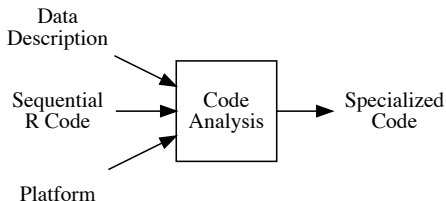


# Parallel Computing Through Code Analysis

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Modern platforms provide incredible computing power.

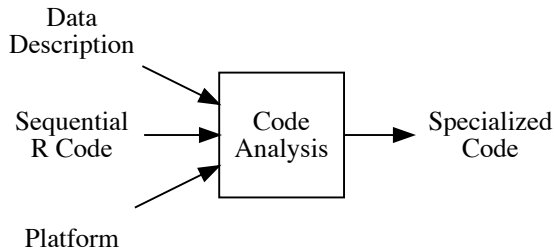


But they require expertise.

The broader goal is for users to write higher level code that also performs better.

- Parallel programming is a means to this end
- Compilation is another way
- Expertise in system rather than end user

We take a holistic approach to the computation.



# The R language offers several benefits.



- Functional languages simplify parallel computing
- Widely used for statistics and data analysis
- Supports metaprogramming aka “programming on the language”

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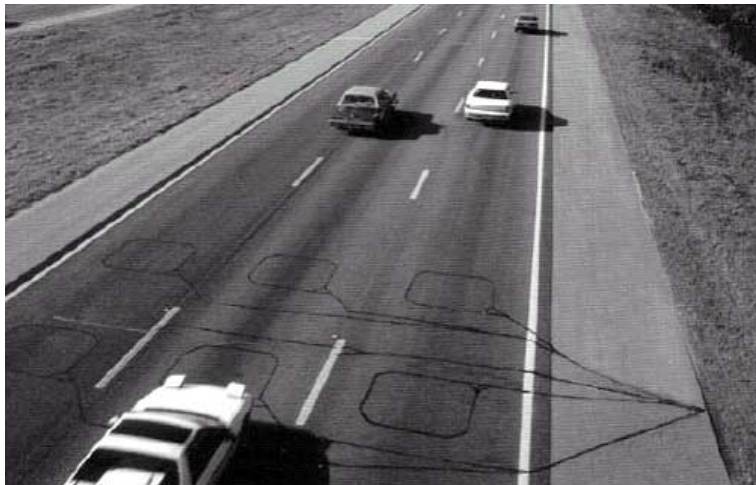
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The purpose of this example is to motivate the proposed research.

- Working with Professor Michael Zhang from Civil Engineering
- Illustrates complexity when computing with larger data sets

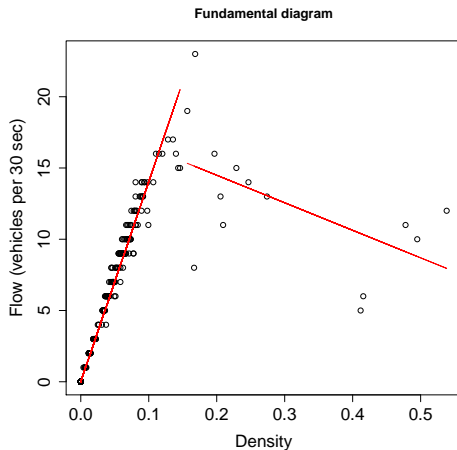


Loop detectors count vehicle flow, measuring velocity and density (time sensor is activated).

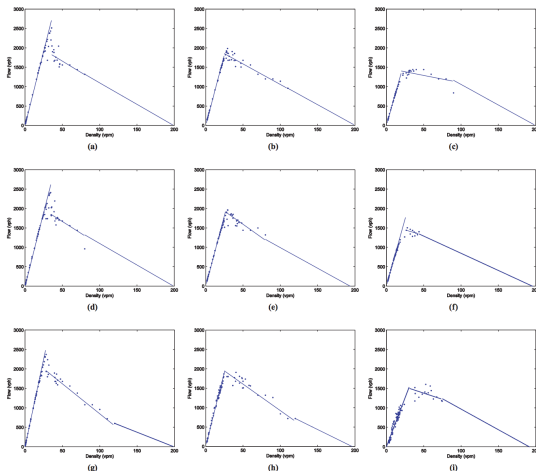


Around 400 million data points per day recorded in California.

