



Chapel Overview



COMPUTE | STORE | ANALYZE



Safe Harbor Statement

This presentation may contain forward-looking statements that are based on our current expectations. Forward looking statements may include statements about our financial guidance and expected operating results, our opportunities and future potential, our product development and new product introduction plans, our ability to expand and penetrate our addressable markets and other statements that are not historical facts. These statements are only predictions and actual results may materially vary from those projected. Please refer to Cray's documents filed with the SEC from time to time concerning factors that could affect the Company and these forward-looking statements.





Motivation for Chapel

Q: Can a single language be...

...as productive as Python?

...as fast as Fortran?

...as portable as C?

...as scalable as MPI?

...as fun as <your favorite language here>?

A: We believe so.





The Challenge

Q: So why don't we have such languages already?

~~**A:** Technical challenges?~~

- while they exist, we don't think this is the main issue...

A: Due to a lack of...

- ...long-term efforts
- ...resources
- ...community will
- ...co-design between developers and users
- ...patience

Chapel is our attempt to reverse this trend



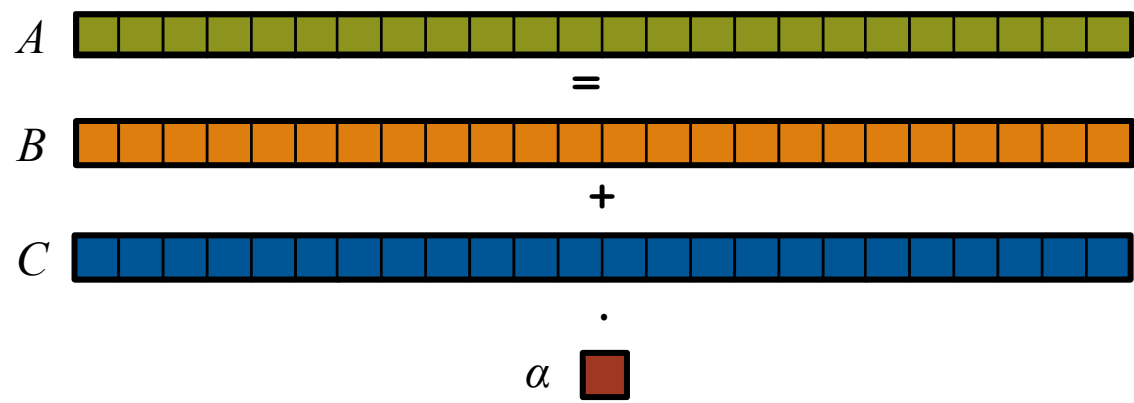


STREAM Triad: a trivial parallel computation

Given: m -element vectors A, B, C

Compute: $\forall i \in 1..m, A_i = B_i + \alpha \cdot C_i$

In pictures:

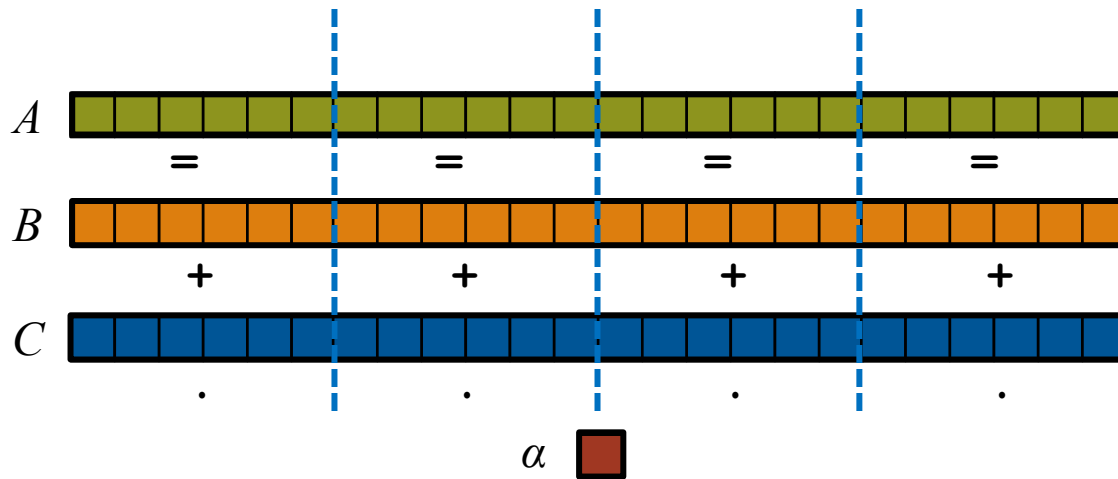


STREAM Triad: a trivial parallel computation

Given: m -element vectors A, B, C

Compute: $\forall i \in 1..m, A_i = B_i + \alpha \cdot C_i$

In pictures, in parallel:

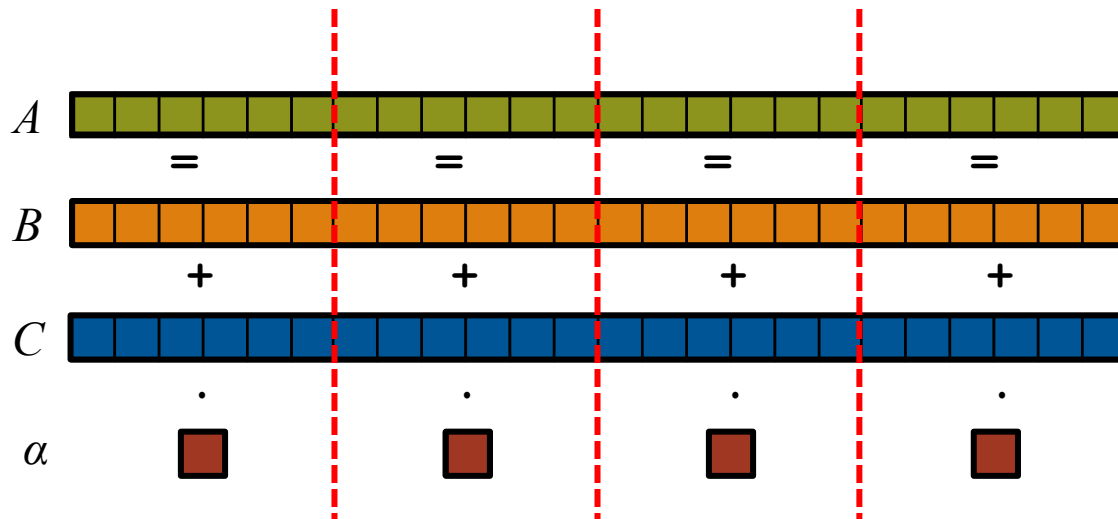


STREAM Triad: a trivial parallel computation

Given: m -element vectors A, B, C

Compute: $\forall i \in 1..m, A_i = B_i + \alpha \cdot C_i$

In pictures, in parallel (distributed memory):

