine. A collection of these names is performed during the first time step of the simulation. The second time step (which could be the second coupling time step when sea-ice or bio-geo-chemistry model is enabled) produces the listing of the calling tree and the size of the exchanges in a dedicated output file called communication_report.txt. We produce below the content of such analysis for an ocean/sea-ice simulation:
Communication pattern report (second oce+sbc time step):

Exchanged halos :  773
3D Exchanged halos :  93
Multi arrays exchanged halos : 163
   from which 3D :  22
Array max size :      20250

lbc_lnk called
- 1 times by subroutine usrdef_sbc
- 1 times by subroutine icedyn_rhg_ev
- 2 times by subroutine icedyn_rdgft
- 483 times by subroutine icedyn_adv_umx
- 1 times by subroutine icedth
- 1 times by subroutine icecor
- 2 times by subroutine iceupdate
- 2 times by subroutine zdfphy
- 4 times by subroutine ldfs1p
- 1 times by subroutine divhor
- 2 times by subroutine domvv1
- 1 times by subroutine dynkeg
- 1 times by subroutine dynvor
- 177 times by subroutine dynspg_ts
- 1 times by subroutine divhor
- 2 times by subroutine domvv1
- 1 times by subroutine traqsr
- 1 times by subroutine trabbc
- 6 times by subroutine traadv_fct
- 2 times by subroutine trannx
- 1 times by subroutine dynx
- 3 times by subroutine domvv1

Global communications :  1
- 1 times by subroutine icedyn_adv_umx

During the second time step, it is possible to substitute the model operations (step routine) by the MPI exchanges of the realistic sequence of halo communications. This skeleton of realistic exchanges is useful to avoid processing the whole simulation and help to save computational resources during tests, as recently experienced by Zheng and Marguinaud, 2018 [3]. In our case, the skeleton can be automatically set up for any NEMO configuration (resolution, namelist, CPP keys) during the first time step and applied during the rest of the simulation. The sequence of the communications can be modified or the communication block even switched off on purpose to estimate the effect of such modification to the model performance.

At the simulation end, we also produce (if ln_timing option is activated in the namctl namelist) a separated counting, per MPI process, of the total duration of (i) halo exchanges for 2D/3D and simple/multiple arrays, (ii) global communication to produce global sum/min/max, (iii) all other MPI independent operations, called ‘computing time’ and (iv) the whole simulation, excluding first and last time step. This analysis is output together with the existing information related to timing in NEMO (timing.output file). We produce below the first lines of this analysis for the same ocean ice simulation:

Report on time spent on waiting MPI messages
 total timing measured between nit000+1 and nitend-1
 warning: includes restarts writing time if output before nitend ...

<table>
<thead>
<tr>
<th>Operation</th>
<th>Time</th>
<th>MPI Rank</th>
</tr>
</thead>
<tbody>
<tr>
<td>Computing time</td>
<td>7.674150</td>
<td>1</td>
</tr>
<tr>
<td>Waiting lbc_lnk</td>
<td>10.760121 (58.4 %)</td>
<td>1</td>
</tr>
<tr>
<td>Waiting global</td>
<td>0.000163 (0.0 %)</td>
<td>1</td>
</tr>
<tr>
<td>Total time</td>
<td>18.437015 (SYPD: 533.887)</td>
<td>1</td>
</tr>
<tr>
<td>Computing time</td>
<td>5.714904</td>
<td>2</td>
</tr>
<tr>
<td>Waiting lbc_lnk</td>
<td>12.721502 (69.0 %)</td>
<td>2</td>
</tr>
</tbody>
</table>
2. **BENCH: a simplified NEMO configuration for realistic performance analysis**

The NEMO framework proposes various configurations based on the core ocean model OPA, global (ORCA grid) or regional, with various horizontal and vertical resolutions. To this core element, other modules can be added: sea-ice (e.g. SI3) or bio-geo-chemistry (e.g. TOP-PISCES). To deliver a synthetic message about the computing speed of the NEMO model, a comprehensive performance analysis requires to investigate several resolutions / module sets. On the other hand, such benchmarking exercise must be kept simple, since it is supposed to be used by people with basic knowledge of model handling and physical oceanography.

