The PARSEC Benchmark Suite: Characterization and Architectural Implications

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ABSTRACT

This paper presents and characterizes the Princeton Application Repository for Shared-Memory Computers (PAR-SEC), a benchmark suite for studies of Chip-Multiprocessors (CMPs). Previous available benchmarks for multiprocessors have focused on high-performance computing applications and used a limited number of synchronization methods. PARSEC includes emerging applications in recognition, mining and synthesis (RMS) as well as systems applications which mimic large-scale multi-threaded commercial programs. Our characterization shows that the benchmark suite is diverse in working set, locality, data sharing, synchronization, and off-chip traffic. The benchmark suite has been made available to the public.

Categories and Subject Descriptors

D.0 [Software]: [benchmark suite]

General Terms

Performance, Measurement, Experimentation

Keywords

benchmark suite, performance measurement, multi-threading, shared-memory computers

1. INTRODUCTION

Benchmarking is the quantitative foundation of computer architecture research. Program execution time is the only accurate way to measure performance[18]. Without a program selection that provides a representative snapshot of the target application space, performance results can be misleading and no valid conclusions may be drawn from an experiment outcome. CMPs require a disruptive change in order for programs to benefit from their full potential. Future applications will have to be parallel, but due to the lack of a representative, multi-threaded benchmark suite most scientists were forced to fall back to existing benchmarks. This usually meant the use of older High-Performance Computing (HPC) workloads, smaller suites with only few programs or unparallelized benchmarks. We consider this trend extremely dangerous for the whole discipline. Representative conclusions require representative experiments and, as we argue in this paper, existing benchmark suites cannot be considered adequate to describe future CMP applications.

Large processor manufacturers have already reacted and developed their own, internal collections of workloads. An example is the Intel RMS benchmark suite[14]. However, these

suites often include proprietary code and are not publicly available. To address this problem, we created the PAR-SEC benchmark suite in collaboration with Intel Corporation. It includes not only a number of important applications from the RMS suite but also several leading-edge applications from Princeton University, Stanford University and the open-source domain. The goal is to create a suite of emerging workloads that can drive CMP research.

This paper makes three contributions:

- We identify shortcomings of commonly used benchmark suites and explain why they should not be used to evaluate CMPs.
- We present and characterize PARSEC, a new benchmark suite for CMPs that is diverse enough in order to allow representative conclusions.
- Based on our characterization of PARSEC, we analyze
 what properties future CMPs must have in order to
 be able to deliver scalable performance for emerging
 applications. In particular, our understanding of the
 behavior of future workloads allows us to quantify how
 CMPs must be built in order to mitigate the effects of
 the memory wall on the next generation of programs.

In Section 2 we describe why existing benchmark suites cannot be considered adequate to describe future CMP applications. In Section 3, we present the PARSEC benchmark suite and explain how it avoids the shortcomings of other collections of benchmarks. The methodology which we use to characterize our workloads is presented in Section 4. In Sections 5 to 8, we analyze the parallelization, working sets, communication behavior and off-chip traffic of the benchmark programs. We conclude our study in Section 9.

2. MOTIVATION

The goal of this work is to define a benchmark suite that can be used to design the next generation of processors. In this section, we first present the requirements for such a suite. We then discuss how the existing benchmarks fail to meet these requirements.

2.1 Requirements for a Benchmark Suite

We have the following five requirements for a benchmark suite:

Multi-threaded Applications Shared-memory CMPs are already ubiquitous. The trend for future processors is to deliver large performance improvements through increasing core counts on CMPs while only providing modest serial performance improvements. Consequently, applications that require additional processing power will need to be parallel.

Emerging Workloads Rapidly increasing processing power is enabling a new class of applications whose computational requirements were beyond the capabilities of the earlier generation of processors[14]. Such applications are significantly different from earlier applications (see Section 3). Future processors will be designed to meet the demands of these emerging applications and a benchmark suite should represent them.

Diverse Applications are increasingly diverse, run on a variety of platforms and accommodate different usage models. They include both interactive applications like computer games, offline applications like data mining programs and programs with different parallelization models. Specialized collections of benchmarks can be used to study some of these areas in more detail, but decisions about general-purpose processors should be based on a diverse set of applications. While a truly representative suite is impossible to create, reasonable effort should be made to maximize the diversity of the program selection. The number of benchmarks must be large enough to capture a sufficient amount of characteristics of the target application space.

Employ State-of-Art Techniques A number of application domains have changed dramatically over the last decade and use very different algorithms and techniques. Visual applications for example have started to increasingly integrate physics simulations to generate more realistic animations[20]. A benchmark should not only represent emerging applications but also use state-of-art techniques.

Support Research A benchmark suite intended for research has additional requirements compared to one used for benchmarking real machines alone. Benchmark suites intended for research usually go beyond pure scoring systems and provide infrastructure to instrument, manipulate, and perform detailed simulations of the included programs in an efficient manner.

2.2 Limitations of Existing Benchmark Suites

In the remaining part of this section we analyze how existing benchmark suites fall short of the presented requirements and must thus be considered unsuitable for evaluating CMP performance.

SPLASH-2 SPLASH-2 is a suite composed of multi-threaded applications [44] and hence seems to be an ideal candidate to measure performance of CMPs. However, its program collection is skewed towards HPC and graphics programs. It thus does not include parallelization models such as the pipeline model which are used in other application areas. SPLASH-2 should furthermore not be considered state-of-art anymore. Barnes

for example implements the Barnes-Hut algorithm for N-body simulation[8]. For galaxy simulations it has largely been superseded by the TreeSPH[19] method, which can also account for mass such as dark matter which is not concentrated in bodies. However, even for pure N-body simulation barnes must be considered outdated. In 1995 Xu proposed a hybrid algorithm which combines the hierarchical tree algorithm and the Fourier-based Particle-Mesh (PM) method to the superior TreePM method[45]. Our analysis shows that similar issues exist for a number of other applications of the suite including raytrace and radiosity.

SPEC CPU2006 and OMP2001 SPEC CPU2006 and SPEC OMP2001 are two of the largest and most significant collections of benchmarks. They provide a snapshot of current scientific and engineering applications. Computer architecture research, however, commonly focuses on the near future and should thus also consider emerging applications. Workloads such as systems programs and parallelization models which employ the producer-consumer model are not included. SPEC CPU2006 is furthermore a suite of serial programs that is not intended for studies of parallel machines.

Other Benchmark Suites Besides these major benchmark suites, several smaller suites exist. They were usually designed to study a specific program area and are thus limited to a single application domain. Therefore they usually include a smaller set of applications than a diverse benchmark suite typically offers. Due to these limitations they are commonly not used for scientific studies which do not restrict themselves to the covered application domain. Examples for these types of benchmark suites are ALPBench[25], BioParallel[22], MediaBench[1], NU-MineBench[23] and PhysicsBench[46]. Because of their different focus we do not discuss these suites in more detail.

3. THE PARSEC BENCHMARK SUITE

One of the goals of the PARSEC suite was to assemble a program selection that is large and diverse enough to be sufficiently representative for scientific studies. It consists of 9 applications and 3 kernels which were chosen from a wide range of application domains. In Table 1 we present a qualitative summary of their key characteristics. PARSEC workloads were selected to include different combinations of parallel models, machine requirements and runtime behaviors.

PARSEC meets all the requirements outlined in Section 2.1:

- Each of the applications has been parallelized.
- The PARSEC benchmark suite is not skewed towards HPC programs, which are abundant but represent only a niche. It focuses on emerging workloads. The algorithms these programs implement are usually considered useful, but their computational demands are prohibitively high on contemporary platforms. As more powerful processors become available in the near future, they are likely to proliferate rapidly.

Program	Application Domain	Paralle	lization	3371-: C-4	Data Usage	
		Model	Granularity	Working Set	Sharing	Exchange
blackscholes	Financial Analysis	data-parallel	coarse	small	low	low
bodytrack	Computer Vision	data-parallel	medium	medium	high	medium
canneal	Engineering	unstructured	fine	unbounded	high	high
dedup	Enterprise Storage	pipeline	medium	unbounded	high	high
facesim	Animation	data-parallel	coarse	large	low	medium
ferret	Similarity Search	pipeline	medium	unbounded	high	high
fluidanimate	Animation	data-parallel	fine	large	low	medium
freqmine	Data Mining	data-parallel	medium	unbounded	high	medium
streamcluster	Data Mining	data-parallel	medium	medium	low	medium
swaptions	Financial Analysis	data-parallel	coarse	medium	low	low
vips	Media Processing	data-parallel	coarse	medium	low	medium
x264	Media Processing	pipeline	coarse	medium	high	high

Table 1: Qualitative summary of the inherent key characteristics of PARSEC benchmarks. Working sets and data usage patterns are explained and quantified in later sections. The pipeline model is a data-parallel model which also uses a functional partitioning. PARSEC workloads were chosen to cover different application domains, parallel models and runtime behaviors.

- The workloads are diverse and were chosen from many different areas such as computer vision, media processing, computational finance, enterprise servers and animation physics.
- Each of the applications chosen represents the stateof-art technique in its area.
- PARSEC supports computer architecture research in a number of ways. The most important one is that for each workload six input sets with different properties are defined. Three of these inputs are suitable for microarchitectural simulation. We explain the different types of input sets in more detail in Section 3.1.

3.1 Input Sets

PARSEC defines six input sets for each benchmark:

test A very small input set to test the basic functionality of the program.

simdev A very small input set which guarantees basic program behavior similar to the real behavior, intended for simulator test and development.

simsmall, simmedium and simlarge Input sets of different sizes suitable for microarchitectural studies with simulators.

native A large input set intended for native execution.

test and simdev are merely intended for testing and development and should not be used for scientific studies. The three simulator inputs for studies vary in size, but the general trend is that larger input sets contain bigger working sets and more parallelism. Finally, the native input set is intended for performance measurements on real machines and exceeds the computational demands which are generally considered feasible for simulation by orders of magnitude. From a scientific point of view, the native input set is the most interesting one because it resembles real program

inputs most closely. The remaining input sets can be considered coarser approximations which sacrifice accuracy for tractability. Table 2 shows a breakdown of instructions and synchronization primitives of the simlarge input set which we used for the characterization study.

