Lecture 20: Domain-Specific Programming Systems

Parallel Computer Architecture and Programming
CMU 15-418/15-618, Spring 2018

Slide acknowledgments:
Pat Hanrahan, Zach Devito (Stanford University)
Jonathan Ragan-Kelley (MIT)
Course themes:

Designing computer systems that **scale**
(running faster given more resources)

Designing computer systems that are **efficient**
(running faster under constraints on resources)

Techniques discussed:

- Exploiting parallelism in applications
- Exploiting locality in applications
- Leveraging hardware specialization (earlier lecture)
Claim: most software uses modern hardware resources inefficiently

- Consider a piece of sequential C code
  - Let’s consider the performance of this code “baseline performance”

- Well-written sequential C code: ~ 5-10x faster
- Assembly language program: another small constant factor faster
- Java, Python, PHP, etc. ??

Credit: Pat Hanrahan


Code performance: relative to C (single core)

Data from: The Computer Language Benchmarks Game: http://shootout.alioth.debian.org

Data from: The Computer Language Benchmarks Game: http://shootout.alioth.debian.org
Even good C code is inefficient

Recall Assignment 1’s Mandelbrot program

Consider execution on a high-end laptop: quad-core, Intel Core i7, AVX instructions...

Single core, with AVX vector instructions: 5.8x speedup over C implementation

Multi-core + hyper-threading + AVX instructions: 21.7x speedup

Conclusion: basic C implementation compiled with -O3 leaves a lot of performance on the table
Making efficient use of modern machines is challenging

(proof by assignments 2, 3, and 4)

In our assignments you only programmed homogeneous parallel computers. And parallelism in that context was not easy.

Assignment 2: GPU cores only

Assignments 3 & 4: shared memory / message passing
Recall: need for efficiency leading to heterogeneous parallel platforms

Integrated CPU + GPU

GPU:
throughput cores + fixed-function

CPU+data-parallel accelerator

Qualcomm Snapdragon SoC 800 PROCESSOR

Mobile system-on-a-chip: CPU+GPU+media processing
Hardware trend: specialization of execution

- **Multiple forms of parallelism**
  - SIMD/vector processing
  - Multi-threading
  - Multi-core
  - Multiple node
  - Multiple server

  - Fine-granularity parallelism: perform same logic on different data
  - Mitigate inefficiencies (stalls) caused by unpredictable data access
  - Varying scales of coarse-granularity parallelism

- **Heterogeneous execution capability**
  - Programmable, latency-centric (e.g., “CPU-like” cores)
  - Programmable, throughput-optimized (e.g., “GPU-like” cores)
  - Fixed-function, application-specific (e.g., image/video/audio processing)

Motivation for parallelism and specialization: maximize compute capability given constraints on chip area, chip energy consumption.

Result: amazingly high compute capability in a wide range of devices!
Hardware diversity is a huge challenge

- Machines with very different performance characteristics

- Even worse: different technologies and performance characteristics within the same machine at different scales
  - **Within a core**: SIMD, multi-threading: fine-granularity sync and communication
  - **Across cores**: coherent shared memory via fast on-chip network
  - **Hybrid CPU+GPU multi-core**: incoherent (potentially) shared memory
  - **Across racks**: distributed memory, multi-stage network
Variety of programming models to abstract HW

- Machines with very different performance characteristics
- Worse: different technologies and performance characteristics within the same machine at different scales
  - Within a core: SIMD, multi-threading: fine grained sync and comm
    - Abstractions: SPMD programming (ISPC, Cuda, OpenCL, Metal, Renderscript)
  - Across cores: coherent shared memory via fast on-chip network
    - Abstractions: OpenMP pragma, Cilk, TBB
  - Hybrid CPU+GPU multi-core: incoherent (potentially) shared memory
    - Abstractions: OpenCL
  - Across racks: distributed memory, multi-stage network
    - Abstractions: message passing (MPI, Go, Spark, Legion, Charm++)

Credit: Pat Hanrahan
This is a huge challenge

- Machines with very different performance characteristics
- Worse: different performance characteristics within the same machine at different scales
- To be efficient, software must be optimized for HW characteristics
  - Difficult even in the case of one level of one machine
  - Combinatorial complexity of optimizations when considering a complex machine, or different machines
  - Loss of software portability
Open computer science question:

How do we enable programmers to productively write software that efficiently uses current and future heterogeneous, parallel machines?
The [magical] ideal parallel programming language

High Performance
(software is scalable and efficient)

Productivity
(ease of development)

Completeness
(applicable to most problems we want to write a program for)

Credit: Pat Hanrahan
Successful programming languages

Here: definition of success = widely used

- High Performance
  (software is scalable and efficient)
- Productivity
  (ease of development)
- Completeness
  (applicable to most problems we want to write a program for)

Credit: Pat Hanrahan
Growing interest in domain-specific programming systems
To realize high performance and productivity: willing to sacrifice completeness

High Performance (software is scalable and efficient)

Domain-specific languages (DSL) and programming frameworks

Productivity (ease of development)

Completeness (applicable to most problems we want to write a program for)

Credit: Pat Hanrahan
Domain-specific programming systems

- **Main idea:** raise level of abstraction for expressing programs

- Introduce high-level programming primitives specific to an application domain
  - **Productive:** intuitive to use, portable across machines, primitives correspond to behaviors frequently used to solve problems in targeted domain
  - **Performant:** system uses domain knowledge to provide efficient, optimized implementation(s)
    - Given a machine: system knows what algorithms to use, parallelization strategies to employ for this domain
    - Optimization goes beyond efficient mapping of software to hardware! The hardware platform itself can be optimized to the abstractions as well

- **Cost:** loss of generality/completeness
Two domain-specific programming examples

1. **Liszt**: for scientific computing on meshes

2. **Halide**: for image processing

What are other domain specific languages? (SQL is another good example)
Example 1:
Lizst: a language for solving PDE’s on meshes

[DeVito et al. Supercomputing 11, SciDac ’11]

Slide credit for this section of lecture:
Pat Hanrahan and Zach Devito (Stanford)

http://liszt.stanford.edu/
What a Liszt program does

A Liszt program is run on a mesh
A Liszt program defines, and compute the value of, fields defined on the mesh

Position is a field defined at each mesh vertex. The field’s value is represented by a 3-vector.

```scala
val Position = FieldWithConst[Vertex, Float3](0.f, 0.f, 0.f)
val Temperature = FieldWithConst[Vertex, Float](0.f)
val Flux = FieldWithConst[Vertex, Float](0.f)
val JacobiStep = FieldWithConst[Vertex, Float](0.f)
```

Color key:
- **Fields**
- **Mesh entity**

Notes:
Fields are a higher-kind type (special function that maps a type to a new type)
Liszt program: heat conduction on mesh

Program computes the value of fields defined on meshes

```liszt
var i = 0;
while ( i < 1000 ) {
    Flux(vertices(mesh)) = 0.f;
    JacobiStep(vertices(mesh)) = 0.f;
    for (e <- edges(mesh)) {
        val v1 = head(e)
        val v2 = tail(e)
        val dP = Position(v1) - Position(v2)
        val dT = Temperature(v1) - Temperature(v2)
        val step = 1.0f/(length(dP))
        Flux(v1) += dT*step
        Flux(v2) -= dT*step
        JacobiStep(v1) += step
        JacobiStep(v2) += step
    }
    i += 1
}
```

Given edge, loop body accesses/modifies field values at adjacent mesh vertices

Access value of field at mesh vertex v2

Color key:
- **Fields**
- **Mesh**
- **Topology functions**
- **Iteration over set**
Liszt’s topological operators

Used to access mesh elements relative to some input vertex, edge, face, etc. Topological operators are the only way to access mesh data in a Liszt program. Notice how many operators return sets (e.g., “all edges of this face”)

```plaintext
BoundarySet\textsuperscript{1}[ME <: MeshElement](name : String) : Set[ME]
vertices(e : Mesh) : Set[Vertex]
cells(e : Mesh) : Set[Cell]
edges(e : Mesh) : Set[Edge]
faces(e : Mesh) : Set[Face]

vertices(e : Vertex) : Set[Vertex]
cells(e : Vertex) : Set[Cell]
edges(e : Vertex) : Set[Edge]
faces(e : Vertex) : Set[Face]

vertices(e : Edge) : Set[Vertex]
facesCCW\textsuperscript{2}(e : Edge) : Set[Face]
cells(e : Edge) : Set[Cell]
head(e : Edge) : Vertex
tail(e : Edge) : Vertex
flip\textsuperscript{4}(e : Edge) : Edge
towards\textsuperscript{5}(e : Edge, t : Vertex) : Edge

vertices(e : Cell) : Set[Cell]
faces(e : Cell) : Set[Face]
edges(e : Cell) : Set[Edge]
cells(e : Face) : Set[Cell]
edgesCCW\textsuperscript{2}(e : Face) : Set[Edge]
faces(e : Face) : Set[Vertex]
inside\textsuperscript{3}(e : Face) : Cell
outside\textsuperscript{3}(e : Face) : Cell
flip\textsuperscript{4}(e : Face) : Face
towards\textsuperscript{5}(e : Face, t : Cell) : Face
```
Liszt programming