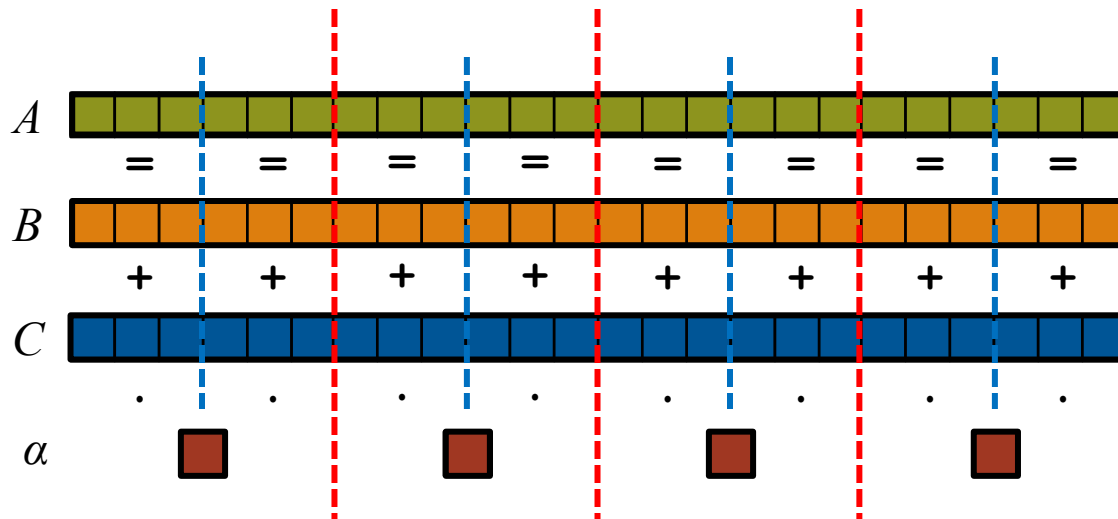


STREAM Triad: a trivial parallel computation

Given: m -element vectors A, B, C

Compute: $\forall i \in 1..m, A_i = B_i + \alpha \cdot C_i$

In pictures, in parallel (distributed memory multicore):



STREAM Triad: MPI



MPI

```
#include <hpcc.h>
```

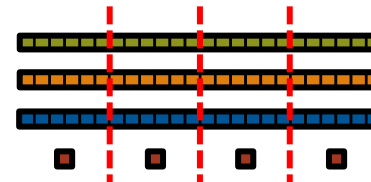
```
static int VectorSize;  
static double *a, *b, *c;
```

```
int HPCC_StarStream(HPCC_Params *params) {  
    int myRank, commSize;  
    int rv, errCount;  
    MPI_Comm comm = MPI_COMM_WORLD;  
  
    MPI_Comm_size( comm, &commSize );  
    MPI_Comm_rank( comm, &myRank );  
  
    rv = HPCC_Stream( params, 0 == myRank );  
    MPI_Reduce( &rv, &errCount, 1, MPI_INT, MPI_SUM,  
        0, comm );  
  
    return errCount;  
}
```

```
int HPCC_Stream(HPCC_Params *params, int doIO) {  
    register int j;  
    double scalar;
```

```
    VectorSize = HPCC_LocalVectorSize( params, 3,  
        sizeof(double), 0 );
```

```
    a = HPCC_XMALLOC( double, VectorSize );  
    b = HPCC_XMALLOC( double, VectorSize );  
    c = HPCC_XMALLOC( double, VectorSize );
```



```
    if (!a || !b || !c) {  
        if (c) HPCC_free(c);  
        if (b) HPCC_free(b);  
        if (a) HPCC_free(a);  
        if (doIO) {  
            fprintf( outFile, "Failed to allocate memory  
                (%d).\n", VectorSize );  
            fclose( outFile );  
        }  
        return 1;  
    }
```

```
    for (j=0; j<VectorSize; j++) {  
        b[j] = 2.0;  
        c[j] = 1.0;  
    }
```

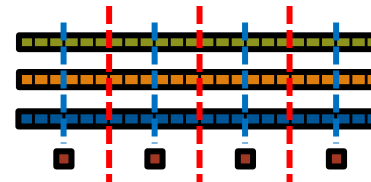
```
    scalar = 3.0;
```

```
    for (j=0; j<VectorSize; j++)  
        a[j] = b[j]+scalar*c[j];
```

```
    HPCC_free(c);  
    HPCC_free(b);  
    HPCC_free(a);
```



STREAM Triad: MPI+OpenMP



MPI + OpenMP

```
#include <hpcc.h>
#ifdef _OPENMP
#include <omp.h>
#endif

static int VectorSize;
static double *a, *b, *c;

int HPCC_StarStream(HPCC_Params *params) {
    int myRank, commSize;
    int rv, errCount;
    MPI_Comm comm = MPI_COMM_WORLD;

    MPI_Comm_size( comm, &commSize );
    MPI_Comm_rank( comm, &myRank );

    rv = HPCC_Stream( params, 0 == myRank );
    MPI_Reduce( &rv, &errCount, 1, MPI_INT, MPI_SUM,
        0, comm );

    return errCount;
}

int HPCC_Stream(HPCC_Params *params, int doIO) {
    register int j;
    double scalar;

    VectorSize = HPCC_LocalVectorSize( params, 3,
        sizeof(double), 0 );

    a = HPCC_XMALLOC( double, VectorSize );
    b = HPCC_XMALLOC( double, VectorSize );
    c = HPCC_XMALLOC( double, VectorSize );
```

```
    if (!a || !b || !c) {
        if (c) HPCC_free(c);
        if (b) HPCC_free(b);
        if (a) HPCC_free(a);
        if (doIO) {
            fprintf( outFile, "Failed to allocate memory
(%d).\n", VectorSize );
            fclose( outFile );
        }
        return 1;
    }

#ifdef _OPENMP
#pragma omp parallel for
#endif
    for (j=0; j<VectorSize; j++) {
        b[j] = 2.0;
        c[j] = 1.0;
    }

    scalar = 3.0;

#ifdef _OPENMP
#pragma omp parallel for
#endif
    for (j=0; j<VectorSize; j++)
        a[j] = b[j]+scalar*c[j];

    HPCC_free(c);
    HPCC_free(b);
    HPCC_free(a);
```