3.2 Workloads

The following workloads are part of the PARSEC suite:

3.2.1 blackscholes

The blackscholes application is an Intel RMS benchmark. It calculates the prices for a portfolio of European options analytically with the Black-Scholes partial differential equation (PDE)[10]

$$\frac{\partial V}{\partial t} + \frac{1}{2}\sigma^2 S^2 \frac{\partial^2 V}{\partial S^2} + rS \frac{\partial V}{\partial S} - rV = 0$$

where V is an option on the underlying S with volatility σ at time t if the constant interest rate is r. There is no closed-form expression for the Black-Scholes equation and as such it must be computed numerically[21]. The blackscholes benchmark was chosen to represent the wide field of analytic PDE solvers in general and their application in computational finance in particular. The program is limited by the amount of floating-point calculations a processor can perform.

blackscholes stores the portfolio with numOptions derivatives in array OptionData. The program includes file optionData.txt which provides the initialization and control reference values for 1,000 options which are stored in array data_init. The initialization data is replicated if necessary to obtain enough derivatives for the benchmark.

The program divides the portfolio into a number of work units equal to the number of threads and processes them concurrently. Each thread iterates through all derivatives in its contingent and calls function BlkSchlsEqEuroNoDiv for each of them to compute its price. If error checking was

Program	Problem Size	Instructions (Billions)				Synchronization Primitives		
		Total	FLOPS	Reads	Writes	Locks	Barriers	Conditions
blackscholes	65,536 options	2.67	1.14	0.68	0.19	0	8	0
bodytrack	4 frames, 4,000 particles	14.03	4.22	3.63	0.95	114,621	619	2,042
canneal	400,000 elements	7.33	0.48	1.94	0.89	34	0	0
dedup	184 MB data	37.1	0	11.71	3.13	158,979	0	1,619
facesim	1 frame,	29.90	9.10	10.05	4.29	14,541	0	3,137
	372,126 tetrahedra							
ferret	256 queries,	23.97	4.51	7.49	1.18	345,778	0	1255
	34,973 images							
fluidanimate	5 frames,	14.06	2.49	4.80	1.15	17,771,909	0	0
	300,000 particles							
freqmine	990,000 transactions	33.45	0.00	11.31	5.24	990,025	0	0
streamcluster	16,384 points per block,	22.12	11.6	9.42	0.06	191	129,600	127
	1 block							
swaptions	64 swaptions,	14.11	2.62	5.08	1.16	23	0	0
	20,000 simulations							
vips	1 image,	31.21	4.79	6.71	1.63	33,586	0	6,361
	2662×5500 pixels							
x264	128 frames,	32.43	8.76	9.01	3.11	16,767	0	1,056
	640×360 pixels							

Table 2: Breakdown of instructions and synchronization primitives for input set simlarge on a system with 8 cores. All numbers are totals across all threads. Numbers for synchronization primitives also include primitives in system libraries. 'Locks' and 'Barriers' are all lock- resp. barrier-based synchronizations, 'Conditions' are all waits on condition variables.

enabled at compile time it also compares the result with the reference price.

The input sets for blackscholes are sized as follows:

test: 1 optionsimdev: 16 options

simsmall: 4,096 options
simmedium: 16,384 options
simlarge: 65,536 options
native: 10,000,000 options

3.2.2 bodytrack

The bodytrack computer vision application is an Intel RMS workload which tracks a 3D pose of a marker-less human body with multiple cameras through an image sequence 13, 6]. bodytrack employs an annealed particle filter to track the pose using edges and the foreground silhouette as image features, based on a 10 segment 3D kinematic tree body model. These two image features were chosen because they exhibit a high degree of invariance under a wide range of conditions and because they are easy to extract. An annealed particle filter was employed in order to be able to search high dimensional configuration spaces without having to rely on any assumptions of the tracked body such as the existence of markers or constrained movements. This benchmark was included due to the increasing significance of computer vision algorithms in areas such as video surveillance, character animation and computer interfaces.

For every frame set Z_t of the input videos at time step t, the bodytrack benchmark executes the following steps:

- 1. The image features of observation Z_t are extracted. The features will be used to compute the likelihood of a given pose in the annealed particle filter.
- 2. Every time step t the filter makes an annealing run through all M annealing layers, starting with layer m=M.
- 3. Each layer m uses a set of N unweighted particles which are the result of the previous filter update step to begin with.

$$S_{t,m} = \{(s_{t,m}^{(1)})...(s_{t,m}^{(N)})\}.$$

Each particle $s_{t,m}^{(i)}$ is an instance of the multi-variate model configuration X which encodes the location and state of the tracked body.

4. Each particle $s_{t,m}^{(i)}$ is then assigned a weight $\pi_{t,m}^{(i)}$ by using weighting function $\omega(Z_t,X)$ corresponding to the likelihood of X given the image features in Z_t scaled by an annealing level factor:

$$\pi_{t,m}^{(i)} \propto \omega(Z_t, s_{t,m}^{(i)}).$$

The weights are normalized so that $\sum_{i=1}^{N} \pi_{t,m}^{(i)} = 1$. The result is the weighted particle set

$$S_{t,m}^{\pi} = \{(s_{t,m}^{(1)}, \pi_{t,m}^{(1)}) ... (s_{t,m}^{(N)}, \pi_{t,m}^{(N)})\}.$$

5. N particles are randomly drawn from set $S_{t,m}^{\pi}$ with a probability equal to their weight $\pi_{t,m}^{(i)}$ to obtain the temporary weighted particle set

$$\bar{S}_{t,m}^{\pi} = \{ (\bar{s}_{t,m}^{(1)}, \pi_{t,m}^{(1)}) ... (\bar{s}_{t,m}^{(N)}, \pi_{t,m}^{(N)}) \}.$$

Each particle $\bar{s}_{t,m}^{(i)}$ is then used to produce particle

$$s_{t m-1}^{(i)} = \bar{s}_{t m}^{(i)} + B_m$$

where B_m is a multi-variate Gaussian random variable. The result is particle set $S_{t,m-1}^{\pi}$ which is used to initialize layer m-1.

- 6. The process is repeated until all layers have been processed and the final particle set $S_{t,0}^{\pi}$ has been computed.
- 7. $S_{t,0}^{\pi}$ is used to compute the estimated model configuration χ_t for time step t by calculating the weighted average of all configuration instances:

$$\chi_t = \sum_{i=1}^{N} s_{t,0}^{(i)} \pi_{t,0}^{(i)}.$$

8. The set $S_{t+1,M}$ is then produced from $S_{t,0}^{\pi}$ using

$$s_{t+1}^{(i)} = s_{t,0}^{(i)} + B_0.$$

In the subsequent time step t+1 the set $S_{t+1,M}$ is used to initialize layer M.

The likelihood $\omega(Z_t, s_{t,m}^{(i)})$ which is used to determine the particle weights $\pi_{t,m}^{(i)}$ is computed by projecting the geometry of the human body model into the image observations Z_t for each camera and determining the error based on the image features. The likelihood is a measure of the 3D body model alignment with the foreground and edges in the images. The body model consists of conic cylinders to represent 10 body parts 2 for each limb plus the torso and the head. Each cylinder is represented by a length and a radius for each end. The body parts are assembled into a kinematic tree based upon the joint angles. Each particle represents the set of joint angles plus a global translation. To evaluate the likelihood of a given particle, the geometry of the body model is first built in 3D space given the angles and translation. Next, each 3D body part is projected onto each of the 2D images as a quadrilateral. A likelihood value is then computed based on the two image features the foreground map and the edge distance map. To compute the foreground term, samples are taken within the interior of each 2D body part projection and compared with the binary foreground map images. Samples that correspond to foreground increase the likelihood while samples that correspond to background are penalized. The edge map gives a measure of the distance from an edge in the image - values closer to an edge have a higher value. To compute the edge term samples are taken along the axis-parallel edges of each 2D body part projection and the edge map values at each sample are summed together. In this way, samples that are closer to edges in the images increase the likelihood while samples farther from edges are penalized.

bodytrack has a persistent thread pool which is implemented in class WorkPoolPthread. The main thread executes the program and sends a task to the thread pool with method SignalCmd whenever it reaches a parallel kernel. It resumes execution of the program as soon as it receives the result from the worker threads. Possible tasks are encoded by enumeration threadCommands in class WorkPoolPthread. The program has three parallel kernels:

Edge detection (Step 1) bodytrack employs a gradient based edge detection mask to find edges. The result is compared against a threshold to eliminate spurious edges. Edge detection is implemented in function GradientMagThreshold. The output of this kernel will be further refined before it is used to compute the particle weights.

Edge smoothing (Step 1) A separable Gaussian filter of size 7×7 pixels is used to smooth the edges in function GaussianBlur. The result is remapped between 0 and 1 to produce a pixel map in which the value of each pixel is related to its distance from an edge. The kernel has two parallel phases, one to filter image rows and one to filter image columns.

Calculate particle weights (Step 4) This kernel evaluates the foreground silhouette and the image edges produced earlier to compute the weights for the particles. This kernel is executed once for every annealing layer during every time step, making it the computationally most intensive part of the body tracker.

The parallel kernels use tickets to distribute the work among threads balance the load dynamically. The ticketing mechanism is implemented in class TicketDispenser and behaves like a shared counter.