- A Liszt program describes operations on fields of an abstract mesh representation
- Application specifies type of mesh (regular, irregular) and its topology
- Mesh representation is chosen by Liszt (not by the programmer)

Well, that’s interesting. I write a program, and the compiler decides what data structure it should use based on what operations my code performs.
Compiling to parallel computers

Recall challenges you have faced in your assignments

1. Identify parallelism
2. Identify data locality
3. Reason about required synchronization

Now consider how to automate this process in the Liszt compiler.
Key: determining program dependencies

1. Identify **parallelism**
   - Absence of dependencies implies code can be executed in parallel

2. Identify **data locality**
   - Partition data based on dependencies (localize dependent computations for faster synchronization)

3. Reason about required **synchronization**
   - Synchronization is needed to respect dependencies (must wait until the values a computation depends on are known)

In general programs, compilers are unable to infer dependencies at global scale: \( a[f(i)] += b[i] \) (must execute \( f(i) \) to know if dependency exists across loop iterations \( i \))
Liszt is constrained to allow dependency analysis

Liszt infers “stencils”: “stencil” = mesh elements accessed in an iteration of loop
= dependencies for the iteration

Statically analyze code to find stencil of each top-level for loop
- Extract nested mesh element reads
- Extract field operations

```scala
for (e <- edges(mesh)) {
  val v1 = head(e)
  val v2 = tail(e)
  val dP = Position(v1) - Position(v2)
  val dT = Temperature(v1) - Temperature(v2)
  val step = 1.0f/(length(dP))
  Flux(v1) += dT*step
  Flux(v2) -= dT*step
  JacobiStep(v1) += step
  JacobiStep(v2) += step
}
```

Edge 6’s read stencil is D and F
Restrict language for dependency analysis

Language restrictions:

- Mesh elements are only accessed through built-in topological functions:
  
  \texttt{cells(mesh)}, \ldots

- Single static assignment:
  
  \texttt{val v1 = head(e)}

- Data in fields can only be accessed using mesh elements:
  
  \texttt{Pressure(v)}

- No recursive functions

Restrictions allow compiler to automatically infer stencil for a loop iteration.
Portable parallelism: use dependencies to implement different parallel execution strategies

I’ll discuss two strategies…

Strategy 1: mesh partitioning

Strategy 2: mesh coloring
Imagine compiling a Lizst program to the (entire) Latedays cluster

(multiple nodes, distributed address space)

How might Liszt distribute a graph across these nodes?
Distributed memory implementation of Liszt

Mesh + Stencil → Graph → Partition

```
for(f <- faces(mesh)) {
  rhoOutside(f) =
    calc_flux(f, rho(outside(f))) +
    calc_flux(f, rho(inside(f)))
}
```

Consider distributed memory implementation
Store region of mesh on each node in a cluster
(Note: ParMETIS is a tool for partitioning meshes)
Each processor also needs data for neighboring cells to perform computation ("ghost cells"). Listz allocates ghost region storage and emits required communication to implement topological operators.
Imagine compiling a Lizst program to a GPU (single address space, many tiny threads)
GPU implementation: parallel reductions

In previous example, one region of mesh assigned per processor (or node in MPI cluster)
On GPU, natural parallelization is one edge per CUDA thread

Threads (each edge assigned to 1 CUDA thread)

<table>
<thead>
<tr>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>B</td>
<td>C</td>
<td>D</td>
<td></td>
<td>E</td>
<td>F</td>
<td>G</td>
<td>H</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Flux field values (per vertex)

```java
for (e <- edges(mesh)) {
  ...
  Flux(v1) += dT*step
  Flux(v2) -= dT*step
  ...
}
```

