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Citation for published version:

Digital Object Identifier (DOI):
10.1007/11549468_83

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Peer reviewed version

Published In:
Euro-Par 2005 Parallel Processing

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Flexible Skeletal Programming with eSkel

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Abstract. We present an overview of eSkel, a library for skeletal parallel programming. eSkel aims to maximise the conceptual flexibility afforded by its component skeletons and to facilitate dynamic selection of skeleton compositions. We present simple examples which illustrate these properties, and discuss the implementation challenges which the model poses.

1 Introduction

The skeletal approach to parallel programming is well documented in the research literature (see [1–4] for surveys). It observes that many parallel algorithms can be characterised and classified by their adherence to one or more of a number of generic patterns of computation and interaction. For instance, a variety of applications in image and signal processing are naturally expressed as process pipelines, with parallelism both between pipeline stages, and within each stage by replication and/or more traditional data parallelism [5].

Skeletal programming proposes that such patterns be abstracted and provided as a programmer’s toolkit, with specifications which transcend architectural variations but implementations which recognise these to enhance performance. This level of abstraction makes it easier for the disciplined programmer to experiment with a variety of parallel structurings for a given application, enabling a clean separation between structural aspects and the application specific details. Meanwhile, the explicit structural information provided by skeletons enables static and dynamic optimisations of the underlying implementation.

In [2] we presented a number of observations and proposals designed to facilitate the popularisation of the skeletal approach, and outlined the first prototype of eSkel, a library of skeleton operations for C and MPI. In the light of subsequent experiences we isolated and described two key concepts, nesting mode and interaction mode, which capture important choices in the design of any skeletal programming framework [6]. This caused us to substantially refine and partly redesign eSkel. While these and other concepts embedded in eSkel may be found in various combinations in previous skeletal systems, we believe that this paper discusses the first attempt to enumerate and compose them systematically and explicitly within a single framework.

This paper describes current work on the latest instantiation of eSkel. We describe the ways in which key concepts have been incorporated into the programming model and API in order to maximise the flexibility with which the
programmer can shape and reshape skeletal programs, and on the substantial implementation challenges which this flexibility creates.

2 Dynamic Selection of Skeletons and Modes

cSkel is designed to provide the skeletal parallel programmer with flexibility in a number of ways. As in the initial prototype [2], a skeleton abstracts the pattern of interactions between a number of component activities, each of which may be internally parallel, and which may choose to invoke further skeletons. The skeletons provided each encompass the ability to specify the use of either localised or distributed data (the data spread), and the choice of either explicit or implicit interaction mode [6] to trigger interactions between the various activities. The aspects of the interface which capture these choices have been rationalised. Guided by the concept of nesting mode [6], the programmer can choose to incorporate both transient and persistent skeleton calls within the same skeleton nest. Crucially, all of these decisions, as well as the choice of overall skeleton structure for each phase of a program, are made dynamically, and can therefore be based, where appropriate, on information gathered during execution (for example, data values or sizes). In summary the significant improvements over cSkel’s first version [2] are as follows.

- The definition of ‘families’ of related skeletons, capturing distinctions in interaction style, have been replaced by the explicit selection of interaction mode. For example, there is now only one pipeline skeleton, which can be parameterized to behave as either of its predecessors.
- All skeletons in the original cSkel were (implicitly) in transient mode. There were no persistent nestings. Both transient and persistent modes are now available (by setting the corresponding parameter).
- The data model has been revised and extended to incorporate the concept of molecules, enabling the expression of skeletons (such as HaloSwap) in which several distinct pieces of data are exchanged at each interaction.

2.1 cSkel interface

The current specification of cSkel defines five skeletons, each with flexibility in the dimensions discussed above. We will focus on the two of these, Pipeline and Deal, which have so far been implemented in the current prototype, since in combination these can illustrate all the points we wish to make. The reader is referred to the cSkel homepage [7] for discussion of the Farm, HaloSwap and Butterfly skeletons.