STREAM Triad: MPI+OpenMP vs. CUDA

MPI + OpenMP

```
#ifdef _OPENMP
#include <omp.h>
#endif

static int VectorSize;
static double *a, *b, *c;

int HPCC_StarStream(HPCC_Params *params) {
    int myRank, commSize;
    int rv, errCount;
    MPI_Comm comm = MPI_COMM_WORLD;

    MPI_Comm_size(comm, &commSize);
    MPI_Comm_rank(comm, &myRank);

    rv = HPCC_Stream(params, 0 == myRank);
    MPI_Reduce(&rv, &errCount, 1, MPI_INT, MPI_SUM, 0, comm);

    return errCount;
}

int HPCC_Stream(HPCC_Params *params, int myRank) {
    double scalar;

    VectorSize = HPCC_LocalVectorSize(params, 3, sizeof(double), 0);

    a = HPCC_XMALLOC(double, VectorSize);
    b = HPCC_XMALLOC(double, VectorSize);
    c = HPCC_XMALLOC(double, VectorSize);

    if (!a || !b || !c) {
        if (c) HPCC_free(c);
        if (b) HPCC_free(b);
        if (a) HPCC_free(a);
        if (doIO) {
            fprintf(outFile, "Failed to allocate memory (%d).\n", VectorSize);
            fclose(outFile);
        }
        return 1;
    }

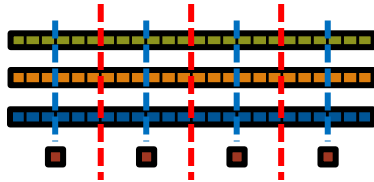
#ifdef _OPENMP
#pragma omp parallel for
#endif
    for (j=0; j<VectorSize; j++) {
        b[j] = 2.0;
        c[j] = 1.0;
    }

    scalar = 3.0;

#ifdef _OPENMP
#pragma omp parallel for
#endif
    for (j=0; j<VectorSize; j++)
        a[j] = b[j]+scalar*c[j];

    HPCC_free(c);
    HPCC_free(b);
    HPCC_free(a);

    return 0;
}
```



CUDA

```
#define N 2000000

int main() {
    float *d_a, *d_b, *d_c;
    float scalar;

    cudaMalloc((void**) &d_a, sizeof(float)*N);
    cudaMalloc((void**) &d_b, sizeof(float)*N);
    cudaMalloc((void**) &d_c, sizeof(float)*N);

    dim3 dimBlock(128);
    if (N % dimBlock.x != 0) dimGrid
        = (N / dimBlock.x) + 1;

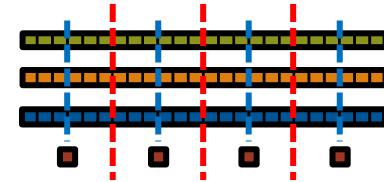
    set_array<<<dimGrid,dimBlock>>>(d_b, .5f, N);
    set_array<<<dimGrid,dimBlock>>>(d_c, .5f, N);

    scalar=3.0f;
    STREAM_Triad<<<dimGrid,dimBlock>>>(d_b, d_c, d_a, scalar, N);
    cudaThreadSynchronize();

    cudaFree(d_a);
    cudaFree(d_b);
    cudaFree(d_c);

    __global__ void set_array(float *a, float value, int len) {
        int idx = threadIdx.x + blockIdx.x * blockDim.x;
        if (idx < len) a[idx] = value;
    }

    __global__ void STREAM_Triad(float *a, float *b, float *c,
                                float scalar, int len) {
        int idx = threadIdx.x + blockIdx.x * blockDim.x;
        if (idx < len) c[idx] = a[idx]+scalar*b[idx];
    }
}
```



HPCC suffers from too many distinct notations for expressing parallelism and locality



Why so many programming models?

HPC tends to approach programming models bottom-up:
Given a system and its core capabilities...
...provide features that can access the available performance.

- portability, generality, programmability: not strictly necessary.

Type of HW Parallelism	Programming Model	Unit of Parallelism
Inter-node	MPI	executable
Intra-node/multicore	OpenMP / pthreads	iteration/task
Instruction-level vectors/threads	pragmas	iteration
GPU/accelerator	CUDA / Open[MP CL ACC]	SIMD function/task

benefits: lots of control; decent generality; easy to implement
downsides: lots of user-managed detail; brittle to changes

Rewinding a few slides...

MPI + OpenMP

```
#ifdef _OPENMP
#include <omp.h>
#endif

static int VectorSize;
static double *a, *b, *c;

int HPCC_StarStream(HPCC_Params *params) {
    int myRank, commSize;
    int rv, errCount;
    MPI_Comm comm = MPI_COMM_WORLD;

    MPI_Comm_size(comm, &commSize);
    MPI_Comm_rank(comm, &myRank);

    rv = HPCC_Stream(params, 0 == myRank);
    MPI_Reduce(&rv, &errCount, 1, MPI_INT, MPI_SUM, 0, comm);

    return errCount;
}

int HPCC_Stream(HPCC_Params *params, int myRank) {
    double scalar;

    VectorSize = HPCC_LocalVectorSize(params, 3, sizeof(double), 0);

    a = HPCC_XMALLOC(double, VectorSize);
    b = HPCC_XMALLOC(double, VectorSize);
    c = HPCC_XMALLOC(double, VectorSize);

    if (!a || !b || !c) {
        if (c) HPCC_free(c);
        if (b) HPCC_free(b);
        if (a) HPCC_free(a);
        if (doIO) {
            fprintf(outFile, "Failed to allocate memory (%d).\n", VectorSize);
            fclose(outFile);
        }
        return 1;
    }

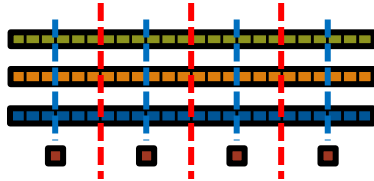
#ifdef _OPENMP
#pragma omp parallel for
#endif
    for (j=0; j<VectorSize; j++) {
        b[j] = 2.0;
        c[j] = 1.0;
    }

    scalar = 3.0;

#ifdef _OPENMP
#pragma omp parallel for
#endif
    for (j=0; j<VectorSize; j++)
        a[j] = b[j]+scalar*c[j];

    HPCC_free(c);
    HPCC_free(b);
    HPCC_free(a);

    return 0;
}
```