Different edges share a vertex: requires atomic update of per-vertex field data
GPU implementation: conflict graph

Threads (each edge assigned to 1 CUDA thread)

Flux field values (per vertex)

Identify mesh edges with colliding writes (lines in graph indicate presence of collision)

Can simply run program once to get this information.
(results valid for subsequent executions provided mesh does not change)
GPU implementation: conflict graph

Threads (each edge assigned to 1 CUDA thread)

Flux field values (per vertex)

“Color” nodes in graph such that no connected nodes have the same color

Can execute on GPU in parallel, without atomic operations, by running all nodes with the same color in a single CUDA launch.
Cluster performance of Lizst program

256 nodes, 8 cores per node (message-passing implemented using MPI)

Important: performance portability!
Same Liszt program also runs with high efficiency on GPU (results not shown here).
But uses a different algorithm when compiled to GPU! (graph coloring)
Liszt summary

- **Productivity:**
  - Abstract representation of mesh: vertices, edges, faces, fields (concepts that a scientist thinks about already!)
  - Intuitive topological operators

- **Portability**
  - Same code runs on large cluster of CPUs (MPI) and GPUs (and combinations thereof!)

- **High-performance**
  - Language is constrained to allow compiler to track dependencies
  - Used for locality-aware partitioning in distributed memory implementation
  - Used for graph coloring in GPU implementation
  - Compiler knows how to choose different parallelization strategies for different platforms
  - Underlying mesh representation can be customized by system based on usage and platform (e.g., don’t store edge pointers if code doesn’t need it, choose struct of arrays vs. array of structs for per-vertex fields)
Example 2:
Halide: a domain-specific language for image processing

Jonathan Ragan-Kelley, Andrew Adams et al.
[SIGGRAPH 2012, PLDI 13]
Halide used in practice

- Halide used to implement Android HDR+ app
- Halide code used to process all images uploaded to Google Photos
A quick tutorial on high-performance image processing
What does this C code do?

```c
int WIDTH = 1024;
int HEIGHT = 1024;
float input[(WIDTH+2) * (HEIGHT+2)];
float output[WIDTH * HEIGHT];

float weights[] = {1.0/9, 1.0/9, 1.0/9,
                   1.0/9, 1.0/9, 1.0/9,
                   1.0/9, 1.0/9, 1.0/9};

for (int j=0; j<HEIGHT; j++) {
    for (int i=0; i<WIDTH; i++) {
        float tmp = 0.f;
        for (int jj=0; jj<3; jj++)
            for (int ii=0; ii<3; ii++)
                tmp += input[(j+jj)*(WIDTH+2) + (i+ii)] * weights[jj*3 + ii];
        output[j*WIDTH + i] = tmp;
    }
}
```
3x3 box blur

(Zoom view)
3x3 image blur

```c
int WIDTH = 1024;
int HEIGHT = 1024;
float input[(WIDTH+2) * (HEIGHT+2)];
float output[WIDTH * HEIGHT];

float weights[] = {1.0/9, 1.0/9, 1.0/9,
                   1.0/9, 1.0/9, 1.0/9,
                   1.0/9, 1.0/9, 1.0/9};

for (int j=0; j<HEIGHT; j++) {
    for (int i=0; i<WIDTH; i++) {
        float tmp = 0.f;
        for (int jj=0; jj<3; jj++)
            for (int ii=0; ii<3; ii++)
                tmp += input[(j+jj)*(WIDTH+2) + (i+ii)] * weights[jj*3 + ii];
        output[j*WIDTH + i] = tmp;
    }
}
```

Total work per image = $9 \times WIDTH \times HEIGHT$

For N\times N filter: $N^2 \times WIDTH \times HEIGHT$
Two-pass 3x3 blur

int WIDTH = 1024;
int HEIGHT = 1024;
float input[(WIDTH+2) * (HEIGHT+2)];
float tmp_buf[WIDTH * (HEIGHT+2)];
float output[WIDTH * HEIGHT];

float weights[] = {1.0/3, 1.0/3, 1.0/3};

for (int j=0; j<(HEIGHT+2); j++)
    for (int i=0; i<WIDTH; i++) {
        float tmp = 0.f;
        for (int ii=0; ii<3; ii++)
            tmp += input[j*(WIDTH+2) + i+ii] * weights[ii];
        tmp_buf[j*WIDTH + i] = tmp;
    }