This is why a new NEMO configuration was specifically developed for this study, and included in the reference code with hope to simplify future benchmark activities related to computing performance. This new configuration, called BENCH, is available in the tests/BENCH directory of the last 4.0 code version. It is based on the existing GYRE configuration, with modifications that make the measurements easier and increase its robustness.

GYRE horizontal grid is a simple cuboid, which dimensions can be easily changed by namelist parameter. The lack of bathymetry ensures a relative numerical stability that allows various numerical tests. All NEMO numerical schemes can be used. A selection of namelist parameters is done to confer to GYRE the numerical properties that correspond to the chosen resolution.

GYRE does not include bathymetry (nor continents). However, in NEMO, calculations are processed at each grid point (then mask is applied to take into account land grid point if any). Consequently, the amount of calculations is the same with or without bathymetry and we assume that this amount in GYRE (and BENCH) configurations is realistic. The only function that our benchmark configuration cannot test is the removal of sub-domain entirely covered by land points. We assume that the communication pattern would not be affected by this characteristic.

The East-West periodical conditions and the North pole folding cannot be activated with GYRE. These characteristics of the set of ORCA global grids (a consequence of the global periodicity and of the bi-polar shape of the NEMO northern hemisphere representation) have a big impact on performance and this is why the new BENCH configuration includes these periodicity conditions, unlike GYRE. A simple change of the \texttt{nn_perio} namelist parameter gives to the BENCH grid the ORCA periodic characteristics and reproduces the same communication pattern between subdomains located on the Northern most part of the grid.

To be sure that these additional communications, compared to GYRE, are not affecting too much the numerical stability of the model, we ensure in BENCH a bigger stability with constant and small external forcing. External forcing is not read in files, which excludes evaluation of input reading procedures from our performance analysis. The initial vertical gradient of the density is minimised, but not removed, to let adjustment processes be launched and keep the associated amount of calculations at a realistic value. In addition, the prognostic variables are
initialised with different values at each grid point, to ensure that reproducibility tests can be effectively processed. Again, for stability reasons, these differences are minimised.

SI3 sea-ice or TOP bio-geo-chemistry can be activated in BENCH. An ORCA-like representation of SI3 processes in the BENCH configuration implies a careful initial temperature/salinity definition, in such a way that 1/5 of the domain would be covered, during the whole simulation, with sea ice.

3. Experimental set-up

3.1. Methodology

A set of short simulations are produced with the Météo-France BULL supercomputer “beaufix2". The total number of time step is adjusted to average enough information and to minimise the exercise cost. Considering that initialisation and finalisation phases are excluded of our measurement, the result dependency with respect to the time step number is supposed to be small. At the opposite, it is important to minimise any disk access during the time loop. All NETCDF Input/Output are disabled (no CPP key_iomput) and ASCII output are reduced to the minimum.

To evaluate the benefit of our future modification, we will take measurement of a set of simulations with various decompositions. Let's emphasise here that performance is strongly linked to the decomposition choice. This is why our BENCH implementation comes with a new calculation of optimum decomposition values. We define the decomposition set which helped us to determine scalability of our configurations, relying on the main assumptions that (i) subdomain perimeter has to be minimised and (ii) the division rest of the total number of subdomain by the number of core per node (CpN) has to be zero or close to CpN.

Part (i) of this algorithm was also implemented in the last official model release for all kind of configurations, and enriched to take into account the land-only subdomains that are automatically switched off.

3.2. BENCH performance analysis

Applying our measurement tool to the BENCH simplified configuration, a better picture of scalability variations with resolution or machine is now available and it is even possible to clarify their origin. On our current supercomputers, a normal usage of resources limits scalability investigations to 1 or 0.25 degrees configurations. As shown in Figure 1-left, “waiting time” (MPI communications + calculation load imbalance, pink line) is the limiting factor for BENCH-1 speed (1 degree resolution, BENCH configuration, red line). The scalability limit is reached when the length of the MPI subdomain sides falls below 10x10 grid points. In addition, the difference between the fastest and the slowest calculations between MPI processes (orange line) gives an idea of the total calculation spread, i.e. load imbalance, between subdomains. Its contribution to the total waiting time is significant.