The *fundamental diagram* in traffic engineering shows the relationship between flow and density.



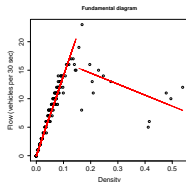
Each station has a fundamental diagram, which can be fit in parallel.



Source: Li, Zhang 2011

Using more data allows new types of analysis.

## R expresses statistical computation well.



```
by(data=single_day, INDICES = station, FUN = my_fd)
```

- For a single day with 400 million observations this can be done on a single machine.
- A sensible way to run this in parallel is to `fork()` the process after reading in the data.
- So you write a bunch of code to do that :)

## Small changes can require totally different computations.

If we compute on one year then this will exceed memory.

```
by(data=one_year, INDICES = station, FUN = my_fd)
```

A different model, such as least squares, may be able to process the data as a stream.

```
by(data=one_year, INDICES = station, FUN = my_lm_fd)
```

Access to the underlying database may allow us to run code directly inside the database.



```
SELECT station, my_fd(...) FROM data GROUP BY station
```

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# This simple example shows how to write parallel code in R.

Consider computing the mean,

$$\bar{x} = \frac{1}{n} \sum_{i=1}^n x_i \quad (1)$$

where the  $x_i$ 's are i.i.d.  $\sim t(d)$ .

In R this code is written:

```
xbar = mean(rt(n, d))
```



We can express the mean as a weighted mean.

Suppose  $n = n_j p$ , where  $n_j$  is the chunk size and  $p$  is the number of chunks.

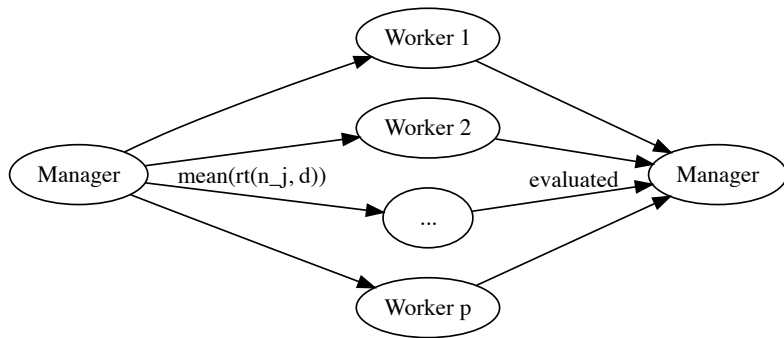
$$\bar{x} = \frac{1}{n} \sum_{j=1}^p \sum_{i=1}^{n_j} x_{ij} = \frac{1}{p} \sum_{j=1}^p \frac{1}{n_j} \sum_{i=1}^{n_j} x_{ij} = \frac{1}{p} \sum_{j=1}^p \bar{x}_{\cdot j} \quad (2)$$

The weighted mean can be directly translated into R code.

```
partial_means = replicate(p, mean(rt(n_j, d)))  
xbar = mean(partial_means)
```

- While not parallel, this effectively removes the memory limits.
- How to choose  $n_j$  and  $p$ ?

The same computation can be evaluated on many workers simultaneously.



Here is one way to parallelize this code.

```
library(parallel)
p = floor(detectCores(logical = FALSE) / 2)
n_j = n / p
cluster = makeCluster(p)

expr = substitute(mean(rt(n_j, d)),
  list(d = d, n_j = n_j))

partial_means = unlist(
  clusterCall(cluster, eval, expr))

xbar = mean(partial_means)
```

We're considering a system that transforms expressions.

**Input:**

```
xbar = mean(rt(n, d))
```

**Output:** (omitting boilerplate)

```
p = floor(detectCores(logical = FALSE) / 2)
```

```
expr = substitute(mean(rt(n_j, d)),  
  list(d = d, n_j = n_j))
```

```
partial_means = clusterCall(cluster, eval, expr)  
xbar = mean(partial_means)
```

This can be difficult because R is implemented in C.

```
> head(rt, 4)
```

```
1 function (n, df, ncp)
2 {
3     if (missing(ncp))
4         .Call(C_rt, n, df)
```

Options:

- Start from `replicate(p, mean(rt(n_j, d)))`
- Allow users to indicate how `rt` is vectorized
- Analyze the preprocessed C code
- Rewrite the C code in R, then analyze the R code