The input sets for bodytrack are defined as follows:

- test: 4 cameras, 1 frame, 5 particles, 1 annealing layer
- simdev: 4 cameras, 1 frame, 100 particles, 3 annealing layers
- simsmall: 4 cameras, 1 frame, 1,000 particles, 5 annealing layers
- simmedium: 4 cameras, 2 frames, 2,000 particles, 5 annealing layers
- simlarge: 4 cameras, 4 frames, 4,000 particles, 5 annealing layers
- native: 4 cameras, 261 frames, 4,000 particles, 5 annealing layers

3.2.3 canneal

This kernel was developed by Princeton University. It uses cache-aware simulated annealing (SA) to minimize the routing cost of a chip design[7]. SA is a common method to approximate the global optimum in a large search space. Canneal pseudo-randomly picks pairs of elements and tries

to swap them. To increase data reuse, the algorithm discards only one element during each iteration which effectively reduces cache capacity misses. The SA method accepts swaps which increase the routing cost with a certain probability to make an escape from local optima possible. This probability continuously decreases during runtime to allow the design to converge. The program was included in the PARSEC program selection to represent engineering workloads, for the fine-grained parallelism with its lock-free synchronization techniques and due to its pseudo-random worst-case memory access pattern.

canneal uses a very aggressive synchronization strategy that is based on data race recovery instead of avoidance. Pointers to the elements are dereferenced and swapped atomically, but no locks are held while a potential swap is evaluated. This can cause disadvantageous swaps if one of the relevant elements has been replaced by another thread during that time. This equals a higher effective probability to accept swaps which increase the routing cost, and the SA method automatically recovers from it. The swap operation employs lock-free synchronization which is implemented with atomic instructions. An alternative implementation which relied on conventional locks turned out to be too inefficient due to excessive locking overhead. The synchronization routines with the atomic instructions are taken from the BSD kernel. Support for most new architectures can be added easily by copying the correct header file from the BSD kernel sources.

The annealing algorithm is implemented in the Run function of the annealer_thread class. Each thread uses the function get_random_element to pseudo-randomly pick one new netlist element per iteration with a Mersenne Twister[31]. calculate_delta_routing_cost is called to compute the change of the total routing cost if the two elements are swapped. accept_move evaluates the change in cost and the current temperature and decides whether the change is to be committed. Finally, accepted swaps are executed by calling swap_locations.

canneal implements an AtomicPtr class which encapsulates a shared pointer to the location of a netlist element. The pointer is atomically accessed and modified with the Get and Set functions offered by the class. A special Swap member function executes an atomic swap of two encapsulated pointers. If an access is currently in progress the functions spin until the operation could be completed. The implementation of Swap imposes a partial order to avoid deadlocks by processing the pointer at the lower memory location first.

We provide the following input sets for canneal:

- test: 5 swaps per temperature step, 100° start temperature, 10 netlist elements
- simdev: 100 swaps per temperature step, 300° start temperature, 100 netlist elements
- simsmall: 10,000 swaps per temperature step, 2,000° start temperature, 100,000 netlist elements
- simmedium: 15,000 swaps per temperature step, 2,000° start temperature, 200,000 netlist elements

- simlarge: 15,000 swaps per temperature step, 2,000° start temperature, 400,000 netlist elements
- native: 15,000 swaps per temperature step, 2,000° start temperature, 2,500,000 netlist elements

3.2.4 dedup

The dedup kernel was developed by Princeton University. It compresses a data stream with a combination of global compression and local compression in order to achieve high compression ratios. Such a compression is called 'deduplication'. The reason for the inclusion of this kernel is that deduplication has become a mainstream method to compress storage footprints for new-generation backup storage systems[36] and to compress communication data for new-generation bandwidth optimized networking appliances[39].

The kernel uses a pipelined programming model to parallelize the compression to mimic real-world implementations. There are five pipeline stages the intermediate three of which are parallel. In the first stage, dedup reads the input stream and breaks it up into coarse-grained chunks to get independent work units for the threads. The second stage anchors each chunk into fine-grained small segments with rolling fingerprinting[29, 11]. The third pipeline stage computes a hash value for each data segment. The fourth stage compresses each data segment with the Ziv-Lempel algorithm and builds a global hash table that maps hash values to data. The final stage assembles the deduplicated output stream consisting of hash values and compressed data segments.

Anchoring is a method which identifies brief sequences in a data stream that are identical with sufficiently high probability. It uses fast Rabin-Karp fingerprints[24] to detect identity. The data is then broken up into two separate blocks at the determined location. This method ensures that fragmenting a data stream is unlikely to obscure duplicate sequences since duplicates are identified on a block basis.

dedup uses a separate thread pool for each parallel pipeline stage. Each thread pool should at least have a number of threads equal to the number of available cores to allow the system to fully work on any stage should the need arise. The operating system scheduler is responsible for a thread schedule which will maximize the overall throughput of the pipeline. In order to avoid lock contention, the number of queues is scaled with the number of threads, with a small group of threads sharing an input and output queue at a time.

 ${\tt dedup}$ employs the following five kernels, one for each pipeline stage:

Coarse-grained fragmentation This serial kernel takes the input stream and breaks it up into work units which can be processed independently from each other by the parallel pipeline stages of dedup. It is implemented in function DataProcess. First, the kernel reads the input file from disk. It then determines the locations where the data is to be split up by jumping a fixed length in the buffer for each chunk. The resulting

data blocks are enqueued in order to be further refined by the subsequent stage.

Fine-grained fragmentation This parallel kernel uses Rabin-Karp fingerprints to break a coarse-grained data chunk up into fine-grained fragments. It scans each input block starting from the beginning. An anchor is found if the lowest 12 bits of the Rabin-Karp hash-sum are 0. The data is then split up at the location of the anchor. On average, this produces blocks of size $2^{12}/8 = 512$ bytes. The fine-grained data blocks are sent to the subsequent pipeline stage to compute their checksum. This kernel is implemented in function FindAllAnchor.

Hash computation To uniquely identify a fine-grained data block, this parallel kernel computes the SHA1 checksum of each chunk and checks for duplicate blocks with the use of a global database. It is implemented in function ChunkProcess. A hash table which is indexed with the SHA1 sum serves as the database. Each bucket of the hash table is associated with an independent lock in order to synchronize accesses. The large number of buckets and therefore locks makes the probability of lock contention very low in practice.

Once the SHA1 sum of a data block is available, the kernel checks whether a corresponding entry already exists in the database. If no entry could be found, the data block is added to the hash table and sent to the compression stage. If an entry already exists the block is classified as a duplicate. The compression stage is omitted and the block is sent directly to the pipeline stage which assembles the output stream.

Compression This kernel compresses data blocks in parallel. It is implemented in function Compress. Once the compressed image of a data block is available it is added to the database and the corresponding data block is sent to the next pipeline stage. Every data block is compressed only once because the previous stage does not send duplicates to the compression stage.

Assemble output stream This serial kernel reorders the data blocks and produces a compressed output stream. It is implemented in the SendBlock function. The stages which fragment the input stream into fine-grained data blocks add sequence numbers to allow a reconstruction of the original order. Because data fragmentation occurs in two different pipeline stages, two levels of sequence numbers have to be considered - one for each granularity level. SendBlock uses a search tree for the first level and a heap for the second level. The search tree allows rapid searches for the correct heap corresponding to the current first-level sequence number. For second-level sequence numbers only the minimum has to be found and hence a heap is used.

Once the next data block in the sequence becomes available it is removed from the reordering structures. If it has not been written to the output stream yet, its compressed image is emitted. Otherwise it is a duplicate and only its SHA1 signature is written as a placeholder. The kernel uses the global hash table to keep track of the output status of each data block.

Each input for dedup is an archive which contains a selection of files. The archives have the following sizes:

• test: 10 KB

• simdev: 1.1 MB

• simsmall: 10 MB
• simmedium: 31 MB

• simlarge: 184 MB

• native: 672 MB

3.2.5 facesim

This Intel RMS application was originally developed by Stanford University. It takes a model of a human face and a time sequence of muscle activations and computes a visually realistic animation of the modeled face by simulating the underlying physics[38, 40]. The goal is to create a visually realistic result. Certain effects such as inertial movements would have only a small visible effect and are not simulated[20]. The workload was included in the benchmark suite because an increasing number of computer games and other forms of animation employ physical simulation to create more realistic virtual environment. Human faces in particular are observed with more attention from users than other details of a virtual world, making their realistic presentation a key element for animations.

The parallelization uses a static partitioning of the mesh. Data that spans nodes belonging to more than one partition is replicated. Every time step the partitions process all elements that contain at least one node owned by the particle, but only results for nodes which are owned by the partition are written.

The iteration which computes the state of the face mesh at the end of each iteration is implemented in function Advance_-One_Time_Step_Quasistatic. facesim employs the fork-join model to process computationally intensive tasks in parallel. It uses the following three parallel kernels for its computations:

Update state This kernel uses the Newton-Raphson method to solve the nonlinear system of equations in order to find the steady state of the simulated mesh. This quasi-static scheme achieves speedups of one to two orders of magnitudes over explicit schemes by ignoring inertial effects. It is not suitable for the simulation of less constrained phenomena such as ballistic motion, but it is sufficiently accurate to simulate effects such as flesh deformation where the material is heavily influenced by contact, collision and self-collision and inertial effects only have a minor impact on the state.

In each Newton-Raphson iteration, the kernel reduces the nonlinear system of equations to a linear system which is guaranteed to be positive definite and symmetric. These two properties allow the use of a fast conjugate gradient solver later on. One iteration step is computed by function <code>Update_Position_Based_State</code>. The matrix of the linear system is sparse and can hence be stored in two one-dimensional arrays - dX_{full} and R_{full} . The matrix is the sum of the contribution of each tetrahedron of the face mesh.

Add forces This module computes the velocity-independent forces acting on the simulation mesh. After the matrix of the linear system with the position-independent state has been computed by the previous kernel, the right-hand side of that system has to be calculated. The kernel does this by iterating over all tetrahedra of the mesh, reading the positions of the vertices and computing the force contribution to each of the four nodes.

Conjugate gradient This kernel uses the conjugate gradient algorithm to solve the linear equation system assembled by the previous two modules. The two arrays dX_full and R_full which store the sparse matrix are sequentially accessed and matrix-vector multiplication is employed to solve the system.

The input sets of facesim all use the same face mesh. Scaling down the resolution of the mesh to create more tractable input sizes is impractical. A reduction of the number of elements in the model would result in under-resolution of the muscle action and cause problems for collision detection[20]. Our input sets for facesim are defined as follows:

• test: Print out help message.

• simdev: 80,598 particles, 372,126 tetrahedra, 1 frame

simsmall: Same as simdevsimmedium: Same as simdev

simlarge: Same as simdev

• native: Same as simdev, but with 100 frames

3.2.6 ferret

This application is based on the Ferret toolkit which is used for content-based similarity search of feature-rich data such as audio, images, video, 3D shapes and so on [27]. It was developed by Princeton University. The reason for the inclusion in the benchmark is that it represents emerging next-generation desktop and Internet search engines for non-text document data types. In the benchmark, we have configured the Ferret toolkit for image similarity search. Ferret is parallelized using the pipeline model with six stages. The first and the last stage are for input and output. The middle four stages are for query image segmentation, feature extraction, indexing of candidate sets with multi-probe Locality Sensitive Hashing (LSH)[28] and ranking. Each stage has its own thread pool and the basic work unit of the pipeline is a query image.