The Pipeline skeleton abstracts classical pipeline parallelism. It allows an indexed set of activities (called “stages”) to be chained together and applied to a sequence of inputs. Data is passed from a stage to its successor following the natural ordering on stage indices. The user can choose between an implicit pipeline, in which the transfer of the data is done automatically, constraining
the stages to produce one result for each input, or an explicit mode, in which a stage can produce arbitrarily many results without necessarily consuming data (for example, a generator or filter stage). In the latter case, the data transfer is controlled by the programmer, by calls to the generic interaction functions Give and Take.

The Deal skeleton is similar to a traditional farm, but with tasks distributed strictly cyclically to workers. Thus it is most appropriately used when the workload associated with individual tasks is expected to be homogeneous. This skeleton is useful nested in a pipeline, in order to internally replicate a stage. Deal semantics require the ordering of outputs from the skeleton to match that of the corresponding inputs, irrespective of the internal speed of the workers.

The price for this flexibility is paid in the complexity of the API, where each skeleton call must specify its choices. In addition to the issues already discussed, the pragmatic decision to borrow and build upon MPI's data model with its already substantial parameter set, leads to rather long parameter lists. However, when understood as a set of groups, each addressing orthogonal issues, the parameters become more conceptually manageable. We now present these groupings informally. Their formal use will be illustrated in Section 2.2.

One set of parameters deals with the called skeleton's data buffers. As with any MPI collective operation, there are distributed input and output buffers, each requiring the standard MPI pointer, type, length definition. These, together with the enclosing MPI communicator, capture the call's interface to the rest of the program. A second set of parameters deals with the call's internal structure and interfaces. For example, in a pipeline, we must describe the number of stages and the types and data modes on their interfaces, and the allocation of processes from the calling group to the distinct stages (captured by borrowing MPI's colouring parameter mechanism to describe the required sub-groups). Finally, a group of parameters describe the details of the skeleton's various activities (stages in a Pipeline, workers in a Deal) with a choice of interaction mode and a pointer to a function for each.

In the interests of regularity, we have chosen a single generic interface for all functions which contain code for skeleton activities. The essence of activities is that they interact in predefined patterns, via the skeleton infrastructure, with each other and with skeleton calling programs. Each such interaction may involve different numbers of data atoms, according to the semantics of the skeleton. For example, “getting the next task” in a Deal skeleton involves a single atom, whereas “updating the halo” in a HaloSwap skeleton will involve two atoms for each neighbour (one arriving, one leaving). The eSkel_molecule_t type collects atoms involved in an interaction into a single sequence. Activity functions both accept and return a single item, of type eSkel_molecule_t *. The specification of each skeleton defines its detailed molecule usage.

2.2 Examples

We present two variants of a toy program in order to concisely illustrate the ease with which the eSkel programmer can describe and revise the parallel structure
of an application. The first version illustrates the use of persistent nesting and implicit interactions. The second uses transient nesting and explicit interactions.

**Example 1: persistent nesting and implicit interactions**

This program calls a Pipeline skeleton, in which the second stage contains a persistently nested Deal. Some of the activities (pipeline stages and deal workers) are assigned several processes, and process data with “global spread” (i.e. the data is distributed across such processes and accessed in SPMD style).

The constant declarations define the number of pipeline stages (STAGES), the number of workers (DEALWORKERS) in the deal, and the number and size of input items (INPUTNB, INPUTSZ). When run with 8 processes, the input to this pipeline consists of a total of \((8^*4^*5)\) integers, which are treated as 4 (INPUTNB) inputs to the pipeline, each of which is constructed from a 5 (INPUTSZ) integer contribution per process. The first stage in the pipeline has been assigned two processes. Thus, since we are in “global spread” mode, each pipeline input will be fed to the two first stage processes as a block of \((8^*5)/2\) integers each.