for (int j=0; j<HEIGHT; j++) {
    for (int i=0; i<WIDTH; i++) {
        float tmp = 0.f;
        for (int jj=0; jj<3; jj++)
            tmp += tmp_buf[(j+jj)*WIDTH + i] * weights[jj];
        output[j*WIDTH + i] = tmp;
    }
}
### Two-pass image blur: locality

```c
int WIDTH = 1024;
int HEIGHT = 1024;
float input[(WIDTH+2) * (HEIGHT+2)];
float tmp_buf[WIDTH * (HEIGHT+2)];
float output[WIDTH * HEIGHT];

float weights[] = {1.0/3, 1.0/3, 1.0/3};

for (int j=0; j<(HEIGHT+2); j++)
    for (int i=0; i<WIDTH; i++) {
        float tmp = 0.f;
        for (int ii=0; ii<3; ii++)
            tmp += input[j*(WIDTH+2) + i+ii] * weights[ii];
        tmp_buf[j*WIDTH + i] = tmp;
    }

for (int j=0; j<HEIGHT; j++) {
    for (int i=0; i<WIDTH; i++) {
        float tmp = 0.f;
        for (int jj=0; jj<3; jj++)
            tmp += tmp_buf[(j+jj)*WIDTH + i] * weights[jj];
        output[j*WIDTH + i] = tmp;
    }
}
```

### Intrinsic bandwidth requirements of algorithm:
Application must read each element of input image and must write each element of output image.

Data from `input` reused three times. (immediately reused in next two i-loop iterations after first load, never loaded again.)
- Perfect cache behavior: never load required data more than once
- Perfect use of cache lines (don’t load unnecessary data into cache)

Data from `tmp_buf` reused three times (but three rows of image data are accessed in between)
- Never load required data more than once... if cache has capacity for three rows of image
- Perfect use of cache lines (don’t load unnecessary data into cache)

Two pass: loads/stores to `tmp_buf` are overhead (this memory traffic is an artifact of the two-pass implementation: it is not intrinsic to computation being performed)
Two-pass image blur, “chunked” (version 1)

```c
int WIDTH = 1024;
int HEIGHT = 1024;
float input[(WIDTH+2) * (HEIGHT+2)];
float tmp_buf[WIDTH * 3];
float output[WIDTH * HEIGHT];
float weights[] = {1.0/3, 1.0/3, 1.0/3};

for (int j=0; j<HEIGHT; j++) {
    for (int j2=0; j2<3; j2++)
        for (int i=0; i<WIDTH; i++) {
            float tmp = 0.f;
            for (int ii=0; ii<3; ii++)
                tmp += input[(j+j2)*(WIDTH+2) + i+ii] * weights[ii];
            tmp_buf[j2*WIDTH + i] = tmp;
        }
    for (int i=0; i<WIDTH; i++) {
        float tmp = 0.f;
        for (int jj=0; jj<3; jj++)
            tmp += tmp_buf[jj*WIDTH + i] * weights[jj];
        output[j*WIDTH + i] = tmp;
    }
}
```

Only 3 rows of intermediate buffer need to be allocated

Produce 3 rows of tmp_buf (only what’s needed for one row of output)

Combine them together to get one row of output

Total work per row of output:
- step 1: 3 x 3 x WIDTH work
- step 2: 3 x WIDTH work

Total work per image = 12 x WIDTH x HEIGHT

Loads from tmp_buffer are cached (assuming tmp_buffer fits in cache)
Two-pass image blur, “chunked” (version 2)

```c
int WIDTH = 1024;
int HEIGHT = 1024;
float input[(WIDTH+2) * (HEIGHT+2)];
float tmp_buf[WIDTH * (CHUNK_SIZE+2)];
float output[WIDTH * HEIGHT];
float weights[] = {1.0/3, 1.0/3, 1.0/3};

for (int j=0; j<HEIGHT; j+CHUNK_SIZE) {
    for (int j2=0; j2<CHUNK_SIZE+2; j2++)
        for (int i=0; i<WIDTH; i++) {
            float tmp = 0.f;
            for (int ii=0; ii<3; ii++)
                tmp += input[(j+j2)*(WIDTH+2) + i+ii] * weights[ii];
            tmp_buf[j2*WIDTH + i] = tmp;
        }
    for (int j2=0; j2<CHUNK_SIZE; j2++)
        for (int i=0; i<WIDTH; i++) {
            float tmp = 0.f;
            for (int jj=0; jj<3; jj++)
                tmp += tmp_buf[(j2+jj)*WIDTH + i] * weights[jj];
            output[(j+j2)*WIDTH + i] = tmp;
        }
}
```