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Idiomatic R already expresses computation in a natural parallel way through “apply” functions.

```
x = replicate(p, mean(rt(n_j, d)), simplify = FALSE)
```

```
partialmeans = lapply(x, mean)
```

```
by(data, INDICES = station, FUN = my_fd)
```

Also

```
apply, sapply, tapply, by, mapply, Map, vapply, outer
```

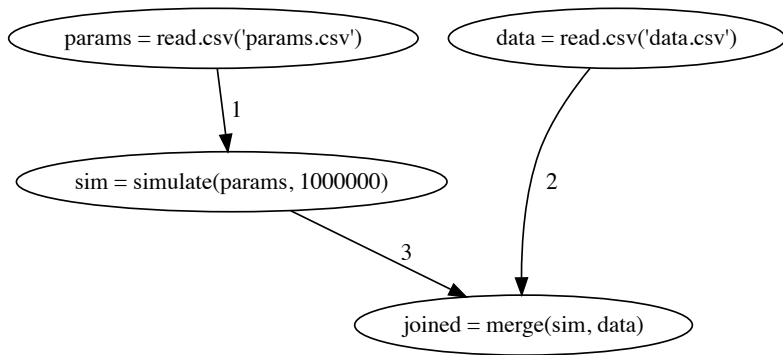


CodeDepends is a tool for analyzing code as a data structure.

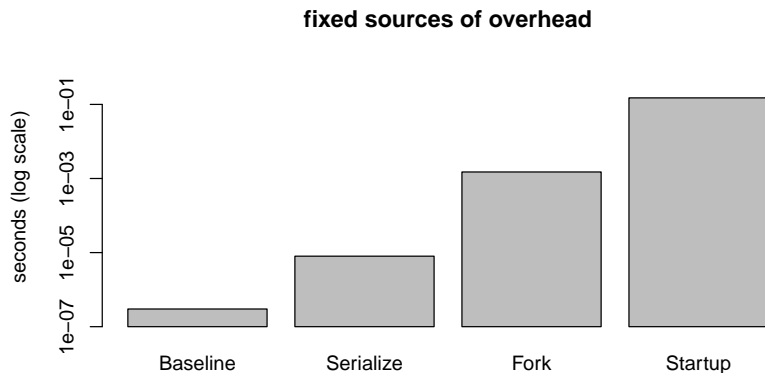
Consider this script:

```
params = read.csv('params.csv')  
data = read.csv('data.csv')  
sim = simulate(params, 1000000)  
joined = merge(data, sim)
```

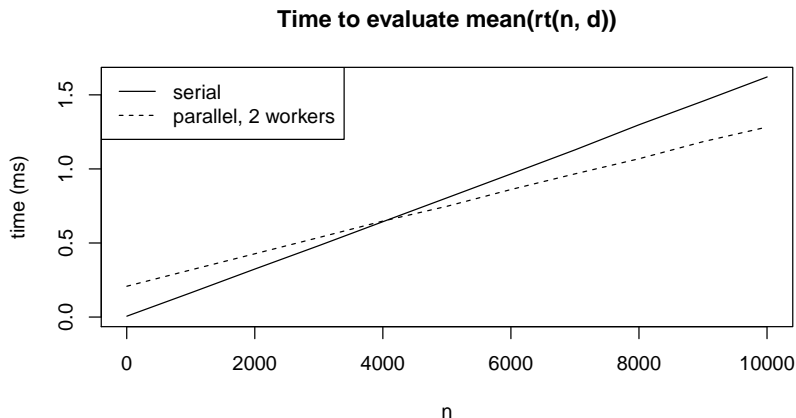
The expression graph represents the dependencies between expressions.



# Is it worth it to go parallel?



Given an existing SNOW cluster with 2 workers we see benefits from parallelization when  $n > 4000$ .

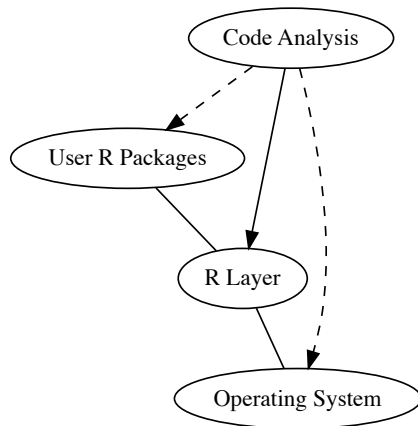


Timings on a 3.4 GHz Intel i3 CPU

Layers mark ways for users to write parallel code for one platform.