The main function (lines 31-57) makes the pipeline call. It defines all the pipeline parameters and assigns processes to the stages. Lines 34-39 describe the structure of the pipeline: all inter-stage data transfers use global spread (PGLOBAL) and involve sequence of integers. All activities except the second stage use “implicit” mode (IMPL) transfers (in other words, handled implicitly by the skeleton). The second stage has “devolved” mode (DEV), indicating that this will be inherited from a persistently nested skeleton (the deal in this case). All stages will execute the SimpleStage activity, except the second, which executes DealStage. Lines 42-43 create some (distributed) input data. Lines 45-48 compute an assignment of processes to stages. Line 50 creates the (distributed) output buffer. The pipeline call itself is on lines 52-54, and is passed the various parameters as created, together with details of the input and output buffers in the style of a conventional MPI collective operation.

Function SimpleStage, on lines 6-12, describes the actions of the first, third and fourth stages. Since implicit interaction mode is used, the function takes an eSkel molecule as a parameter and returns a molecule. Here we simply add 1000 to each integer in the block and return the modified molecule. In contrast, the second stage is described by DealStage. It computes an assignment of processes (from the stage) to workers in the internal deal, then calls the Deal skeleton on lines 26-28. The data mode is stream (STRM) indicating that the data are streamed into the deal from the parent skeleton. The input and output buffer related parameters are therefore redundant (all the NULLs and 0s). Deal workers run the function DealWorker. In our simple example this is identical to SimpleStage. We leave the duplication to emphasise that these functions can be programmed independently, if appropriate.

Figure 2 illustrates the requested assignment of processes to stages and workers, and the different communicators involved. This will help to clarify the discussion of eSkel implementation in Section 3. We have a 4-stage pipeline where the second stage is a deal with two workers, of which one has two processes. Px
```c
#define STAGES 4   // Number of pipeline stages
#define DEALERS 2  // Number of workers in the deal
#define INPUTS 4   // Input multiplicity
#define OUTPUTS 5  // Size of each input datum, per process

eSkel_molecule_t * SimpleStage (eSkel_molecule_t *thingy) {
    int i;
    for (i=0; i<thingy->len[0]; i++) {
        ((int *) thingy->data[0][i]) += 1000;
    }
    return thingy;
}

eSkel_molecule_t * DealWorker (eSkel_molecule_t *thingy) {
    int i;
    for (i=0; i<thingy->len[0]; i++) {
        ((int *) thingy->data[0][i]) += 1000;
    }
    return thingy;
}

void DealStage (void) {
    int workercol, outcol;
    if ((myrank() < 1)) workercol = 0; else workercol = 1;
    Deal (DEALERS, INPUTS, DealWorker, workercol, STAGE,
        NULL, 0, 0, SGLOBAL, MPI_ANY, MPI_ANY, 0, NULL);
    Deal (DEALERS, INPUTS, DealStage, workercol, STAGE,
        NULL, 0, 1, SGLOBAL, MPI_ANY, MPI_ANY, 0, NULL);
    Deal (DEALERS, INPUTS, DealStage, workercol, STAGE,
        NULL, 0, 2, SGLOBAL, MPI_ANY, MPI_ANY, 0, NULL);
    Deal (DEALERS, INPUTS, DealStage, workercol, STAGE,
        NULL, 0, 3, SGLOBAL, MPI_ANY, MPI_ANY, 0, NULL);
}

int main (int argc, char *argv[]) {
    int i, j, p, next, outcol, mystagemum, inputs, results;
    int inodes[STAGES+1] = {SGLOBAL, SGLOBAL, SGLOBAL, SGLOBAL, SGLOBAL};
    int mystagemum[STAGES+1] = {MPI_ANY, MPI_ANY, MPI_ANY, MPI_ANY, MPI_ANY};
    int *inputs[INPUTS];
    int *results[INPUTS];
    MPI_Init(&argc, &argv); SkellInit();
    inputs = (int *) malloc (sizeof(int)*INPUTS*INPUTS);
    results = (int *) malloc (sizeof(int)*INPUTS*INPUTS); // Create output buffer
    Pipeline (STAGES,   inodes, stage, mystagemum,   inputs, types,
        void * inputs, INPUTS, 0, results, INPUTS, &outcol,
        INPUTS, INPUTS, mycomm());
    MPI_Finalize(); return 0;
}
```