Segmentation is the process of decomposing an image into separate areas which display different objects. The rationale behind this step is that in many cases only parts of an image are of interest, such as the foreground. Segmentation allows the subsequent stages to assign a higher weight to image parts which are considered relevant and seem to belong together. After segmentation, ferret extracts a feature vector

from every segment. A feature vector is a multi-dimensional mathematical description of the segment contents. It encodes fundamental properties such as color, shape and area. Once the feature vectors are known, the indexing stage can query the image database to obtain a candidate set of images. The database is organized as a set of hash tables which are indexed with multi-probe LSH[28]. This method uses hash functions which map similar feature vectors to the same hash bucket with high probability. Because the number of hash buckets is very high, multi-probe LSH first derives a probing sequence which considers the success probabilities for finding a candidate image in a bucket. It then employs a step-wise approach which indexes buckets with a higher success probability first. After a candidate set of images has been obtained by the indexing stage, it is sent to the ranking stage which computes a detailed similarity estimate and orders the images according to their calculated rank. The similarity estimate is derived by analyzing and weighing the pair-wise distances between the segments of the query image and the candidate images. The underlying metric employed is the Earth Mover's Distance (EMD)[37]. For two images X and Y, it is defined as

$$EMD(X,Y) = min \sum_{i} \sum_{j} f_{ij} d(X_i, Y_j)$$

where X_i and Y_j denote segments of X and Y and f_{ij} is the extent to which X_i is matched to Y_j .

The first and the last pipeline stage of ferret are serial. The remaining four modules are parallel:

Image segmentation This kernel uses computer vision techniques to break an image up into non-overlapping segments. The pipeline stage is implemented in function t_seg, which calls image_segment for every image. This function uses statistical region merging (SRM)[33] to segment the image. This method organizes the pixels of an image in sets, starting with a fine-granular decomposition. It repeatedly merges them until the final segmentation has been reached.

Feature extraction This module computes a 14-dimensional feature vector for each image segment. The features extracted are the bounding box of the segment (5 dimensions) and its color moments (9 dimensions). A bounding box is the minimum axis-aligned rectangle which includes the segment. Color moments is a compact representation of the color distribution. It is conceptually similar to a histogram but uses fewer dimensions. Segments are assigned a weight which is proportional to the square root of its size. This stage is implemented in function t_extract. It calls image_extract_helper to compute the feature vectors for every image.

Indexing The indexing stage queries the image database to obtain no more than twice the number of images which are allowed to appear in the final ranking. This stage is implemented in function t_vec. ferret manages image data in tables which have type cass_table_t. Tables can be queried with function cass_table_query. The indexing stage uses this function to access the database in order to generate the candidate set of type <code>cass_result_t</code> for the current query image. Indexing employs LSH for the probing which is implemented in function LSH_query.

Ranking This module performs a detailed similarity computation. From the candidate set obtained by the indexing stage it chooses the final set of images which are most similar to the query image and ranks them. The ranking stage is implemented in function t_rank. It employs cass_table_query to analyze the candidate set and to compute the final ranking with EMD. The type of query that cass_table_query is to perform can be described with a structure of type cass_query_t.

The number of query images determine the amount of parallelism. The working set size is dominated by the size of the image database. The input sets for ferret are sized as follows:

- test: 1 image queries, database with 1 image, find top 1 image
- simdev: 4 image queries, database with 100 images, find top 5 images
- simsmall: 16 image queries, database with 3,544 images, find top 10 images
- simmedium: 64 image queries, database with 13,787 images, find top 10 images
- simlarge: 256 image queries, database with 34,973 images, find top 10 images
- native: 3,500 image queries, database with 59,695 images, find top 50 images

3.2.7 fluidanimate

This Intel RMS application uses an extension of the Smoothed Particle Hydrodynamics (SPH) method to simulate an incompressible fluid for interactive animation purposes[32]. Its output can be visualized by detecting and rendering the surface of the fluid. The force density fields are derived directly from the Navier-Stokes equation. fluidanimate uses special-purpose kernels to increase stability and speed. fluidanimate was included in the PARSEC benchmark suite because of the increasing significance of physics simulations for computer games and other forms of real-time animations.

A simplified version of the Navier-Stokes equation for incompressible fluids[35] which formulates conservation of momentum is

$$\rho(\frac{\partial v}{\partial t} + v \cdot \nabla v) = -\nabla p + \rho g + \mu \nabla^2 v$$

where v is a velocity field, ρ a density field, p a pressure field, g an external force density field and μ the viscosity of the fluid. The SPH method uses particles to model the

state of the fluid at discrete locations and interpolates intermediate values with radial symmetrical smoothing kernels. An advantage of this method is the automatic conservation of mass due to a constant number of particles, but it alone does not guarantee certain physical principals such as symmetry of forces which have to be enforced separately. The SPH algorithm derives a scalar quantity A at location r by a weighted sum of all particles:

$$A_S(r) = \sum_{j} m_j \frac{A_j}{\rho_j} W(r - r_j, h).$$

In the equation, j iterates over all particles, m_j is the mass of particle j, r_j its position, ρ_j the density at its location and A_j the respective field quantity. $W(r-r_j,h)$ is the smoothing kernel to use for the interpolation with core radius h. Smoothing kernels are employed in order to make the SPH method stable and accurate. Because each particle i represents a volume with constant mass m_i , the density ρ_i appears in the equation and has to be recomputed every time step. The density at a location r can be calculated by substituting A with ρ in the previous equation:

$$\rho_S(r) = \sum_j m_j W(r - r_j, h)$$

Applying the SPH interpolation equation to the pressure term $-\nabla p$ and the viscosity term $\mu \nabla^2$ of the Navier-Stokes equation yields the equations for the pressure and viscosity forces, but in order to solve the force symmetry problems of the SPH method, fluidanimate employs slightly modified formulas:

$$f_i^{pressure} = -\sum_i m_j \frac{p_i + p_j}{2\rho_j} \nabla W(r_i - r_j, h)$$

$$f_i^{viscosity} = \mu \sum_i m_j \frac{v_i - v_j}{\rho_j} \nabla^2 W(r_i - r_j, h)$$

Stability, accuracy and speed of fluidanimate are highly dependent on its smoothing kernels. In all cases but the pressure and viscosity computations the program uses the following kernel:

$$W_{poly6}(r,h) = \frac{315}{64\pi h^9} \begin{cases} (h^2 - r^2)^3 & 0 \le r \le h \\ 0 & else \end{cases}$$

One feature of this kernel is that the distance r only appears squared. The computation of square roots is thus not necessary to evaluate it. For pressure computations, fluidanimate uses Desbrun's spiky kernel $W_{spiky}[12]$ and $W_{viscosity}$ for viscosity forces:

$$W_{spiky}(r,h) = \frac{15}{\pi h^6} \begin{cases} (h-r)^3 & 0 \le r \le h \\ 0 & else \end{cases}$$

$$W_{viscosity}(r,h) = \frac{15}{2\pi h^3} \begin{cases} -\frac{r^3}{2h^3} + \frac{r^2}{h^2} + \frac{h}{2r} - 1 & 0 \le r \le h \\ 0 & else \end{cases}$$

The scene geometry employed by fluidanimate is a box in which the fluid resides. All collisions are handled by adding forces in order to change the direction of movement of the involved particles instead of modifying the velocity directly. The workload uses Verlet integration[42] to update the position of the particles. This scheme does not store the velocity of the particles explicitly, but their previous location in addition to the current position. The current velocity can thus be deduced from the distance traveled since the last time step. The force and mass are then used to compute the acceleration and subsequently the new velocity. This scheme is more robust because the velocity is implicitly given.

Every time step, fluidanimate executes five kernels, the first two of which were further broken up into several smaller steps:

Rebuild spatial index Because the smoothing kernels $W(r-r_j,h)$ have finite support h, particles can only interact with each other up to the distance h. The program uses a spatial indexing structure in order to exploit proximity information and limit the number of particles which have to be evaluated. Functions ClearParticles and RebuildGrid build this acceleration structure which is used by the subsequent steps.

Compute densities This kernel estimates the fluid density at the position of each particle by analyzing how closely particles are packed in its neighborhood. In a region in which particles are packed together more closely, the density will be higher. This kernel has 3 phases which are implemented in the functions Init-DensitiesAndForces, ComputeDensities and ComputeDensities2.

Compute forces Once the densities are known, they can be used to compute the forces. This step happens in function ComputeForces. The kernel evaluates pressure, viscosity and also gravity as the only external influence. Collisions between particles are handled implicitly during this step, too.

Handle collisions with scene geometry The next kernel updates the forces in order to handle collisions of particles with the scene geometry. This step is implemented in function ProcessCollisions.

Update positions of particles Finally, the forces can be used to calculate the acceleration of each particle and update its position. fluidanimate uses a Verlet integrator [42] for these computations which is implemented in function AdvanceParticles.

The input sets for fluidanimate are sized as follows:

• test: 5,000 particles, 1 frame

• simdev: 15,000 particles, 3 frames

• simsmall: 35,000 particles, 5 frames

• simmedium: 100,000 particles, 5 frames

simlarge: 300,000 particles, 5 frames
native: 500,000 particles, 500 frames

3.2.8 frequine

The frequine application employs an array-based version of the FP-growth (Frequent Pattern-growth) method[15] for Frequent Itemset Mining (FIMI). It is an Intel RMS benchmark which was originally developed by Concordia University. FIMI is the basis of Association Rule Mining (ARM), a very common data mining problem which is relevant for areas such as protein sequences, market data or log analysis. The serial program this benchmark is based on won the FIMI'03 best implementation award for its efficiency. frequine was included in the PARSEC benchmark suite because of the increasing demand for data mining techniques which is driven by the rapid growth of the volume of stored information.