CUDA

```
#define N 2000000

int main() {
    float *d_a, *d_b, *d_c;
    float scalar;

    cudaMalloc((void**) &d_a, sizeof(float)*N);
    cudaMalloc((void**) &d_b, sizeof(float)*N);
    cudaMalloc((void**) &d_c, sizeof(float)*N);

    dim3 dimBlock(128);
    if (N % dimBlock.x != 0) dimGrid
        = (N / dimBlock.x) + 1;

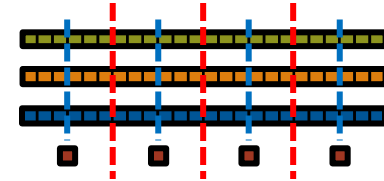
    set_array<<<dimGrid,dimBlock>>>(d_b, .5f, N);
    set_array<<<dimGrid,dimBlock>>>(d_c, .5f, N);

    scalar=3.0f;
    STREAM_Triad<<<dimGrid,dimBlock>>>(d_b, d_c, d_a, scalar, N);
    cudaThreadSynchronize();

    cudaFree(d_a);
    cudaFree(d_b);
    cudaFree(d_c);

    __global__ void set_array(float *a, float value, int len) {
        int idx = threadIdx.x + blockIdx.x * blockDim.x;
        if (idx < len) a[idx] = value;
    }

    __global__ void STREAM_Triad(float *a, float *b, float *c,
                                float scalar, int len) {
        int idx = threadIdx.x + blockIdx.x * blockDim.x;
        if (idx < len) c[idx] = a[idx]+scalar*b[idx];
    }
}
```



HPC suffers from too many distinct notations for expressing parallelism and locality

STREAM Triad: Chapel

MPI + OpenMP

```
#include <hpcc.h>
#ifdef _OPENMP
#include <omp.h>
#endif

static int VectorSize;
static double *a, *b, *c;

int HPCC_StarStream(HPCC_Params *params)
{
    int myRank, commSize;
    int rv, errCount;
    MPI_Comm comm = MPI_COMM_WORLD;

    MPI_Comm_size( comm, &commSize );
    MPI_Comm_rank( comm, &myRank );

    rv = HPCC_Stream( params, 0 == myRank );
    MPI_Reduce( &rv, &errCount, 1, MPI_INT, 0, comm );

    return errCount;
}

int HPCC_Stream(HPCC_Params *params,
    register int j;
    double scalar;

    VectorSize = HPCC_LocalVectorSize( params );
    a = HPCC_XMALLOC( double, VectorSize );
    b = HPCC_XMALLOC( double, VectorSize );
    c = HPCC_XMALLOC( double, VectorSize );

    if (!a || !b || !c) {
        if (c) HPCC_free(c);
        if (b) HPCC_free(b);
        if (a) HPCC_free(a);
    }
    if (doIO) {
        fprintf( outFile, "Failed to allocate memory (%d).\n", VectorSize );
        fclose( outFile );
    }
}
```

Chapel

```
config const m = 1000,
              alpha = 3.0;

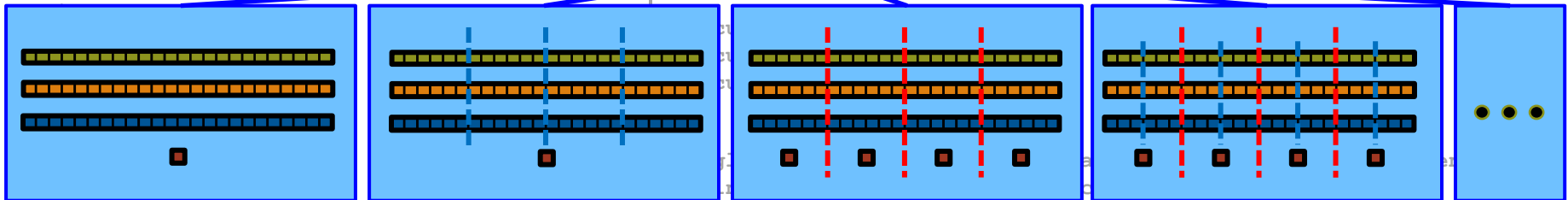
const ProblemSpace = {1..m} dmapped ...;

var A, B, C: [ProblemSpace] real;

B = 2.0;
C = 1.0;

A = B + alpha * C;
```

the special sauce



Philosophy: Good, *top-down* language design can tease system-specific implementation details away from an algorithm, permitting the compiler, runtime, applied scientist, and HPC expert to each focus on their strengths.



What is Chapel?

Chapel: A productive parallel programming language

- portable
- open-source
- a collaborative effort

Goals:

- Support general parallel programming
 - “any parallel algorithm on any parallel hardware”
- Make parallel programming at scale far more productive





What does “Productivity” mean to you?

Recent Graduates:

“something similar to what I used in school: Python, Matlab, Java, ...”

Seasoned HPC Programmers:

“that sugary stuff that I don’t need because I ~~was born to suffer~~
want full control
to ensure performance”

Computational Scientists:

“something that lets me express my parallel computations
without having to wrestle with architecture-specific details”

Chapel Team:

“something that lets computational scientists express what they want,
without taking away the control that HPC programmers want,
implemented in a language as attractive as recent graduates want.”





Chapel is Portable

- **Chapel is designed to be hardware-independent**
- **The current release requires:**
 - a C/C++ compiler
 - a *NIX environment (Linux, OS X, BSD, Cygwin, ...)
 - POSIX threads
 - UDP, MPI, or RDMA (if distributed memory execution is desired)
- **Chapel can run on...**
 - ...laptops and workstations
 - ...commodity clusters
 - ...the cloud
 - ...HPC systems from Cray and other vendors
 - ...modern processors like Intel Xeon Phi, GPUs*, etc.

* = academic work only; not yet supported in the official release





Chapel is Open-Source

- **Chapel's development is hosted at GitHub**
 - <https://github.com/chapel-lang>
- **Chapel is licensed as Apache v2.0 software**
- **Instructions for download + install are online**
 - see <http://chapel.cray.com/download.html> to get started



The Chapel Team at Cray (May 2016)



Chapel Community R&D Efforts



THE GEORGE
WASHINGTON
UNIVERSITY
WASHINGTON, DC



Lawrence Berkeley
National Laboratory



Sandia National Laboratories



Yale

(and several others...)