Fig. 1. Example 1
represents process $x$ ($0 \leq x \leq 7$), as indexed in the original MPI_COMM_WORLD. $S_i$ represents stage $i$ ($0 \leq i \leq 3$), and $W_j$ represents worker $j$ ($0 \leq j \leq 1$).

**Example 2: transient nesting and explicit interactions**

Our second example demonstrates the use transient nesting, explicit interactions and local data spread, and also illustrates the ease with which the programmer can experiment with the high level structure of an application. Working from the program of example 1, the programmer decides to

- turn the second stage (formerly the Deal) into a simple single process stage;
- reallocate the two spare processes from the second stage to the first stage;
- adapt the first stage (which now has four processes) so that every second input is processed by a dynamically created four stage pipeline, with one process per stage. These are transiently nested pipelines, in contrast to the persistently nested Deal of example 1. Other inputs are treated as in the original simple stages.

The structural changes at the outer level are captured in lines of 3, 61, 63, 64, 70 and 71 of Figure 2. Line numbers in parentheses are those of the corresponding lines in Figure 2. The rest of main is unchanged, and is omitted here. The behaviour of the new first stage, and its sub-stages are described by the new functions StrangeStage and SubStage. StrangeStage creates and calls the nested pipeline for every second input (controlled by line 44). The nested sub-stages, called with explicit (EXPL) interaction mode (line 35), use cSkel functions Give and Take to control interactions with neighbouring stages (lines 21 and 26).

### 3 Implementation challenges

We now consider the challenges raised by cSkel's implementation, focusing particularly on the call tree, which is built dynamically, and allows processes to find communication partners. We also discuss the management of the input and output buffers.
```c
// Number of stages in the nested pipeline

void SubStage (void) {
  int i, **lem;
  eSkal_molecule_t *tempitem;
  len = (int **) malloc (sizeof (int *));
  temptime = (eSkal_molecule_t *) malloc (sizeof (eSkal_molecule_t));
  while (temptime->data = Take(len)) {
    temptime->len = *len;
    for (i=0;i<temptime->len[0];i++) {
      ((int *) temptime->data[0])[i] += 250;
    }
    Give(temptime->data, temptime->len);
  }
  eSkal_molecule_t *StrangeStage (eSkal_molecule_t *thingy) {
    static int const=0;
    int i, outxml;
    spread_t spread[STAGES+1] = {SPLOCAL, SPLOCAL, SPLOCAL, SPLOCAL, SPLOCAL};
    MPIDataType types[STAGES+1] = {MPI_INT, MPI_INT, MPI_INT, MPI_INT, MPI_INT};
    Inode_t inodes[STAGES] = {EPL, EPL, EPL, EPL};
    eSkal_molecule_t *(*stages[STAGES]) (eSkal_molecule_t *) =
      { (eSkal_molecule_t *) (* (eSkal_molecule_t *)) SubStage,
        (eSkal_molecule_t *) (* (eSkal_molecule_t *)) SubStage,
        (eSkal_molecule_t *) (* (eSkal_molecule_t *)) SubStage,
        (eSkal_molecule_t *) (* (eSkal_molecule_t *)) SubStage};
    if (count % 2) { /* handle it directly */
      for (i=0;i<thingy->len[0];i++) {
        ((int *) thingy->data[0])[i] += 1000;
      }
    } else { /* handle with a transient pipeline */
      Pipeline (STAGES, inodes, stages, nyrank(), BWF, spreads, types,
        thingy->data[0], i, thingy->len[0], thingy->data[0], i, outxml,
        thingy->len[0], nycom());
    }
    return thingy;
  }

  inodes[0] = {MPI, MPI, MPI, MPI};
  return thingy;
}
```