FP-growth stores all relevant frequency information of the transaction database in a compact data structure called FPtree (Frequent Pattern-tree)[16]. An FP-tree is composed of three parts: First, a prefix tree encodes the transaction data such that each branch represents a frequent itemset. The nodes along the branches are stored in decreasing order of frequency of the corresponding item. The prefix tree is a more compact representation of the transaction database because overlapping itemsets share prefixes of the corresponding branches. The second component of the FP-tree is a header table which stores the number of occurrences of each item in decreasing order of frequency. Each entry is also associated with a pointer to a node of the FP-tree. All nodes which are associated with the same item are linked to a list. The list can be traversed by looking up the corresponding item in the header table and following the links to the end. Each node furthermore contains a counter that encodes how often the represented itemset as seen from the root to the current node occurs in the transaction database. The third component of the FP-tree is a lookup table which stores the frequencies of all 2-itemsets. A row in the lookup table gives all occurrences of items in itemsets which end with the associated item. This information can be used during the mining phase to omit certain FP-tree scans and is the major improvement of the implemented algorithm. The lookup table is especially effective if the dataset is sparse which is usually the case. The FP-trees are then very big due to the fact that only few prefixes are shared. In that case tree traversals are more expensive, and the benefit from being able to omit them is greater. The initial FP-tree can be constructed with only two scans of the original database, the first one to construct the header table and the second one to compute the remaining parts of the FP-tree.

In order to mine the data for frequent itemsets, the FP-growth method traverses the FP-tree data structure and recursively constructs new FP-trees until the complete set of frequent itemsets is generated. To construct a new FP-tree

 $T_{X \cup \{i\}}$ for an item i in the header of an existing FP-tree T_X , the algorithm first obtains a new pattern base from the lookup table. The base is used to initialize the header of the new tree $T_{X \cup \{i\}}$. Starting from item i in the header table of the existing FP-tree T_X , the algorithm then traverses the associated linked list of all item occurrences. The patterns associated with the visited branches are then inserted into the new FP-tree $T_{X \cup \{i\}}$. The resulting FP-tree is less bushy because it was constructed from fewer itemsets. The recursion terminates when an FP-tree was built which has only one path. The properties of the algorithm guarantee that this is a frequent itemset.

frequine has been parallelized with OpenMP. It employs three parallel kernels:

Build FP-tree header This kernel scans the transaction database and counts the number of occurrences of each item. It performs the first of two database scans necessary to construct the FP-tree. The result of this operation is the header table for the FP-tree which contains the item frequency information. This kernel has one parallelized loop and is implemented in function scan1_DB.

Construct prefix tree The next kernel builds the initial tree structure of the FP-tree. It performs the second and final scan of the transaction database necessary to build the data structures which will be used for the actual mining operation. The kernel has four parallelized loops. It is implemented in function scan2_DB which contains two of them. The remaining two loops are in its helper function database_tiling.

Mine data The last kernel uses the data structures previously computed and mines them to recursively obtain the frequent itemset information. It is an improved version of the conventional FP-growth method[16]. This module has similarities with the previous two kernels which construct the initial FP-tree because it builds a new FP-tree for every recursion step.

The module is implemented in function FP_growth_first. It first derives the initial lookup table from the current FP-tree by calling first_transform_FPTree_into_FP-Array. This function executes the first of two parallelized loops. After that the second parallelized loop is executed in which the recursive function FP_growth is called. It is the equivalent of FP_growth_first. Each thread calls FP_growth independently so that a number of recursions up to the number of threads can be active.

The input sets for freqmine are defined as follows:

- test: Database with 3 synthetic transactions, minimum support 1.
- simdev: Database with 1,000 synthetic transactions, minimum support 3.
- simsmall: Database with 250,000 anonymized click streams from a Hungarian online news portal, minimum support 220.

- simmedium: Same as simmall but with 500,000 click streams, minimum support 410.
- simlarge: Same as simsmall but with 990,000 click streams, minimum support 790.
- native: Database composed of spidered collection of 250,000 web HTML documents[26], minimum support 11,000.

3.2.9 streamcluster

This RMS kernel was developed by Princeton University and solves the online clustering problem[34]: For a stream of input points, it finds a predetermined number of medians so that each point is assigned to its nearest center. The quality of the clustering is measured by the sum of squared distances (SSQ) metric. Stream clustering is a common operation where large amounts or continuously produced data has to be organized under real-time conditions, for example network intrusion detection, pattern recognition and data mining. The program spends most of its time evaluating the gain of opening a new center. This operation uses a parallelization scheme which employs static partitioning of data points. The program is memory bound for low-dimensional data and becomes increasingly computationally intensive as the dimensionality increases. Due to its online character the working set size of the algorithm can be chosen independently from the input data. streamcluster was included in the PARSEC benchmark suite because of the importance of data mining algorithms and the prevalence of problems with streaming characteristics.

The parallel gain computation is implemented in function pgain. Given a preliminary solution, the function computes how much cost can be saved by opening a new center. For every new point, it weighs the cost of making it a new center and reassigning some of the existing points to it against the savings caused by minimizing the distance

$$d(x, y) = |x - y|^2$$

between two points x and y for all points. The distance computation is implemented in function \mathtt{dist} . If the heuristic determines that the change would be advantageous the results are committed.

The amount of parallelism and the working set size of a problem are dominated by the block size. The input sets of swaptions are defined as follows:

- test: 10 input points, block size 10 points, 1 point dimension, 2–5 centers, up to 5 intermediate centers allowed
- simdev: 16 input points, block size 16 points, 3 point dimensions, 3–10 centers, up to 10 intermediate centers allowed
- simsmall: 4,096 input points, block size 4,096 points, 32 point dimensions, 10–20 centers, up to 1,000 intermediate centers allowed

- simmedium: 8,192 input points, block size 8,192 points, 64 point dimensions, 10-20 centers, up to 1,000 intermediate centers allowed
- simlarge: 16,384 input points, block size 16,384 points, 128 point dimensions, 10-20 centers, up to 1,000 intermediate centers allowed
- native: 1,000,000 input points, block size 200,000 points, 128 point dimensions, 10–20 centers, up to 5,000 intermediate centers allowed

3.2.10 swaptions

The swaptions application is an Intel RMS workload which uses the Heath-Jarrow-Morton (HJM) framework to price a portfolio of swaptions. The HJM framework describes how interest rates evolve for risk management and asset liability management[17] for a class of models. Its central insight is that there is an explicit relationship between the drift and volatility parameters of the forward-rate dynamics in a no-arbitrage market. Because HJM models are non-Markovian the analytic approach of solving the PDE to price a derivative cannot be used. Swaptions therefore employs Monte Carlo (MC) simulation to compute the prices. The workload was included in the benchmark suite because of the significance of PDEs and the wide use of Monte Carlo simulation.

The program stores the portfolio in the swaptions array. Each entry corresponds to one derivative. Swaptions partitions the array into a number of blocks equal to the number of threads and assigns one block to every thread. Each thread iterates through all swaptions in the work unit it was assigned and calls the function HJM_Swaption_Blocking for every entry in order to compute the price. This function invokes HJM_SimPath_Forward_Blocking to generate a random HJM path for each MC run. Based on the generated path the value of the swaption is computed.

The following input sets are provided for swaptions:

• test: 1 swaption, 5 simulations

• simdev: 3 swaptions, 50 simulations

• simsmall: 16 swaptions, 5,000 simulations

• simmedium: 32 swaptions, 10,000 simulations

ullet simlarge: 64 swaptions, 20,000 simulations

• native: 128 swaptions, 1,000,000 simulations

3.2.11 *vips*

This application is based on the VASARI Image Processing System (VIPS)[30] which was originally developed through several projects funded by European Union (EU) grants. The benchmark version is derived from a print on demand service that is offered at the National Gallery of London, which is also the current maintainer of the system. The benchmark includes fundamental image operations such as an affine transformation and a convolution. It was chosen because image transformations are a common task on desktop computers and for the ability of the VASARI system to

construct multi-threaded image processing pipelines transparently on the fly. Future libraries might use concepts such as the ones employed by VIPS to make multi-threaded functionality available to the user.

The image transformation pipeline of the vips benchmark has 18 stages. It is implemented in the VIPS operation im_benchmark. The stages can be grouped into the following kernels:

Crop The first step of the pipeline is to remove 100 pixels from all edges with VIPS operation im_extract_area.

Shrink Next, vips shrinks the image by 10%. This affine transformation is implemented as the matrix operation

$$f(\vec{x}) = \begin{bmatrix} 0.9 & 0 \\ 0 & 0.9 \end{bmatrix} \vec{x} + \begin{bmatrix} 0 \\ 0 \end{bmatrix}$$

in VIPS operation im_affine. The transformation uses bilinear interpolation to compute the output values.

Adjust white point and shadows To improve the perceived visual quality of the image under the expected target conditions, vips brightens the image, adjusts the white point and pulls the shadows down. These operations require several linear transformations and a matrix multiplication, which are implemented in im_lintra, im_lintra_vec and im_recomb.

Sharpen The last step slightly exaggerates the edges of the output image in order to compensate for the blurring caused by printing and to give the image a better overall appearance. This convolution employs a Gaussian blur filter with mask radius 11 and a subtraction in order to isolate the high-frequency signal component of the image. The intermediate result is transformed via a look-up table shaped as

$$f(x) = \begin{cases} 0.5x & |x| \le 2.5\\ 1.5x + 2.5 & x < -2.5\\ 1.5x - 2.5 & x > 2.5 \end{cases}$$

and added back to the original image to obtain the sharpened image. Sharpening is implemented in VIPS operation im_sharpen.

The VASARI Image Processing System fuses all image operations to construct an image transformation pipeline that can operate on subsets of an image. VIPS can automatically replicate the image transformation pipeline in order to process multiple image regions concurrently. This happens transparently for the user of the library. Actual image processing and any I/O is deferred as long as possible. Intermediate results are represented in an abstract way by partial image descriptors. Each VIPS operation can specify a demand hint which is evaluated to determine the work unit size of the combined pipeline. VIPS uses memory-mapped I/O to load parts of an input image on demand. After the requested part of a file has been loaded, all image operations

are applied to the image region before the output region is written back to disk.