1 https://www.top500.org/system/178075
Figure 1, left: Scalability of BENCH-1 degree resolution configuration, considering total time to solution, in second per simulated year, for several side length of MPI subdomains. Red line shows the total time to solution, pink line shows the time spent to wait halo arrays (MPI messages), orange shows the difference between slowest and fastest total calculations (excluding waiting time, see Figure 1, right) amongst MPI processes. Figure 1, right: total time to solution but excluding time spent to wait halo arrays, for each MPI process, same unit, same model configuration than left figure. The 6 measures are done with 6 different domain decompositions (domain side length in legend).

Figure 2, left: Scalability of BENCH-1 configuration, considering total time to solution and time spent waiting MPI messages (halos), same experiment than Figure 1-left (red and pink lines). In a second experiment, communications are switched off. Total time to solution of this second experiment in shown in dark blue, with corresponding calculations spread in light blue. Figure 3, right. In a third experiment, realistic communications are performed every two time step only. Total time to solution of this second experiment in shown in dark green, with corresponding waiting time in light green.

Figure 1-right illustrates the imbalance of calculation time between the MPI processes. Processes of the last two nodes shows extra calculations (arrays copies) of polar folding. They are the origin of most of the total imbalance as plotted in Figure1-left (orange line). Additional analysis (not shown) reveals that it is also at the origin of most of the extra waiting time (MPI
communication time). These analysis lead to prioritise our current model performance enhancement work on polar folding effect reduction.

We also took benefit of the possibility to easily modulate the amount of communication to sharply estimate the MPI exchange effect on scalability. Figures 2 compare time to solution and calculation spread of our reference experiment with two others. In the first one, all halo exchanges were switched off. In the second one, exchanges were performed every two time steps only.

The first experiment quantifies the dominant effect of communication on performance at every position of the scalability curve. As far as our measurement can be lead, the remaining load imbalance cannot limit the scalability. The second experiment estimates the maximum benefit of a possible implementation of a communication-reduced algorithm such as the so called double halo (a two point wide halo is transmitted every two time step).

4. Enhancement of North folding exchanges

The so-called North folding exchanges are made necessary by the particular tri-polar structure of the ORCA global grids (see [4]). The uppermost subdomains located along the North pole exchange their north-south halos between them. In this North folding area, the grid points of the last line are duplicated (symmetry of type 5 or 6, currently used for ORCA1 discretisation). With symmetry of type 3 or 4, used for ORCA025 or ORCA12 discretisation, the grid points of the last but one line are symmetrically duplicated and the grid points of the last line are duplicated on the last but two line. In addition, depending on the MPI decomposition, a subdomain must have to be connected with two subdomains. These particularities, combined with the C-Arakawa structure of variables, add extra arrays copies and extra communications to the halo exchanges of the North fold area. Up to four lines, instead of one, are communicated in the original NEMO code.

Simulation limited to realistic halo exchanges (skeleton) reveals that the size of exchanges data has an impact on both calculation (mainly arrays copies) and MPI communications. A fine study of North-South / East-West communication pattern reveals that only one or two lines (depending on the grid type and the chose symmetry) are really necessary to update all grid points in the North fold area. We call “moderate Reduced North folding (RN)” exchanges the implementation that limits the exchanges to 1 to 2 lines. This implementation does not affect the results.

A more aggressive reduction to one single communicated line is also implemented (“full RN” exchanges). This can theoretically change the results because of the possible heterogeneity of values that should be identical in restart variables on one hand, and the “triad” algorithm, used e. g. in the total vorticity trend computation – energy and enstrophy conserving scheme - that leads to reverse the operation order in the last 3 northernmost lines of the grid and introduce a small difference in calculation results of two North-fold-symmetric subdomains, on the other hand.
The two implementations are tested in the 4.0 version of the code (specific branch). The experimental design is detailed at the previous chapter. The configuration is limited to the ocean model only.