<http://chapel.cray.com/collaborations.html>



COMPUTE | STORE | ANALYZE



Outline

- ✓ Chapel Motivation and Background
- Chapel in a Nutshell
- Chapel Project: Past, Present, Future



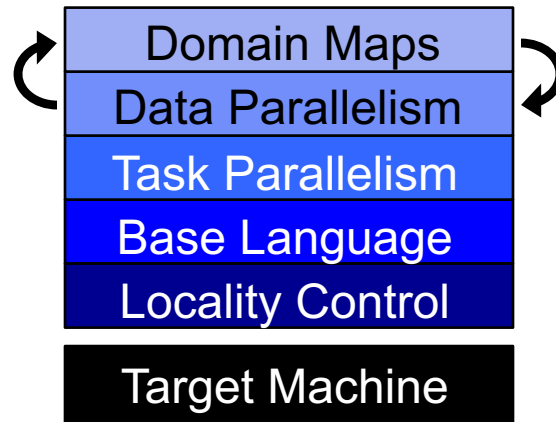


Chapel's Multiresolution Philosophy

Multiresolution Design: Support multiple tiers of features

- higher levels for programmability, productivity
- lower levels for greater degrees of control

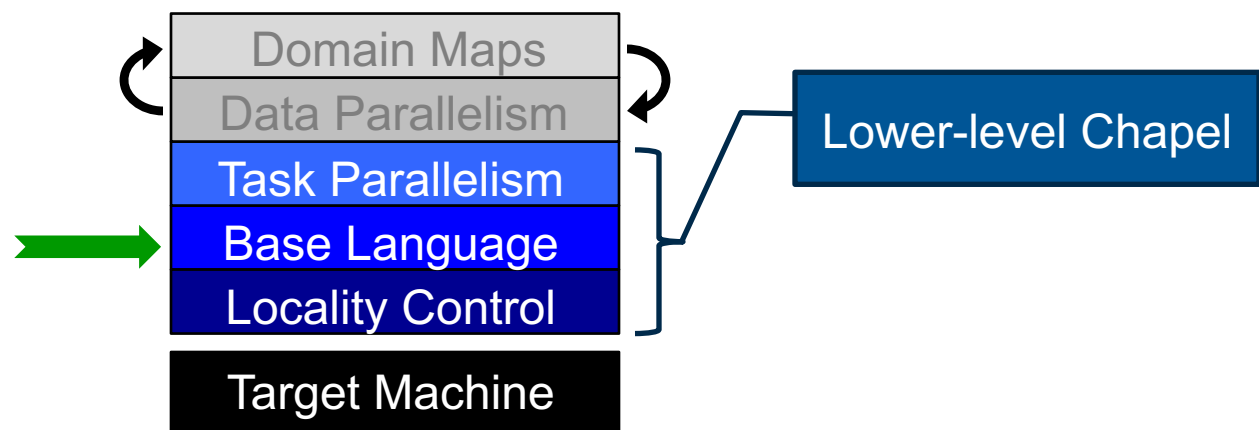
Chapel language concepts



- build the higher-level concepts in terms of the lower
- permit the user to intermix layers arbitrarily



Base Language





Base Language Features, by example

```
iter fib(n) {  
    var current = 0,  
        next = 1;  
  
    for i in 1..n {  
        yield current;  
        current += next;  
        current <=> next;  
    }  
}
```

```
config const n = 10;  
  
for f in fib(n) do  
    writeln(f);
```

```
0  
1  
1  
2  
3  
5  
8  
...
```



Base Language Features, by example

Modern iterators

```

iter fib(n) {
    var current = 0,
        next = 1;

    for i in 1..n {
        yield current;
        current += next;
        current <=> next;
    }
}

```

```

config const n = 10;

for f in fib(n) do
    writeln(f);

```

```

0
1
1
2
3
5
8
...

```

Base Language Features, by example

Configuration declarations
(to avoid command-line argument parsing)
`./a.out --n=1000000`

```
iter fib(n) {
    var current = 0,
        next = 1;

    for i in 1..n {
        yield current;
        current += next;
        current <=> next;
    }
}
```

```
config const n = 10;

for f in fib(n) do
    writeln(f);
```

```
0
1
1
2
3
5
8
...
```

Base Language Features, by example

Static type inference for:

- arguments
- return types
- variables

```
iter fib(n) {
  var current = 0,
    next = 1;

  for i in 1..n {
    yield current;
    current += next;
    current <=> next;
  }
}
```

```
config const n = 10;

for f in fib(n) do
  writeln(f);
```

```
0
1
1
2
3
5
8
...
```

Base Language Features, by example

Zippered iteration

```
iter fib(n) {
    var current = 0,
        next = 1;

    for i in 1..n {
        yield current;
        current += next;
        current <=> next;
    }
}
```

```
config const n = 10;

for (i, f) in zip(0..#n, fib(n)) do
    writeln("fib #", i, " is ", f);
```

```
fib #0 is 0
fib #1 is 1
fib #2 is 1
fib #3 is 2
fib #4 is 3
fib #5 is 5
fib #6 is 8
...
```

Base Language Features, by example

Range types and operators

```
iter fib(n) {
  var current = 0;
  next = 1;

  for i in 1..n {
    yield current;
    current += next;
    current <=> next;
  }
}
```

```
config const n = 10;

for (i, f) in zip(0..#n, fib(n)) do
  writeln("fib #", i, " is ", f);
```

```
fib #0 is 0
fib #1 is 1
fib #2 is 1
fib #3 is 2
fib #4 is 3
fib #5 is 5
fib #6 is 8
...
```

Base Language Features, by example

tuples

```
iter fib(n) {
    var current = 0,
        next = 1;

    for i in 1..n {
        yield current;
        current += next;
        current <=> next;
    }
}
```

```
config const n = 10;

for (i, f) in zip(0..#n, fib(n)) do
    writeln("fib #", i, " is ", f);
```

```
fib #0 is 0
fib #1 is 1
fib #2 is 1
fib #3 is 2
fib #4 is 3
fib #5 is 5
fib #6 is 8
...
```

Base Language Features, by example

```
iter fib(n) {
    var current = 0,
        next = 1;

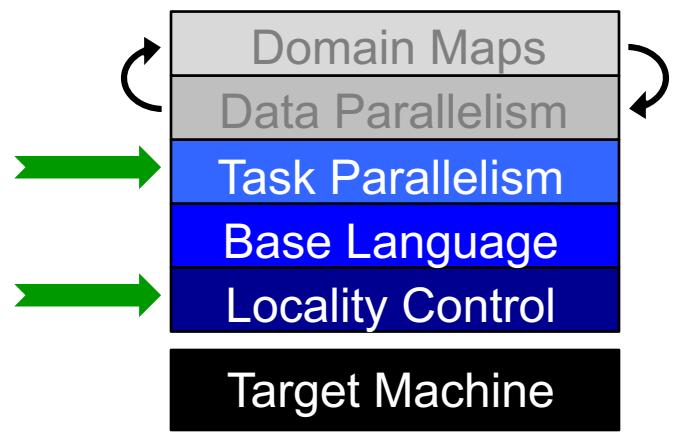
    for i in 1..n {
        yield current;
        current += next;
        current <=> next;
    }
}
```

```
config const n = 10;

for (i, f) in zip(0..#n, fib(n)) do
    writeln("fib #", i, " is ", f);
```

```
fib #0 is 0
fib #1 is 1
fib #2 is 1
fib #3 is 2
fib #4 is 3
fib #5 is 5
fib #6 is 8
...
```