A VIPS operation is composed of the main function which provides the public interface employed by the users, the generate function which implements the actual image operation, as well as a start and a stop function. The main functions register the operation with the VIPS evaluation system. Start functions are called by the runtime system to perform any per-thread initialization. They produce a sequence value which is passed to all generate functions and the stop function. Stop functions handle the shutdown at the end of the evaluation phase and destroy the sequence value. The VIPS system guarantees the mutually exclusive execution of start and stop functions, which can thus be used to communicate between threads during the pipeline initialization or shutdown phase. The generate functions transform the image and correspond to the pipeline stages.

The sizes of the images used for the input sets for vips are:

test: 256 × 288 pixels
 simdev: 256 × 288 pixels

• simsmall: $1,600 \times 1,200$ pixels

 \bullet simmedium: $2,336\times 2,336~\mathrm{pixels}$

• simlarge: $2,662 \times 5,500$ pixels • native: $18,000 \times 18,000$ pixels

$3.2.12 \quad x264$

The x264 application is an H.264/AVC (Advanced Video Coding) video encoder. In the 4th annual video codec comparison[41] it was ranked 2nd best codec for its high encoding quality. It is based on the ITU-T H.264 standard which was completed in May 2003 and which is now also part of ISO/IEC MPEG-4. In that context the standard is also known as MPEG-4 Part 10. H.264 describes the lossy compression of a video stream[43]. It improves over previous video encoding standards with new features such as increased sample bit depth precision, higher-resolution color information, variable block-size motion compensation (VB-SMC) or context-adaptive binary arithmetic coding (CABAC). These advancements allow H.264 encoders to achieve a higher output quality with a lower bit-rate at the expense of a significantly increased encoding and decoding time. The flexibility of H.264 allows its use in a wide range of contexts with different requirements, from video conferencing solutions to high-definition (HD) movie distribution. Nextgeneration HD DVD or Blu-ray video players already require H.264/AVC encoding. The flexibility and wide range of application of the H.264 standard and its ubiquity in nextgeneration video systems are the reasons for the inclusion of x264 in the PARSEC benchmark suite.

H.264 encoders and decoders operate on macroblocks of pixels which have the fixed size of 16×16 pixels. Various techniques are used to detect and eliminate data redundancy. The most important one is motion compensation. It is employed to exploit temporal redundancy between successive frames. Motion compensation is usually the most expensive

operation that has to be executed to encode a frame. It has a very high impact on the final compression ratio. The compressed output frames can be encoded in one of three possible ways:

- I-Frame An I-Frame includes the entire image and does not depend on other frames. All its macroblocks are encoded using intra prediction. In intra mode, a prediction block is formed using previously encoded blocks. This prediction block is subtracted from the current block prior to encoding.
- P-Frame These frames include only the changed parts of an image from the previous I- or P-frame. A P-Frame is encoded with intra prediction and inter prediction with at most one motion-compensated prediction signal per prediction block. The prediction model is formed by shifting samples from previously encoded frames to compensate for motion such as camera pans.
- **B-Frame** B-Frames are constructed using data from the previous and next I- or P-Frame. They are encoded like a P-frame but using inter prediction with two motion-compensated prediction signals. B-Frames can be compressed much more than other frame types.

The enhanced inter and intra prediction techniques of H.264 are the main factors for its improved coding efficiency. The prediction schemes can operate on block of varying size and shapes which can be as small as 4×4 pixels.

The parallel algorithm of x264 uses the pipeline model with one stage per input video frame. This results in a virtual pipeline with as many stages as there are input frames. x264 processes a number of pipeline stages equal to the number of encoder threads in parallel, resulting in a sliding window which moves from the beginning of the pipeline to its end. For P- and B-Frames the encoder requires the image data and motion vectors from the relevant region of the reference frames in order to encode the current frame, and so each stage makes this information available as it is calculated during the encoding process. Fast upward movements can thus cause delays which can limit the achievable speedup of x264 in practice. In order to compensate for this effect, the parallelization model requires that x264 is executed with a number of threads greater than the number of cores to achieve maximum performance.

x264_encoder_encode to encode another frame. x264_encoder_encode uses function x264_slicetype_decide to determine as which type the frame will be encoded and calls all necessary functions to produce the correct output. It also manages the threading functionality of x264. Threads use the functions x264_frame_cond_broadcast and x264_frame_cond_wait to inform each other of the encoding progress and to make sure that no data is accessed while it is not yet available.

The videos used for the input sets have been derived from the uncompressed version of the short film "Elephants Dream" [3]. The number of frames determines the amount of parallelism. The exact characteristics of the input sets are: • test: 32×18 pixels, 1 frame

• simdev: 64×36 pixels, 3 frames

• simsmall: 640×360 pixels ($\frac{1}{3}$ HDTV resolution), 8 frames

• simmedium: 640×360 pixels ($\frac{1}{3}$ HDTV resolution), 32 frames

• simlarge: 640×360 pixels ($\frac{1}{3}$ HDTV resolution), 128 frames

• native: $1,920 \times 1,080$ pixels (HDTV resolution), 512 frames

4. METHODOLOGY

In this section we explain how we characterized the PAR-SEC benchmark suite. We are interested in the following characteristics:

Parallelization PARSEC benchmarks use different parallel models which have to be analyzed in order to know whether the programs can scale well enough for the analysis of CMPs of a certain size.

Working sets and locality Knowledge of the cache requirements of a workload are necessary to identify benchmarks suitable for the study of CMP memory hierarchies.

Communication to computation ratio and sharing The communication patterns of a program determine the potential impact of private caches and the on-chip network on performance.

Off-chip traffic The off-chip traffic requirements of a program are important to understand how off-chip bandwidth limitations of a CMP can affect performance.

In order to characterize all applications, we had to make several trade-off decisions. Given a limited amount of computational resources, higher accuracy comes at the expense of a lower number of experiments. We followed the approach of similar studies[44, 22] and chose faster but less accurate execution-driven simulation to characterize the PAR-SEC workloads. This approach is feasible because we limit ourselves to study fundamental program properties which should have a high degree of independence from architectural details. Where possible we supply measurement results from real machines. This methodology made it possible to gather the large amount of data which we present in this study. We preferred machine models comparable to real processors over unrealistic models which might have been a better match for the program needs.

4.1 Experimental Setup

We used CMP\$im[22] for our workload characterization. CMP\$im is a plug-in for Pin[2] that simulates the cache hierarchy of a CMP. Pin is similar to the ATOM toolkit for Compaq's Tru64 Unix on Alpha processors. It uses dynamic binary instrumentation to insert routines at arbitrary points in the instruction stream. For the characterization we simulate a single-level cache hierarchy of a CMP and vary its

parameters. The baseline cache configuration was a shared 4-way associative cache with 4 MB capacity and 64 byte lines. By default the workloads used 8 cores. All experiments were conducted on a set of Symmetric Multiprocessor (SMP) machines with x86 processors and Linux. The programs were compiled with gcc 4.2.1.

Because of the large computational cost we could not perform simulations with the native input set, instead we used the simlarge inputs for all simulations and analytically describe any differences between the two sets of which we know.

4.2 Methodological Limitations and Error Margins

For their characterization of the SPLASH-2 benchmark suite, Woo et al. fixed a timing model which they used for all experiments[44]. They give two reasons: First, nondeterministic programs would otherwise be difficult to compare because different execution paths could be taken, and second, the characteristics they study are largely independent from an architecture. They also state that they believe that the timing model should have only a small impact on the results. While we use similar characteristics and share this belief, we think a characterization study of multi-threaded programs should nevertheless analyze the impact of nondeterminism on the reported data. Furthermore, because our methodology is based on execution on real machines combined with dynamic binary instrumentation, it can introduce additional latencies, and a potential concern is that the nondeterministic thread schedule is altered in a way that might affect our results in unpredictable ways. We therefore conducted a sensitivity analysis to quantify the impact of nondetermin-

Alameldeen and Wood studied the variability of nondeterministic programs in more detail and showed that even small pseudo-random perturbations of memory latencies are effective to force alternate execution paths[5]. We adopted their approach and modified CMP\$im to add extra delays to its analysis functions. Because running all experiments multiple times as Alameldeen and Wood did would be prohibitively expensive, we instead decided to randomly select a subset of all experiments for each metric which we use and report its error margins.

The measured quantities deviated by no more than $\pm 0.04\%$ from the average, with the following two exceptions. The first exception is metrics of data sharing. In two cases (bodytrack and swaptions) the classification is noticeably affected by the nondeterminism of the program. This is partially caused because shared and thread-private data contend aggressively for a limited amount of cache capacity. The high frequency of evictions made it difficult to classify lines and accesses as shared or private. In these cases, the maximum deviation of the number of accesses from the average was as high as $\pm 4.71\%$, and the amount of sharing deviated by as much as $\pm 15.22\%$. We considered this uncertainty in our study and did not draw any conclusions where the variation of the measurements did not allow it. The second case of high variability is when the value of the measured quantity is very low (below 0.1% miss rate or corresponding ratio). In these cases the nondeterministic noise

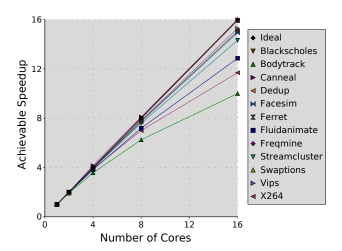


Figure 1: Upper bound for speedup of PARSEC workloads with input set simlarge based on instruction count. Limitations are caused by serial sections, growing parallelization overhead and redundant computations.

made measurements difficult. We do not consider this a problem because in this study we focus on trends of ratios, and quantities that small do not have a noticeable impact. It is however an issue for the analysis of working sets if the miss rate falls below this threshold and continues to decrease slowly. Only few programs are affected, and our estimate of their working set sizes might be slightly off in these cases. This is primarily an issue inherent to experimental working set analysis, since it requires well-defined points of inflection for conclusive results. Moreover, we believe that in these cases the working set size varies nondeterministically, and researchers should expect slight variations for each benchmark run.

The implications of these results are twofold: First, they show that our methodology is not susceptible to the non-deterministic effects of multi-threaded programs in a way that might invalidate our findings. Second, they also confirm that the metrics which we present in this paper are fundamental program properties which cannot be distorted easily. The reported application characteristics are likely to be preserved on a large range of architectures.