Effect on restitution time for each MPI process (subdomain) is plotted in Figure 3. We limit the number of MPI ranks plotted to the very last 3 or 4 nodes, considering that North folding effect only affects subdomains of the last 2 or 3 nodes. The reduction in elapsed time in sensible at both BENCH-1 and BENCH-025 resolution. Resolution BENCH-12 is not shown due to the lack of resources necessary to reach a sufficient communication time/calculation time ratio, but the effect is supposed to be identical to the BENCH-025 one. From figures 4, we can deduce that subdomain sizes of 5x9 (BENCH-1) and 17x24 (BENCH-025) are needed to estimate the effect of our implementation.

The computing time is reduced by our modification, in same proportion with both moderate and full RN strategy: the additional time needed by North folding computations (mainly array...
copies) cannot be estimated by comparison to a reference simulation led without any North folding exchanges.

The reduction of MPI communication time is more difficult to estimate but can be roughly deduced from waiting time measurements. At BENCH-1 resolution, it is hard to evaluate the extra effect of full RN. At the opposite, a further reduction of waiting time is observed at BENCH-025 resolution. This probably explains the extra speed up observed on Figure 4 (right) from 35.2 SYPD (moderate RN) to 42.5 SYPD (full RN). We suppose that the reduction of MPI messages size is more efficient with 17x24 subdomain size (BENCH-025) than with 7x9 (BENCH-1).

In both cases, a non negligible speed up (~x 1.4) is obtained thanks to our strategy. The effect is sensible at every decomposition but brings the better gains at scalability limit. Due to non reproducibility of results in particular configuration (AGRIFF), the moderate RN is eventually included in the official release.

We deduce from Figure 3 that the North fold effect is no more the bottleneck that limits the model scalability. At considered horizontal resolutions, the tri-polar grid does not introduce more substantial extra load imbalance and could be compared to icosahedral grids.

![Figure 4: Scalability of BENCH-1 (left) and BENCH-025 (right) configurations, considering total time to solution and time spent waiting MPI messages (halos). Performance of the initial implementation (red/orange) is compared with the new algorithm of North fold halo exchange in moderate (blue/cyan) and full (green/lime) modes. Same configuration with no exchange at North fold (black) is also presented.](image)

5. **Other enhancements of OPA scalability**

Other modifications are applied following two strategies: to group halo exchanges and to remove unnecessary global communication.

During vorticity (dynvar.F90) calculation, halo communication are performed together for all vertical levels (3D fields communication). Similarly, during tracer advection (traadv_fct.F90), the 3 spatial components are grouped during communication.
Regarding global communications, a new procedure is implemented to check, at every time step, that model prognostic quantities are bounded to realistic values. In this procedure, a local (instead of global) check is performed and the decision to stop the simulation, after a local output of the standard `output.abort.nc` file, is only taken by the MPI process affected by the failure diagnostic. After the output of pre-abort prognostic variables, this single process ends the simulation, stopping all MPI processes of the `MPI_COMM_WORLD` communicator. This simultaneously ensures (i) a clean simulation stop, even in coupled mode, (ii) the output of local quantities relevant to identify the crash origin (local `output.abort.nc` file) and (iii) the removal of a global communication performed at each time step.

In addition, the number of unnecessary writing in disk is reduced:
- The `ln_ctl` logical (unused in production mode) must be true to output the `run.stat` files
- The `time.step` file is now written only by the first subdomain

6. **Enhancement of SI3 scalability**

An analysis of the location and call frequency of halo exchanges and global communications in the code (see figure 5, outer donut) reveals the major contribution of three routines of the sea ice module (advection, rheology and thermodynamics). Only a change of the rheology numerical scheme (iterative procedure needed to solve the momentum equation) could limit the halo exchange frequency. Further analysis are necessary to this change. Modification of the two other parts of the code are presented below.

![Figure 5: Occurrence of MPI-communication calls in an ocean/sea-ice configuration before (outer donut) and after (inner donut) optimisation](image)

Advection trend calculation (TVD, `ice_dyn_adv_umx` routine), which includes 2D halo exchange, is called for every layer (and every category). The reduction of communication
volume is here quite straightforward. We modify the `adv_umx` subroutine call: a single call is processed for all layers/categories and 3D (instead of 2D) arrays are exchanged during halo update.