Sized to fit in cache (capture all producer-consumer locality)

Produce enough rows of tmp_buf to produce a CHUNK_SIZE number of rows of output

Produce CHUNK_SIZE rows of output

Total work per chunk of output:
(assume CHUNK_SIZE = 16)
- Step 1: 18 x 3 x WIDTH work
- Step 2: 16 x 3 x WIDTH work
Total work per image: \( \frac{34}{16} \times 3 \times WIDTH \times HEIGHT \)

= 6.4 \times WIDTH \times HEIGHT

Trends to idea 6 x WIDTH x HEIGHT as CHUNK_SIZE is increased!

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Conflicting goals (once again...)

- Want to be work efficient (perform fewer operations)
- Want to take advantage of locality when present
  - Otherwise work-efficient code will be bandwidth bound
  - Ideally: bandwidth cost of implementation is very close to intrinsic cost of algorithm: data is loaded from memory once and reused as much as needed prior to being discarded from processor’s cache
- Want to execute in parallel (multi-core, SIMD within core)
Optimized C++ code: 3x3 image blur

Good: 10x faster: on a quad-core CPU than my original two-pass code
Bad: specific to SSE (not AVX2), CPU-code only, hard to tell what is going on at all!

```c++
void fast_blur(const Image &in, Image &blurred) {
    __m128i one_third = _mm_set1_epi16(21846);
    #pragma omp parallel for
    for (int yTile = 0; yTile < in.height(); yTile += 32) {
        __m128i a, b, c, sum, avg;
        __m128i tmp[(256/8)*(32+2)];
        for (int xTile = 0; xTile < in.width(); xTile += 256) {
            __m128i *tmpPtr = tmp;
            for (int y = -1; y < 32+1; y++) {
                const uint16_t *inPtr = &in(xTile, yTile+y);
                for (int x = 0; x < 256; x += 8) {
                    a = _mm_loadu_si128((__m128i*)(inPtr-1));
                    b = _mm_loadu_si128((__m128i*)(inPtr+1));
                    c = _mm_load_si128((__m128i*)(inPtr));
                    sum = _mm_add_epi16(_mm_add_epi16(a, b), c);
                    avg = _mm_mulhi_epi16(sum, one_third);
                    _mm_store_si128(tmpPtr++, avg);
                    inPtr += 8;
                }
            }
            tmpPtr = tmp;
            for (int y = 0; y < 32; y++) {
                __m128i *outPtr = (__m128i *)&blurred(xTile, yTile+y));
                for (int x = 0; x < 256; x += 8) {
                    a = _mm_load_si128(tmpPtr+(2*256)/8);
                    b = _mm_load_si128(tmpPtr+256/8);
                    c = _mm_load_si128(tmpPtr++);
                    sum = _mm_add_epi16(_mm_add_epi16(a, b), c);
                    avg = _mm_mulhi_epi16(sum, one_third);
                    _mm_store_si128(outPtr++, avg);
                }
            }
        }
    }
}
```
Halide blur (algorithm description)

// Halide 3x3 blur program definition
Func halide_blur(Func in) {
    Func blurx, out;
    Var x, y;

    blurx(x,y) = (in(x-1, y) + in(x,y) + in(x+1,y)) / 3.0f;
    out(x,y)   = (blurx(x,y-1) + blurx(x,y) + blurx(x,y+1)) / 3.0f;
    return out;
}

// top-level calling code
Image<uint8_t> input = load_image("myimage.png"); // define input image
Func my_program = halide_blur(input); // define pipeline
Image<uint8_t> output = my_program.realize(input.width(), input.height(), input.channels()); // execute pipeline
output.save("myblurredimage.png");

NOTE: execution order and storage are unspecified by the abstraction. The implementation can evaluate, reevaluate, cache individual points as desired!
Think of a Halide program as a pipeline