Task Parallelism



Task Parallelism, Locality Control, by example

taskParallel.chpl

```
coforall loc in Locales do
  on loc {
    const numTasks = here.maxTaskPar;
    coforall tid in 1..numTasks do
      writef("Hello from task %n of %n "+
            "running on %s\n",
            tid, numTasks, here.name);
    }
  }
```

```
prompt> chpl taskParallel.chpl -o taskParallel
prompt> ./taskParallel --numLocales=2
Hello from task 1 of 2 running on n1033
Hello from task 2 of 2 running on n1032
Hello from task 2 of 2 running on n1033
Hello from task 1 of 2 running on n1032
```

Task Parallelism, Locality Control, by example

Abstraction of
System Resources

taskParallel.chpl

```
coforall loc in Locales do
  on loc {
    const numTasks = here.maxTaskPar;
    coforall tid in 1..numTasks do
      writef("Hello from task %n of %n "+
            "running on %s\n",
            tid, numTasks, here.name);
    }
  }
```

```
prompt> chpl taskParallel.chpl -o taskParallel
prompt> ./taskParallel --numLocales=2
Hello from task 1 of 2 running on n1033
Hello from task 2 of 2 running on n1032
Hello from task 2 of 2 running on n1033
Hello from task 1 of 2 running on n1032
```

Task Parallelism, Locality Control, by example

High-Level
Task Parallelism

taskParallel.chpl

```
coforall loc in Locales do
  on loc {
    const numTasks = here.maxTaskPar;
    coforall tid in 1..numTasks do
      writef("Hello from task %n of %n "+
            "running on %s\n",
            tid, numTasks, here.name);
    }
  }
```

```
prompt> chpl taskParallel.chpl -o taskParallel
prompt> ./taskParallel --numLocales=2
Hello from task 1 of 2 running on n1033
Hello from task 2 of 2 running on n1032
Hello from task 2 of 2 running on n1033
Hello from task 1 of 2 running on n1032
```

Task Parallelism, Locality Control, by example

taskParallel.chpl

```
coforall loc in Locales do
  on loc {
    const numTasks = here.maxTaskPar;
    coforall tid in 1..numTasks do
      writef("Hello from task %n of %n "+
            "running on %s\n",
            tid, numTasks, here.name);
    }
  }
```

Control of Locality/Affinity

```
prompt> chpl taskParallel.chpl -o taskParallel
prompt> ./taskParallel --numLocales=2
Hello from task 1 of 2 running on n1033
Hello from task 2 of 2 running on n1032
Hello from task 2 of 2 running on n1033
Hello from task 1 of 2 running on n1032
```

Task Parallelism, Locality Control, by example

Abstraction of
System Resources

taskParallel.chpl

```
coforall loc in Locales do
  on loc {
    const numTasks = here.maxTaskPar;
    coforall tid in 1..numTasks do
      writef("Hello from task %n of %n "+
            "running on %s\n",
            tid, numTasks, here.name);
    }
  }
```

```
prompt> chpl taskParallel.chpl -o taskParallel
prompt> ./taskParallel --numLocales=2
Hello from task 1 of 2 running on n1033
Hello from task 2 of 2 running on n1032
Hello from task 2 of 2 running on n1033
Hello from task 1 of 2 running on n1032
```

Task Parallelism, Locality Control, by example

High-Level
Task Parallelism

taskParallel.chpl

```
coforall loc in Locales do
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    coforall tid in 1..numTasks do
      writef("Hello from task %n of %n "+
            "running on %s\n",
            tid, numTasks, here.name);
    }
  }
```

```
prompt> chpl taskParallel.chpl -o taskParallel
prompt> ./taskParallel --numLocales=2
Hello from task 1 of 2 running on n1033
Hello from task 2 of 2 running on n1032
Hello from task 2 of 2 running on n1033
Hello from task 1 of 2 running on n1032
```

Task Parallelism, Locality Control, by example

taskParallel.chpl

```
coforall loc in Locales do
  on loc {
    const numTasks = here.maxTaskPar;
    coforall tid in 1..numTasks do
      writef("Hello from task %n of %n "+
            "running on %s\n",
            tid, numTasks, here.name);
    }
  }
```

Not seen here:

Data-centric task coordination
via atomic and full/empty vars

```
prompt> chpl taskParallel.chpl -o taskParallel
prompt> ./taskParallel --numLocales=2
Hello from task 1 of 2 running on n1033
Hello from task 2 of 2 running on n1032
Hello from task 2 of 2 running on n1033
Hello from task 1 of 2 running on n1032
```

Task Parallelism, Locality Control, by example

taskParallel.chpl

```
coforall loc in Locales do
  on loc {
    const numTasks = here.maxTaskPar;
    coforall tid in 1..numTasks do
      writef("Hello from task %n of %n "+
            "running on %s\n",
            tid, numTasks, here.name);
    }
  }
```

```
prompt> chpl taskParallel.chpl -o taskParallel
prompt> ./taskParallel --numLocales=2
Hello from task 1 of 2 running on n1033
Hello from task 2 of 2 running on n1032
Hello from task 2 of 2 running on n1033
Hello from task 1 of 2 running on n1032
```




Parallelism and Locality: Orthogonal in Chapel

- This is a **parallel**, but local program:

```
coforall i in 1..msgs do
  writeln("Hello from task ", i);
```

- This is a **distributed**, but serial program:

```
writeln("Hello from locale 0!");
on Locales[1] do writeln("Hello from locale 1!");
on Locales[2] do writeln("Hello from locale 2!");
```

- This is a **distributed parallel** program:

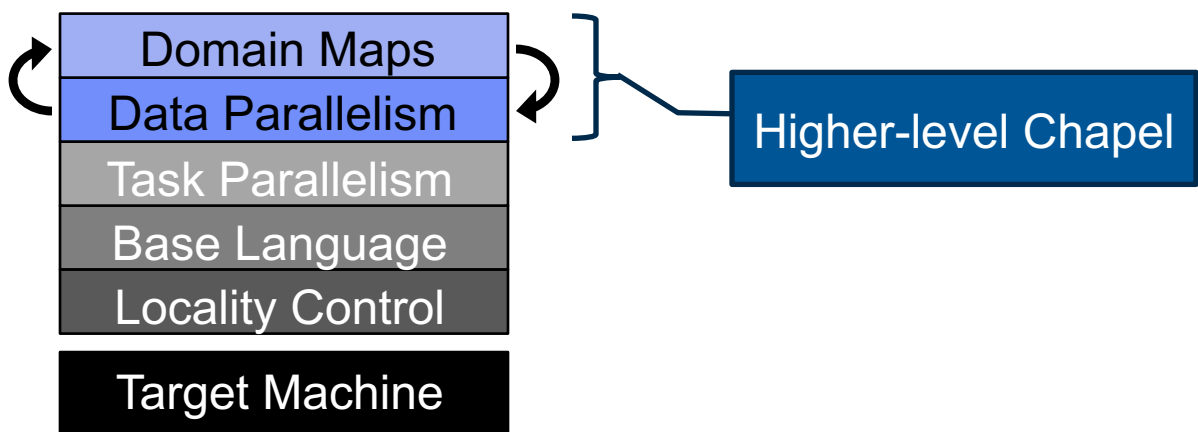
```
coforall i in 1..msgs do
  on Locales[i%numLocales] do
    writeln("Hello from task ", i,
           " running on locale ", here.id);
```





Higher-Level Features

Chapel language concepts



Data Parallelism, by example

dataParallel.chpl

```
config const n = 1000;
var D = {1..n, 1..n};

var A: [D] real;
forall (i,j) in D do
    A[i,j] = i + (j - 0.5)/n;
writeln(A);
```

```
prompt> chpl dataParallel.chpl -o dataParallel
prompt> ./dataParallel --n=5
1.1 1.3 1.5 1.7 1.9
2.1 2.3 2.5 2.7 2.9
3.1 3.3 3.5 3.7 3.9
4.1 4.3 4.5 4.7 4.9
5.1 5.3 5.5 5.7 5.9
```

Data Parallelism, by example

Domains (Index Sets)

dataParallel.chpl

```
config const n = 1000;
var D = {1..n, 1..n};

var A: [D] real;
forall (i,j) in D do
    A[i,j] = i + (j - 0.5)/n;
writeln(A);
```

```
prompt> chpl dataParallel.chpl -o dataParallel
prompt> ./dataParallel --n=5
1.1 1.3 1.5 1.7 1.9
2.1 2.3 2.5 2.7 2.9
3.1 3.3 3.5 3.7 3.9
4.1 4.3 4.5 4.7 4.9
5.1 5.3 5.5 5.7 5.9
```

Data Parallelism, by example

Arrays

dataParallel.chpl

```
config const n = 1000;
var D = {1..n, 1..n};

var A: [D] real;
forall (i,j) in D do
    A[i,j] = i + (j - 0.5)/n;
writeln(A);
```

```
prompt> chpl dataParallel.chpl -o dataParallel
prompt> ./dataParallel --n=5
1.1 1.3 1.5 1.7 1.9
2.1 2.3 2.5 2.7 2.9
3.1 3.3 3.5 3.7 3.9
4.1 4.3 4.5 4.7 4.9
5.1 5.3 5.5 5.7 5.9
```

Data Parallelism, by example

Data-Parallel Forall Loops

dataParallel.chpl

```
config const n = 1000;
var D = {1..n, 1..n};

var A: [D] real;
forall (i,j) in D do
    A[i,j] = i + (j - 0.5)/n;
writeln(A);
```

```
prompt> chpl dataParallel.chpl -o dataParallel
prompt> ./dataParallel --n=5
1.1 1.3 1.5 1.7 1.9
2.1 2.3 2.5 2.7 2.9
3.1 3.3 3.5 3.7 3.9
4.1 4.3 4.5 4.7 4.9
5.1 5.3 5.5 5.7 5.9
```

Distributed Data Parallelism, by example

Domain Maps
(Map Data Parallelism to the System)

dataParallel.chpl

```
use CyclicDist;
config const n = 1000;
var D = {1..n, 1..n}
        dmapped Cyclic(startIdx = (1,1));
var A: [D] real;
forall (i,j) in D do
    A[i,j] = i + (j - 0.5)/n;
writeln(A);
```

```
prompt> chpl dataParallel.chpl -o dataParallel
prompt> ./dataParallel --n=5 --numLocales=4
1.1 1.3 1.5 1.7 1.9
2.1 2.3 2.5 2.7 2.9
3.1 3.3 3.5 3.7 3.9
4.1 4.3 4.5 4.7 4.9
5.1 5.3 5.5 5.7 5.9
```

Distributed Data Parallelism, by example

dataParallel.chpl

```
use CyclicDist;
config const n = 1000;
var D = {1..n, 1..n}
        dmapped Cyclic(startIdx = (1,1));
var A: [D] real;
forall (i,j) in D do
    A[i,j] = i + (j - 0.5)/n;
writeln(A);
```

```
prompt> chpl dataParallel.chpl -o dataParallel
prompt> ./dataParallel --n=5 --numLocales=4
1.1 1.3 1.5 1.7 1.9
2.1 2.3 2.5 2.7 2.9
3.1 3.3 3.5 3.7 3.9
4.1 4.3 4.5 4.7 4.9
5.1 5.3 5.5 5.7 5.9
```




Outline

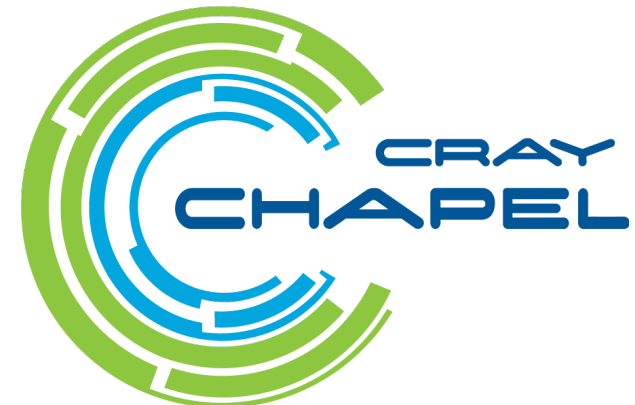
- ✓ Chapel Motivation and Background
- ✓ Chapel in a Nutshell
- **Chapel Project: Past, Present, Future**



Chapel's Origins: HPCS

DARPA HPCS: High Productivity Computing Systems

- **Goal:** improve productivity by a factor of 10x
- **Timeframe:** Summer 2002 – Fall 2012
- Cray developed a new system architecture, network, software stack...
 - this became the very successful Cray XC30™ Supercomputer Series