In ice thermodynamics, the computing of time evolution of snow and sea-ice temperature profiles (ice_thd_zdf_BL99 routine) needed a separate MPI communicator, dedicated to communications only between processor cover by sea-ice. Creation/removal of the communicator was performed at each time step and for each ice category. A global maximum, to evaluate the maximum value of a converging quantity, was the single function of the communicator. This global information ensured a number of convergence iterations that was independent of the MPI decomposition. We can keep this independence by calculating the convergence criterion and stop the convergence calculation for each grid point, independently of the others. This avoids the final global communication.

The remaining global communications necessary to check, in sea-ice advection scheme, whether time splitting is necessary or not to avoid instability, are delayed by one time step, in order to perform one time step computations in the meantime. This strategy implies the coding of the checkpoint/restart procedure to preserve the “restartability” of the model. Finally, the extra cost of these “delayed communications” is practically reduced to zero.

As show in Figure 5 (inner donut), all global communications and 1/3 of the halo exchanges are removed from ocean and ice code. As seen in Figure 6, this clearly push away the scalability limit. For clearer results, 10 categories and 5 layers are set in SI3 (instead of the usual 5 and 2 values), which is called at every ocean time step (instead of every 3 time step usually). The overall NEMO benefit depends on the SI3/OPA relative weight, which depends on the number of SI3 categories/layers, but the performance enhancement of the single SI3 model is mainly independent of these parameters.
7. Enhancement in TOP-PISCES model

Relying on the tools described above, the comprehensive study of global and boundary communications in the TOP tracer module and biogeochemistry model PISCES reveals the intensive use of global communications in 4 routines (see figure 7, outer donut).

![Figure 7: Occurrence of MPI-communication calls in TOP/PISCES modules before (outer donut) and after (inner donut) optimisation](image)

A removal of small negative quantities (numerical errors produced during tracer advection) is performed in the \texttt{trcrad} routine. A global mean is necessary to compensate at every grid point the quantities created by the negative values removal. A new system of compensation is implemented: assuming that the negative values are always small, they can be spread on a 3x3 grid point local stencil, that can always be found (possibly on halo grid points) close to the faulty values. This strategy leads to the removal of a global communication (but needs an additional boundary exchange).

Following the same strategy than for ice and ocean, global checks performed in \texttt{trcstp} are disabled in standard mode, and preserved only when the \texttt{ln_ctl} logical is set. Similarly, global means, only needed for diagnostics, are called under IF statements: this removes the communication call in \texttt{pk4zflx} routine. The \texttt{p4zsink} routine was entirely rewritten by PISCES developers, that removes the last two global communications of the module. Finally, the remaining halo communications in \texttt{trcsbc} can be grouped and reduced to one.

8. Performance

We evaluate the benefit induced by the whole set of modifications between version 3.6 stable, revision 10164 and version 4 of the code. Two set of scalability measurements are performed on beaufix2 Météo-France and irene\textsuperscript{2} TGCC supercomputers, using the BENCH configuration (resolution 1 and 0.25 degrees). This output-less configuration allows to isolate performance of

2 https://www.top500.org/system/179411
calculation and communications, without any interference of disk accesses. In parallel, performance of the widely used ORCA2_ICE_PISCES configuration is also evaluated to see how the impact of XIOS writing and disk accesses affect the results.

Figure 8: Scalability of BENCH-1 configuration, before and after optimisation described in this document, ocean stand alone (red and orange), including sea-ice model with 5 categories and 2 layers, called every 2 ocean time steps (dark and light blue) or including TOP-PISCES biogeochemistry, 24 tracers, called every time step (dark and light green), on Météo-France (left) and CEA-TGCC (right) supercomputers.

In any cases, even though our processes are not sharing resources with other users jobs on our exclusive computing nodes, results are largely dependant of the per-user bandwidth availability, i.e. the machine load. Measurement error margin can be greater than 100% at extreme scalability (>10,000 cores), when MPI communication time is a big contributor to the total restitution time. Scalability plots presented in Figure 8 and 9 are established with the fastest runs we could measure during the period of machine biggest availability (Christmas holidays). This strongly mitigates the enhancements we got in any case (between x 1.15 to more than x 2 faster) but proves the ability of NEMO “polar-folding” communication scheme to lead 1 degree ocean-only global simulations closer to the 1,000 SYPD target and to use ¼ degree ocean-only global model under near perfect scalability conditions on the present day machines.