// Halide 3x3 blur program definition
Func halide_blur(Func in) {

    Func blurx, out;
    Var  x, y;

    blurx(x,y) = (in(x-1, y) + in(x,y) + in(x+1,y)) / 3.0f;
    out(x,y)   = (blurx(x,y-1) + blurx(x,y) + blurx(x,y+1)) / 3.0f;
    return out;
}

Halide schedule describes how to execute a pipeline

// Halide program definition
Func halide_blur(Func in) {

    Func blurx, out;
    Var x, y, xi, yi

    // the "algorithm description" (what to do)
    blurx(x,y) = (in(x-1, y) + in(x,y) + in(x+1,y)) / 3.0f;
    out(x,y) = (blurx(x,y-1) + blurx(x,y) + blurx(x,y+1)) / 3.0f;

    // "the schedule" (how to do it)
    out.tile(x, y, xi, yi, 256, 32).vectorize(xi,8).parallel(y);
    blurx.chunk(x).vectorize(x, 8);

    return out;
}
Given a schedule, Halide carries out mechanical process of implementing the specified schedule.
Halide: two domain-specific co-languages

- **Functional language** for describing image processing operations
- **Domain-specific language** for describing schedules

**Design principle:** separate “algorithm specification” from its schedule
- Programmer’s responsibility: provide a high-performance schedule
- Compiler’s responsibility: carry out mechanical process of generating threads, SIMD instructions, managing buffers, etc.
- **Result:** enable programmer to rapidly explore space of schedules
  - (e.g., “tile these loops”, “vectorize this loop”, “parallelize this loop across cores”)

**Domain scope:**
- All computation on regular N-D coordinate spaces
- Only feed-forward pipelines (includes special support for reductions and fixed recursion depth)
- All dependencies inferable by compiler
Producer/consumer scheduling primitives

Four basic scheduling primitives shown below

**Breadth First**: each function is entirely evaluated before the next one.

**In-line**: values are computed on the fly each time that they are needed.

**Sliding Window**: values are computed when needed then stored until not useful anymore.

**Tiles**: overlapping regions are processed in parallel, functions are evaluated one after another.
Producer/consumer scheduling primitives

```
// Halide program definition
Func halide_blur(Func in) {
  Func blurx, out;
  Var   x, y, xi, yi

  // the "algorithm description" (what to do)
  blurx(x,y) = (in(x-1, y) + in(x,y) + in(x+1,y)) / 3.0f;
  out(x,y) = (blurx(x,y-1) + blurx(x,y) + blurx(x,y+1)) / 3.0f;

  // "the schedule" (how to do it)
  blurx.compute_at(ROOT);
  return out;
}

// Halide program definition
void halide_blur(uint8_t* in, uint8_t* out) {
  uint8_t blurx[WIDTH * HEIGHT];
  for (int y=0; y<HEIGHT; y++) {
    for (int x=0; y<WIDTH; x++) {
      blurx[] = ...
    }
  }
}
```

"Root":
compute all points of the producer,
then run consumer (minimal locality)

```
void halide_blur(uint8_t* in, uint8_t* out) {
  for (int y=0; y<HEIGHT; y++) {
    for (int x=0; y<WIDTH; x++) {
      out[] = (((in[(y-1)*WIDTH+x-1] +
               in[(y-1)*WIDTH+x] +
               in[(y-1)*WIDTH+x+1]) / 3) +
               ((in[y*WIDTH+x-1] +
                 in[y*WIDTH+x] +
                 in[y*WIDTH+x+1]) / 3) +
               ((in[(y+1)*WIDTH+x-1] +
                 in[(y+1)*WIDTH+x] +
                 in[(y+1)*WIDTH+x+1]) / 3));
    }
  }
```

"Inline":
revaluate producer at every use site
in consumer (maximal locality)

```
void halide_blur(uint8_t* in, uint8_t* out) {
  for (int y=0; y<HEIGHT; y++) {
    for (int x=0; y<WIDTH; x++) {
      out[] = ...
    }
  }
```

CMU 15-418/618, Spring 2018
Domain iteration primitives

Specify both order and how to parallelize (multi-thread, SIMD vector)

2D blocked iteration order

serial y, serial x

serial x, serial y

serial y, vectorized x

parallel y, vectorized x

split x into \(2x_o + x_i\)

split y into \(2y_o + y_i\)

serial \(y_o, x_o, y_i, x_i\)
Example Halide results

- Camera RAW processing pipeline
  (Convert RAW sensor data to RGB image)
  - Original: 463 lines of hand-tuned ARM NEON assembly
  - Halide: 2.75x less code, 5% faster

- Bilateral filter
  (Common image filtering operation used in many applications)
  - Original 122 lines of C++
  - Halide: 34 lines algorithm + 6 lines schedule
    - CPU implementation: 5.9x faster
    - GPU implementation: 2x faster than hand-written CUDA
Stepping back: what is Halide?