**Fig. 3. Example 2 - changes from example 1**
3.1 Building the call tree dynamically

In order to correctly implement the communications which have been abstracted by the skeleton calls, it is necessary for the processes assigned to the various activities to understand their position within the overall skeleton nest. Since the structure of the nest emerges dynamically, and independently within distinct branches, this investigation can only be performed dynamically, at the point at which no further changes to the structure are possible. In eSkel this occurs when each activity calls its first explicit (or implicit) interaction. Building the call tree requires collaborative communications between all processes in the nest.

![Call tree diagrams]

**Fig. 4.** Call trees for the two examples

The root of the tree corresponds to the main skeleton call. The children describe the different activities, and if there are some persistently nested skeletons, they appear in the tree. The transiently nested structure is not built in the main tree. The sub-tree will be built dynamically when the transient skeleton call is performed. Fig. 4 represents the call tree for the examples introduced in the previous section. The tree of the nested pipeline of Fig 4b is created at each call of this pipeline.

3.2 Finding communication partners

Communication partners are found by traversing the call tree. For example, in Fig. 4a, we discover that Wo1 takes data from S0 and gives data to S2. Interactions with a neighbouring deal require different partners at each interaction (because of the cyclic nature of interactions in that skeleton). In our example, when S0 performs a Give, it needs to give data alternately to Wo0 and Wo1. To ensure this cyclic distribution we keep track of the number of Give and Take calls performed. This allows us to find the appropriate partner.
Particular attention must be paid to boundary cases. When S0 performs a Take, it cannot find a communication partner in the tree. It needs instead to take the data from the input buffer to the pipeline call. Similarly, S3’s Give must store data in the output buffer. We detail the management of these buffers below.

3.3 Managing the input and output buffers

The input buffer must be created collaboratively by all the processes, since each of them potentially has some of the data at the point of the skeleton call. It is created just after the call tree has been built, during the first interaction of each activity. At that time, the processes which have to take data from the input buffer (processes of S0 in our pipeline examples) collect all the data through group communication, and build the input buffer.

The output buffer is created by the processes which write results into the buffer (processes of S3 in the first example). After completion of the skeleton nest, the results are distributed among all the processes.

More care must be taken when the group of processes dealing with the input or output buffer are part of a deal. For example, if the nested deal was in S0 instead of S1, both Wo0 and Wo1 need to take data from the input buffer while maintaining the cyclic order. To address this issue, we dynamically create a thread which distributes the data to the workers. A symmetrical approach is taken for the output buffer.

3.4 Context of use

The eSkel library operates within the context of a running MPI program which has already created its fixed number of processes. Moreover, a fully thread safe version of MPI (in the sense of MPI_THREAD_MULTIPLE) must be used. We are developing eSkel using the Los Alamos Message Passing Interface LA-MPI [8] and the Sun HPC Cluster Tools [9].

4 Future Work

A full implementation of eSkel is in progress, with the current status reported on the project web pages [7]. After completion of the initial prototype, work will focus on development of a set of demonstrator applications, internal optimisations and expansion of the skeleton set. We will also consider ways in which the API can be simplified, for example by moving to a more powerfully descriptive host language such as Java, as and when the Java-MPI combination gains more widespread practical acceptance. Current work on eSkel is conducted under the umbrella of the Enhance project [10], which exploits the modelling power of the Performance Evaluation Process Algebra (PEPA) [11] to predict the performance of skeletaly structured Grid applications, with a view to assisting in their scheduling and re-scheduling in the presence of the dynamically heterogeneous performance and availability characteristics of Grid computing platforms.
Acknowledgements

The authors are supported by the Enhance project (“Enhancing the Performance
Predictability of Grid Applications with Patterns and Process Algebras”) funded
by the Engineering and Physical Sciences Research Council under grant number
GR/S21717/01.

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