5. PARALLELIZATION

In this section we discuss the parallelization of the PAR-SEC suite. As we will see in Section 6, several PARSEC benchmarks (canneal, dedup, ferret and freqmine) have working sets so large they should be considered unbounded for an analysis. These working sets are only limited by the amount of main memory in practice and they are actively used for inter-thread communication. The inability to use caches efficiently is a fundamental property of these program and affects their concurrent behavior. Furthermore, dedup and ferret use a complex, heterogeneous parallelization model in which specialized threads execute different functions with different characteristics at the same time. These programs employ a pipeline with dedicated thread pools for each parallelized pipeline stage. Each thread pool

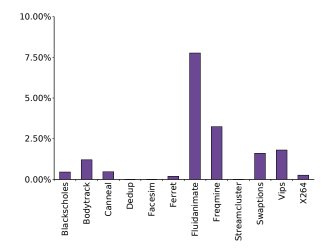


Figure 2: Parallelization overhead of PARSEC benchmarks. The chart shows the slowdown of the parallel version on 1 core over the serial version.

has enough threads to occupy the whole CMP, and it is the responsibility of the scheduler to assign cores to threads in a manner that maximizes the overall throughput of the pipeline. Over time, the number of threads active for each stage will converge against the inverse throughput ratios of the individual pipeline stages relative to each other.

Woo et al. use an abstract machine model with a uniform instruction latency of one cycle to measure the speedups of the SPLASH-2 programs[44]. They justify their approach by pointing out that the impact of the timing model on the characteristics which they measure - including speedup - is likely to be low. Unfortunately, this is not true in general for PARSEC workloads. While we have verified in Section 4.2 that the fundamental program properties such as miss rate and instruction count are largely not susceptible to timing shocks, the synchronization and timing behavior of the programs is. Using a timing model with perfect caches significantly alters the behavior of programs with unbounded working sets, for example how long locks to large, shared data structures are held. Moreover, any changes of the timing model have a strong impact on the number of active threads of programs which employ thread specialization. It will thus affect the load balance and synchronization behavior of these workloads. We believe it is not possible to discuss the timing behavior of these programs without also considering for example different schedulers, which is beyond the scope of this paper. Similar dependencies of commercial workloads on their environment are already known[9, 4].

Unlike Woo et al. who measured actual concurrency on an abstract machine, we therefore decided to analyze inherent concurrency and its limitations. Our approach is based on the number of executed instructions in parallel and serial regions of the code. We neglect any delays due to blocking on contended locks and load imbalance. This methodology is feasible because we do not study performance, our interest is in fundamental program characteristics. The presented data is largely timing-independent and a suitable measure of the concurrency inherent in a workload. The results in

Figure 1 show the maximum achievable speedup measured that way. The numbers account for limitations such as unparallelized code sections, synchronization overhead and redundant computations. PARSEC workloads can achieve actual speedups close to the presented numbers. We verified on a large range of architectures that lock contention and other timing-dependent factors are not limiting factors, but we know of no way to show it in a platform-independent way given the complications outlined above. The maximum speedup of bodytrack, x264 and streamcluster is limited by serial sections of the code. fluidanimate is primarily limited by growing parallelization overhead. On real machines, x264 is furthermore bound by a data dependency between threads, however this has only a noticeable impact on machines larger than the ones described here. It is recommended to run x264 with more threads than cores, since modeling and exposing these dependencies to the scheduler is a fundamental aspect of its parallel algorithm, comparable to the parallel algorithms of dedup and ferret. Figure 2 shows the slowdown of the parallel version on 1 core over the serial version. The numbers show that all workloads use efficient parallel algorithms which are not substantially slower than the corresponding serial algorithms.

PARSEC programs scale well enough to study CMPs. We believe they are also useful on machines larger than the ones analyzed here. The PARSEC suite exhibits a wider variety of parallelization models than previous benchmark suites such as the pipeline model. Some of its workloads can adapt to different timing models and can use threads to hide latencies. It is important to analyze these programs in the context of the whole system.

6. WORKING SETS AND LOCALITY

The temporal locality of a program can be estimated by analyzing how the miss rate of a processor's cache changes as its capacity is varied. Often the miss rate does not decrease continuously as the size of a cache is increased, but stays on a certain level and then makes a sudden jump to a lower level when the capacity becomes large enough to hold the next important data structure. For CMPs an efficient functioning of the last cache level on the chip is crucial because a miss in the last level will require an access to off-chip memory.

To analyze the working sets of the PARSEC workloads we studied a cache shared by all processors. Our results are presented in Figure 3. In Table 3 we summarize the important characteristics of the identified working sets. Most workloads exhibit well-defined working sets with clearly identifiable points of inflection. Compared to SPLASH-2, PARSEC working sets are significantly larger and can reach hundreds of megabytes such as in the cases of canneal and frequine.

Two types of workloads can be distinguished: The first group contains benchmarks such as bodytrack and swaptions which have working sets no larger than 16 MB. These workloads have a limited need for caches with a bigger capacity, and the latest generation of CMPs often already has caches sufficiently large to accommodate most of their working sets. The second group of workloads is composed of the benchmarks canneal, ferret, facesim, fluidanimate and freqmine. These programs have very large working sets of sizes 65 MB and more, and even with a relatively con-

strained input set such as simlarge, their working sets can reach hundreds of megabytes. Moreover, the need of those workloads for cache capacity is nearly insatiable and grows with the amount of data which they process. In Table 3 we give our estimates for the largest working set of each PAR-SEC workload for the native input set. In several cases they are significantly larger and can even reach gigabytes. These large working sets are often the consequence of an algorithm that operates on large amounts of collected input data. ferret for example keeps a data base of feature vectors of images in memory to find the images most similar to a given query image. The cache and memory needs of these applications should be considered unbounded, as they become more useful to their users if they can work with increased amounts of data. Programs with unbounded working sets are canneal, dedup, ferret and freqmine.

In Figure 4 we present our analysis of the spatial locality of the PARSEC workloads. The data shows how the miss rate of a shared cache changes with line size. All programs benefit from larger cache lines, but to different extents. facesim, fluidanimate and streamcluster show the greatest improvement as the line size is increased, up to the the maximum value of 256 bytes which we used. These programs have streaming behavior, and an increased line size has a prefetching effect which these workloads can take advantage of. facesim for example spends most of its time updating the position-based state of the model, for which it employs an iterative Newton-Raphson algorithm. The algorithm iterates over the elements of a sparse matrix which is stored in two one-dimensional arrays, resulting in a streaming behavior. All other programs also show good improvement of the miss rate with larger cache lines, but only up to line sizes of about 128 bytes. The miss rate is not substantially reduced with larger lines. This is due to a limited size of the basic data structures employed by the programs. They represent independent logical units, each of which is intensely worked with during a computational phase. For example, x264 operates on macroblocks of 8 × 8 pixels at a time, which limits the sizes of the used data structures. Processing a macroblock is computationally intensive and largely independent from other macroblocks. Consequently, the amount of spatial locality is bounded in these cases.

For the rest of our analysis we chose a cache capacity of 4 MB for all experiments. We could have used a matching cache size for each workload, but that would have made comparisons very difficult, and the use of very small or very large cache sizes is not realistic. Moreover, in the case of the workloads with an unbounded working set size, a working set which completely fits into a cache would be an artifact of the limited simulation input size and would not reflect realistic program behavior.

7. COMMUNICATION-TO-COMPUTATION RATIO AND SHARING

In this section we discuss how PARSEC workloads use caches to communicate. Most PARSEC benchmarks share data intensely. Two degrees of sharing can be distinguished: Shared data can be read-only during the parallel phase, in which case it is only used for lookups and analysis. Input data is frequently used in such a way. But shared data can also be used for communication between threads, in which

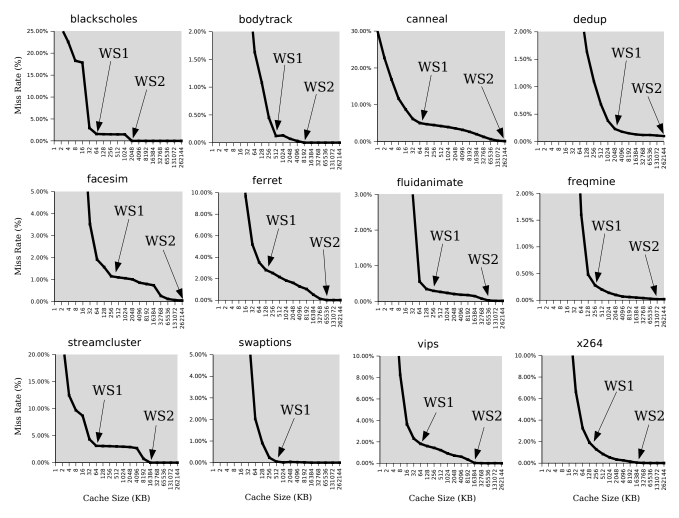


Figure 3: Miss rates versus cache size. Data assumes a shared 4-way associative cache with 64 byte lines. WS1 and WS2 refer to important working sets which we analyze in more detail in Table 3. Cache requirements of PARSEC benchmark programs can reach hundreds of megabytes.

		Input Set					
Program	simlarge						native
	Working Set 1			Working Set 2			Working Set 2
	Data	Size Growth		Data	Size	Growth	Size
	Structure(s)		Rate	Structure(s)		Rate	Estimate
blackscholes	options	64 KB	С	portfolio data	2 MB	С	same
bodytrack	edge maps	512 KB	const.	input frames	8 MB	const.	same
canneal	elements	64 KB	С	netlist	256 MB	DS	2 GB
dedup	data chunks	2 MB	C	hash table	256 MB	DS	2 GB
facesim	tetrahedra	$256~\mathrm{KB}$	C	face mesh	256 MB	DS	same
ferret	images	128 KB	С	data base	64 MB	DS	128 MB
fluidanimate	cells	128 KB	C	particle data	64 MB	DS	128 MB
freqmine	transactions	256 KB	C	FP-tree	128 MB	DS	1 GB
streamcluster	data points	64 KB	C	data block	16 MB	user-def.	256 MB
swaptions	swaptions	512 KB	C	same as WS1	same	same	same
vips	image data	64 KB	C	image data	16 MB	C	same
x264	macroblocks	128 KB	С	reference frames	16 MB	С	same

Table 3: Important working sets and their growth rates. DS represents the data set size and C is the number of cores. Working set sizes are taken from Figure 3. Values for native input set are analytically derived estimates. Working sets that grow proportional to the number of cores C are aggregated private working sets and can be split up to fit into correspondingly smaller, private caches.