- **Halide** is a DSL for **helping good developers optimize image processing code more rapidly**
  - Halide doesn’t decide how to optimize a program for a novice programmer
  - Halide provides primitives for a programmer (that has strong knowledge of code optimization, such as a 418 student) to rapidly express what optimizations the system should apply
  - Halide carries out the nitty-gritty of mapping that strategy to a machine
Automatically generating Halide schedules

[CMU 15-418/618, Spring 2018]

Extend Halide compiler to automatically generate schedule for programmer
- Compiler input: Halide program + size of expected input/output images

<table>
<thead>
<tr>
<th>Function</th>
<th>1 thread</th>
<th>6 threads</th>
<th>12 threads</th>
</tr>
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<tbody>
<tr>
<td>BLUR</td>
<td><img src="image1" alt="Graph" /></td>
<td><img src="image2" alt="Graph" /></td>
<td><img src="image3" alt="Graph" /></td>
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<tr>
<td>UNSHARP</td>
<td><img src="image4" alt="Graph" /></td>
<td><img src="image5" alt="Graph" /></td>
<td><img src="image6" alt="Graph" /></td>
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<td>HARRIS</td>
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<td><img src="image41" alt="Graph" /></td>
<td><img src="image42" alt="Graph" /></td>
</tr>
</tbody>
</table>

- Naive schedule
- Expert manual schedule
- Automatically generated schedule (no autotuning, ~ seconds)
- Automatically generated, with auto-tuning (~ 10 minutes)
- Automatically generated, auto-tuning over 3 days

[Mullapudi, CMU 2016]
“Racing” top Halide programmers

Halide auto-scheduler produced schedules that were better than those of expert Google Halide programmers in two of three cases (it got beat in one!)
Darkroom/Rigel

- Directly synthesize FPGA implementation of image processing pipeline from a high-level description (a constrained “Halide-like” language)

```
bx = im(x,y) 
(I(x-1,y) + 
I(x,y) + 
I(x+1,y))/3
end
by = im(x,y) 
(bx(x,y-1) + 
bx(x,y) + 
bx(x,y+1))/3
end
sharpened = im(x,y) 
I(x,y) + 0.1* 
(I(x,y) - by(x,y))
end
```

Stencil Language

- Goal: ultra high efficiency image processing
Many other recent domain-specific programming systems

**Hadoop**

Less domain specific than examples given today, but still designed specifically for:
data-parallel computations on big data for distributed systems ("Map-Reduce")

**GraphLab**

DSL for graph-based machine learning computations

Also see Green-Marl, Ligra
(DSLs for describing operations on graphs)

**Rails**

Model-view-controller paradigm for web-applications

**Ongoing efforts in many domains...**

Simit: a language for physical simulation [MIT]
Domain-specific programming system development

- Can develop DSL as a stand-alone language
  - Graphics shading languages
  - MATLAB, SQL

- “Embed” DSL in an existing generic language
  - e.g., C++ library (GraphLab, OpenGL host-side API, Map-Reduce)
  - Lizst syntax above was all valid Scala code

- Active research idea:
  - Design generic languages that have facilities that assist rapid embedding of new domain-specific languages
  - “What is a good language for rapidly making new DSLs?”
Summary

- **Modern machines: parallel and heterogeneous**
  - Only way to increase compute capability in energy-constrained world

- **Most software uses small fraction of peak capability of machine**
  - Very challenging to tune programs to these machines
  - Tuning efforts are not portable across machines

- **Domain-specific programming environments trade-off generality to achieve productivity, performance, and portability**
  - Case studies today: Liszt, Halide
  - Common trait: languages provide abstractions that make dependencies known
    - Understanding dependencies is necessary but not sufficient: need domain restrictions and domain knowledge for system to synthesize efficient implementations