...and a new programming language: Chapel



Chapel's 5-year push

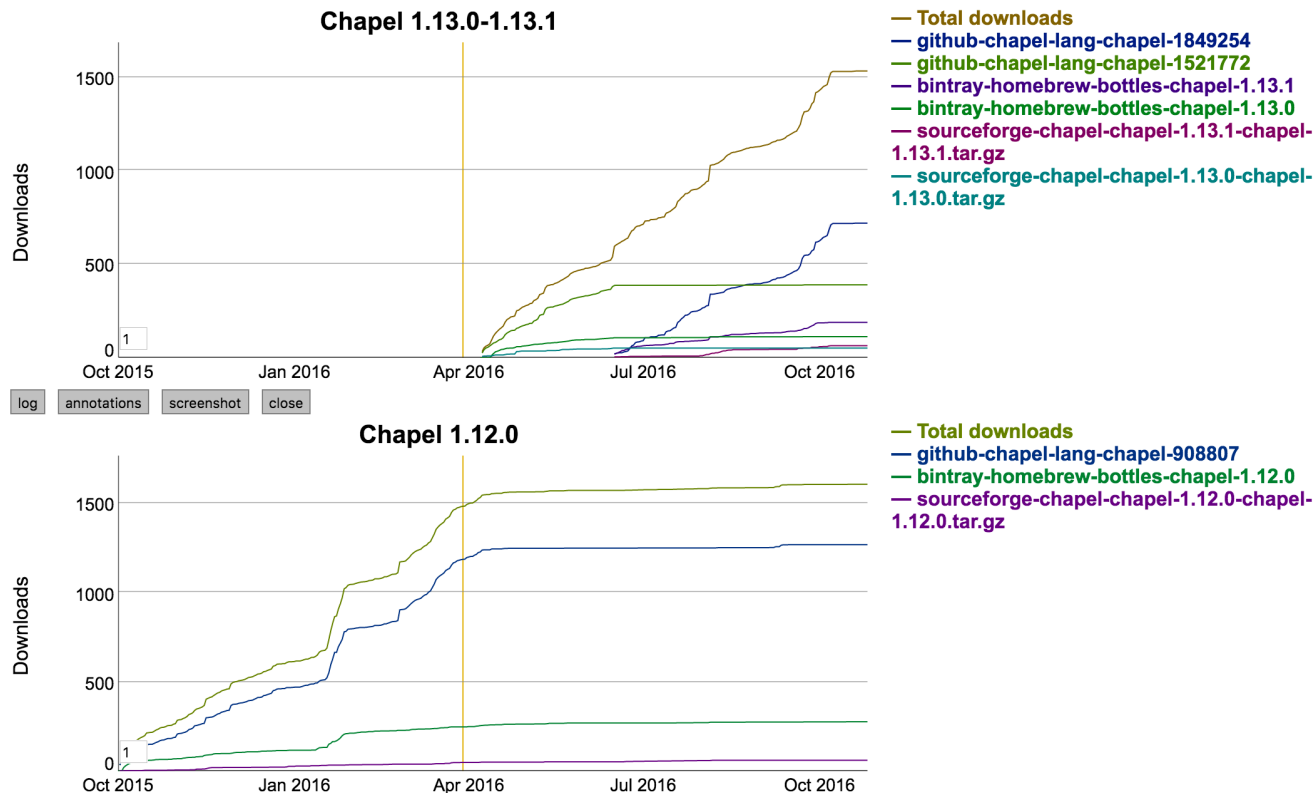
- Based on positive user response to Chapel under HPCS, Cray undertook a five-year effort to improve it
 - we've recently completed our third year
- Focus Areas:
 1. Improving **performance** and scaling
 2. **Fixing** immature aspects of the language and implementation
 - e.g., strings, memory management, error handling, ...
 3. **Porting** to emerging architectures
 - Intel Xeon Phi, accelerators, heterogeneous processors and memories, ...
 4. Improving **interoperability**
 5. Growing the Chapel user and developer **community**
 - including non-scientific computing communities
 6. Exploring transition of Chapel **governance** to a neutral, external body





Chapel is a Work-in-Progress

- Currently being picked up by early adopters
 - 3000+ downloads per year across two releases



- Users who try it generally like what they see




A notable early adopter


Chapel in the (Cosmological) Wild


1:00 – 2:00

Nikhil Padmanabhan, *Yale University Professor, Physics & Astronomy*

Abstract: This talk aims to present my personal experiences using Chapel in my research. My research interests are in observational cosmology; more specifically, I use large surveys of galaxies to constrain the evolution of the




[Chapel Parallel Programming Language](#)
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CHIOW 2016 keynote: "Chapel in the (Cosmological) Wild", Nikhil Padmanabhan

Chapel Parallel Programming Language
1 month ago • 86 views

This is Nikhil Padmanabhan's keynote talk from CHIOW 2016: the 3rd Annual Chapel Implementers and Users workshop. The slides are availabl...

56:14



Chapel is a Work-in-Progress

- **Currently being picked up by early adopters**
 - Last two releases got ~3500 downloads total in a year
 - Users who try it generally like what they see
- **Most core features are functional and working well**
 - some areas need improvements, e.g., error-handling, constructors
- **Performance varies, but is continually improving**
 - shared memory performance is typically competitive with C+OpenMP
 - distributed memory performance tends to be more hit-and-miss
 - PAW workshop talk tomorrow: LLNL got 87% of reference version for CoMD
- **We are actively working on addressing these lacks**





Chapel-related Events at SC16

Today: This tutorial

Today: Women in HPC Workshop (all day)

- *Array initialization improvements in Chapel:* Lydia Duncan (Cray)

This evening: CHUG (Chapel Users Group) happy hour

- 7th annual meet-up, everyone's welcome to attend
- 5:30pm Settebello Pizzeria Napoletana

Monday afternoon: PGAS Applications Workshop

- *CoMD study in Chapel:* Dave Richards and Riyaz Haque (LLNL)
- *ISx study in SHMEM and Chapel:* Jake Hemstad (U Minn / Sandia), Ulf Hanebutte (Intel), Ben Harshbarger and Brad Chamberlain (Cray)
- *PGAS Applications panel:* chaired by Brad Chamberlain (Cray)

Wednesday: PGAS BoF, 12:15pm

Thursday: Talk to a Chapel developer, PGAS booth, 10am-noon

all week: PGAS Booth Poster on Chapel CoMD study, Meet by Request

additional details at <http://chapel.cray.com/events.html>



High-level Questions about Chapel?





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