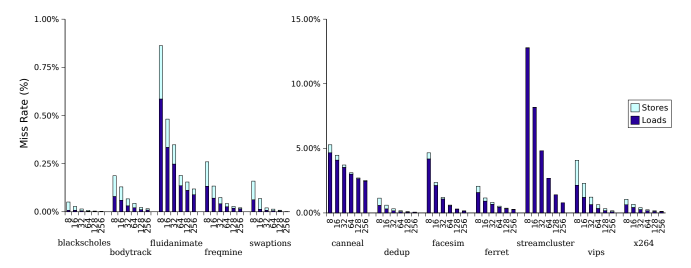


Figure 4: Miss rates as a function of line size. Data assumes 8 cores sharing a 4-way associative cache with 4 MB capacity. Miss rates are broken down to show the effect of loads and stores.

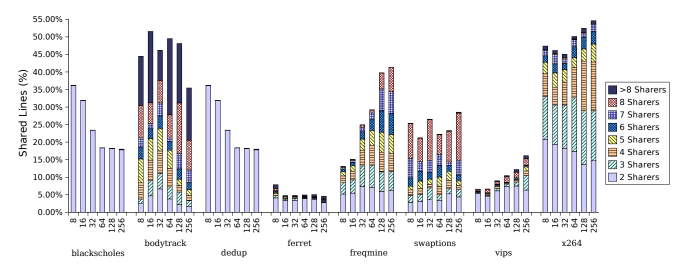


Figure 5: Portion of a 4-way associative cache with 4 MB capacity which is shared. The line size is varied from 8 to 256 bytes. Data assumes 8 cores and is broken down to show the number of threads sharing the lines.

case it is also modified during the parallel phase. In Figure 5we show how the line size affects sharing. The data combines the effects of false sharing and the access pattern of the program due to constrained cache capacity. In Figure 6, we analyze how the program uses its data. The chart shows what data is accessed and how intensely it is used. The information is broken down in two orthogonal ways, resulting in four possible types of accesses: Read and write accesses and accesses to thread-private and shared data. Additionally, we give numbers for true shared accesses. An access is a true access if the last reference to that line came from another thread. True sharing does not count repeated accesses by the same thread. It is a useful metric to estimate the requirements for the cache coherence mechanism of a CMP: A true shared write can trigger a coherence invalidate or update, and a true shared read might require the replication of data. All programs exhibit very few true shared writes. Cache misses are not included in Figure 6, we analyze them

separately when we discuss off-chip traffic in Section 8.

Four programs (canneal, facesim, fluidanimate and streamcluster) showed only trivial amounts of sharing. They have therefore not been included in Figure 5. In the case of canneal, this is a results of the small cache capacity. Most of its large working set is shared and actively worked with by all threads. However, only a minuscule fraction of it fits into the cache, and the probability that a line is accessed by more than one thread before it gets replaced is very small in practice. With a 256 MB cache, 58% of its cached data is shared. blackscholes shows a substantial amount of sharing, but almost all its shared data is only accessed by two threads. This is a side-effect of the parallelization model: At the beginning of the program, the boss threads initializes the portfolio data before it spawns worker threads which process parts of it in a data-parallel way. As such, the entire portfolio is shared between the boss thread and its workers, but

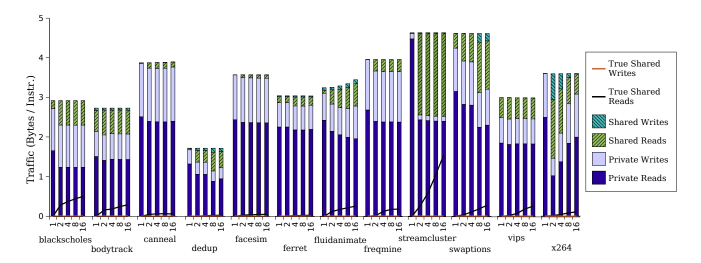


Figure 6: Traffic from cache in bytes per instruction for 1 to 16 cores. Data assumes a shared 4-way associative cache with 64 byte lines. Results are broken down in accesses to private and shared data. True accesses do not count repeated accesses from the same thread.

the worker threads can process the options independently from each other and do not have to communicate with each other. ferret shows a modest amount of data sharing. Like the sharing behavior of canneal, this is caused by severely constrained cache capacity. ferret uses a database that is scanned by all threads to find entries similar to the query image. However, the size of the database is practically unbounded, and because threads do not coordinate their scans with each other it is unlikely that a cache line gets accessed more than once. bodytrack and freqmine exhibit substantial amounts of sharing due to the fact that threads process the same data. The strong increase of sharing of freqmine is caused by false sharing, as the program uses an arraybased tree as its main data structure. Larger cache lines will contain more nodes, increasing the chance that the line is accessed by multiple threads. vips has some shared data which is mostly used by only two threads. This is also predominantly an effect of false sharing since image data is stored in a consecutive array which is processed in a dataparallel way by threads. x264 uses significant amounts of shared data, most of which is only accessed by a low number of threads. This data is the reference frames, since a thread needs this information from other stages in order to encode the frame it was assigned. Similarly, the large amount of shared data of dedup is the input which is passed from stage to stage.

Most PARSEC workloads use a significant amount of communication, and in many cases the volume of traffic between threads can be so high that efficient data exchange via a shared cache is severely constrained by its capacity. An example for this is x264. Figure 6 shows a large amount of writes to shared data, but contrary to intuition its share diminishes rapidly as the number of cores is increased. This effect is caused by a growth of the working sets of x264: Table3 shows that both working set WS1 and WS2 grow proportional to the number of cores. WS1 is mostly composed of thread-private data and is the one which is used more intensely. WS2 contains the reference frames and is used

for inter-thread communication. As WS1 grows, it starts to displace WS2, and the threads are forced to communicate via main memory. Two more programs which communicate intensely are dedup and ferret. Both programs use the pipeline parallelization model with dedicated thread pools for each parallel stage, and all data has to be passed from stage to stage. fluidanimate also shows a large amount of inter-thread communication, and its communication needs grow as the number of threads increase. This is caused by the spatial partitioning that fluidanimate uses to distribute the work to threads. Smaller partitions mean a worse surface to volume ratio, and communication grows with the surface.

Overall, most PARSEC workloads have complex sharing patterns and communicate actively. Pipelined programs can require a large amount of bandwidth between cores in order to communicate efficiently. Shared caches with insufficient capacity can limit the communication efficiency of workloads, since shared data structures might get displaced to memory.

8. OFF-CHIP TRAFFIC

In this section we analyze what the off-chip bandwidth requirements of PARSEC workloads are. Our goal is to understand how the traffic of an application grows as the number of cores of a CMP increases and how the memory wall will limit performance. We again simulated a shared cache and analyze how traffic develops as the number of cores increases. Our results are presented in Figure 7.

The data shows that the off-chip bandwidth requirements of blackscholes are small enough so that memory bandwidth is unlikely to be an issue. bodytrack, dedup, fluidanimate, freqmine, swaptions and x264 are more demanding. Moreover, these programs exhibit a growing bandwidth demand per instruction as the number of cores increases. In the case of bodytrack, most off-chip traffic happens in short, intense bursts since the off-chip communication predominantly

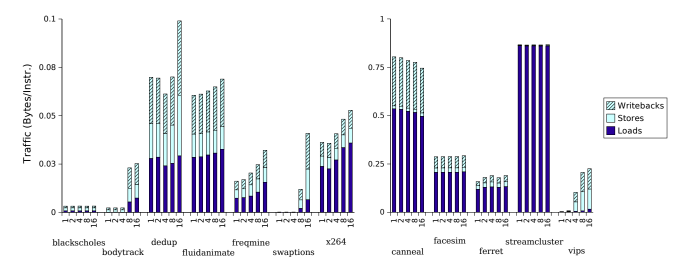


Figure 7: Breakdown of off-chip traffic into bytes for loads, stores and write-backs per instructions. Results are shown for 1 to 16 cores. Data assumes a 4-way associative 4 MB cache with 64 byte lines, allocate-on-store and write-back policy.

takes place during the edge map computation. This phase is only a small part of the serial runtime, but on machines with constrained memory bandwidth it quickly becomes the limiting factor for scalability. The last group of programs is composed of canneal, facesim, ferret, streamcluster and vips. These programs have very high bandwidth requirements and also large working sets. canneal shows a decreasing demand for data per instruction with more cores. This behavior is caused by improved data sharing.

It is important to point out that these numbers do not take the increasing instruction throughput of a CMP into account as its number of cores grows. A constant traffic amount in Figure 7 means that the bandwidth requirements of an application which scales linearly will grow exponentially. Since many PARSEC workloads have high bandwidth requirements and working sets which exceed conventional caches by far, off-chip bandwidth will be their most severe limitation of performance. Substantial architectural improvements are necessary to allow emerging workloads to take full advantage of larger CMPs.

9. CONCLUSIONS

The PARSEC benchmark suite is designed to provide parallel programs for the study for CMPs. It focuses on emerging desktop and server applications and does not have the limitations of other benchmark suites. It is diverse enough to be considered representative, it is not skewed towards HPC programs, it uses state-of-art algorithms and it supports research. In this study we characterized the PARSEC workloads to provide the basic understanding necessary to allow other researchers the effective use of PARSEC for their studies. We analyzed the parallelization, the working sets and locality, the communication-to-computation ratio and the off-chip traffic of its workloads. The high cost of microarchitectural simulation forced us to use smaller machine and problem sizes than we would really like to evaluate.

Our analysis shows that current CMPs are not sufficiently

prepared for the demands of future applications. Many architectural problems still have to be solved to create microprocessors which are ready for the next generation of workloads. Large working sets and high off-chip bandwidth demands might limit the scalability of future programs. PAR-SEC can be used to drive research efforts by application demands.

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