Comparison between 1 and 0.25 degrees configurations shows that scalability limit is only reached at low resolution. From that, we deduce that only low resolution tests help us to investigate the scalability limit of the model. The only extra information that BENCH-025 carries is that our present supercomputers can only process BENCH-025, BENCH-12 and BENCH-beyond calculations below the scalability limit and other strategy of time to solution reduction must be found elsewhere. Another consequence of this result is that the increase of resolution is not a solution for a better use of the supercomputer resources: at 10,000 cores limit, scalability will be nearly perfect for any global configuration above 0.25 degrees. At the opposite, at 10,000 cores limit, the BENCH-12 subdomains, larger than the BENCH-025 ones, will saturate the node memory caches and downgrade the performance.
An uncertainty on measurement of the same kind than with BENCH (reinforced by disk accessibility constraints) is observed with the full I/O low resolution configuration of the model (see Fig 10). However, our communication reduction work is still sensible in this case.

9. Conclusion and perspectives

The modifications we made lead to comfortable gains only if the model is used under very special conditions: perfect condition of bandwidth use (nearly empty machine), no disk accesses, model resolution fitted to reach scalability limit in present day supercomputers. At low
resolution, maximum speed is reached when the maximum per user number of resources are requested. Considering that large requests are always delayed for longer periods than smaller jobs, the actual performance (measured in ASYPD) of our enhanced model is probably less alluring. As expressed in [5], a larger set of production runs would be necessary to better estimate this difference. But it is clear that our enhancements are sensible mainly when production conditions are unfavourable. At higher resolution, the scalability limit cannot be reached, which strongly limit the impact of our modifications. To summarise, our enhancements guarantees that the scalability limits of the tri-polar grided NEMO model can theoretically be compared to other models/grids but, in practice, they lead to a modest 20 to 30% gain in production conditions. This is why we consider that further optimisation should not focus on scalability limits but investigate the enhancement conditions well below this limit.

Figure 11: Restitution (elapsed) time needed by the BENCH model to perform calculations per grid point (arbitrary unit), for various sizes (square side) of the horizontal subdomain (including halo), allocated on each core of the 4x5/6x8 core node of beaufix2/irene supercomputers, in ocean-only, 75 vertical levels (upper-left plot), ocean+SI3 ice module called every 2 time steps (upper-right plot), ocean+bio-geo-chemistry module including 24 tracers (lower-left plot) and all configurations together (lower-right plot).

In particular, a comparison of models (particularly ocean only) scalability with ideal scaling in Figures 8 and 9 shows a super-scalability in the first part (below 500 to 1000 cores) of the plot. A complementary analysis is produced to better identify its origin. In one node of our two
machines, we vary the size of each square subdomain attributed to each of the node cores (40 on beaufix2, 48 on irene) and we calculate the time spent to process calculations only (communications are switched off) for a fixed number of grid points. Results from Figure 11 prove that it is easier for a node, whatever NEMO configuration is (ocean only, ocean+ice or ocean+bio-geo-chemistry), to perform calculations with 10x10 than 100x100 horizontal subdomains. This probably reveals the sensibility of the model to the cache effect of the machine node. From this, we can infer that a judicious tiling of NEMO computations (the splitting of the sub-domain in smaller subdomain and execution within a loop, instrumented or not to add further OpenMP/OpenACC parallelism) could bring nearly a new factor two to the performance at low resource positions of the scalability plot at low global resolution (ORCA1) and any position for high resolutions. This work is complementary to and cumulative with the precision reduction already explored by [6].

We can also notice from our results that the maximum gain our implementation brings is comparable, or even larger, to the gain delivered by the machine upgrade (irene was made available 4 years after beaufix2), which shows the necessity of such algorithm adjustments to compensate the contemporary Moor's law slowing down. It also makes clearer that such gain is now obtained with much less facility than one decade ago and, considering that sources of additional gain in the code are fewer and fewer, supercomputing should no more be considered as a cornucopia and strategies to limit new costly development in the code planned.

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