- **User R Packages:** foreach, future, partools, ddR, biganalytics, RevoScaleR
- **R Layer:** SNOW, parallel, bigmemory, Rmpi, RCUDA
- **Operating System:** threads, processes, \*NIX fork(), memory maps, network sockets, MPI

# How can we transform R code into a lower layer?



Knowledge of the data allows us to generate more specialized code.

- File size
- Dimensions of table / matrix / array
- Column classes
- Randomized rows
- Sorted / grouped
- Possible values for factor
- Indexed
- Including sufficient statistics

## Example: a data format that facilitates sampling.

station, flow, occupancy, time

1	12	0.087	09:57:00
---	----	-------	----------

1	14	0.092	14:29:30
---	----	-------	----------

...

7	14	0.088	16:32:30
---	----	-------	----------

7	11	0.090	17:12:00
---	----	-------	----------

- ASCII fixed width format file,  $c$  characters (bytes) per row
- sorted on station, then occupancy
- $r$  rows per station
- $\implies$  new stations begin at byte  $i \times c \times r$



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We propose using {Code, Data, Platform} to determine a parallel execution strategy.

```
1 library(BenfordTests)
2
3 # Using data downloaded from
4 # http://www.fec.gov/finance/disclosure/ftpdet.shtml
5 # Individual contributions file for 2017 is 4GB
6
7 r = edist.benftest(x)
8
9 header = as.character(read.table("indiv_header_file.csv", sep = ",",
10                                stringsAsFactors = FALSE))
11
12 keepers = c("CITE_ID", "ENTITY_ID", "TRANSACTION_AMT")
13 keep = header %>% keepers
14
15 cc = rep("NULL", length(header))
16 cc[keep] = c("factor", "factor", "numeric")
17
18 ltcont = read.table("~/data/ltcont.txt",
19                    sep = "\t",
20                    colClasses = cc,
21                    nrows = 200)
22
23
```



## Related Work

- CS literature on automatic parallelization mostly focused on algorithmic applications and lower level languages
- Dask, Theano, Tensorflow: User builds computation / data flow graphs
- Hadoop, Spark, Databases: Powerful, but less flexible than R

# The next step is to build a prototype of the system.

Specifically beginning with:

- *code*: Apply family of functions
- *data*: Files on disk exceeding main memory
- *platform*: Single server

Then test it with the traffic sensor data.

Next I'd like to extend this to platforms that store data

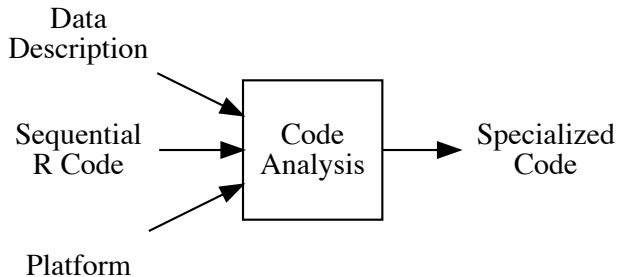
```
SELECT station, my_rlm(...) FROM data GROUP BY station
```

- Better data locality
- Many practical use cases
- Connection to compilation

# Acknowledgements

- Faculty members
- Data Science Institute Affiliates and statistics students for applications and feedback
- Special thanks to Professors Duncan Temple Lang and Michael Zhang

# Questions?



# Additional Slides



## Follow up on the sensor example

The transformed program should:

- ➊ Remove any observations it can
- ➋ Reorganize files on disk based on station ID
- ➌ Apply function to each station ID file

# Caltrans Performance Measurement System (PeMS)

records loop detector data for the whole state.

- Each sensor measures 3 quantities
- Data point every 30 seconds
- 43,680 sensors in California
- $\implies$  377 million data points per day

# Using more data allows new types of analyses.

- Effect of policy such as speed limits and carpool lanes
- Impact of road features such as on/off ramps
- Effect of weather
- Clustering detectors

# Factors to consider when switching to parallel programs

## Parameters

- number of processor cores to use
- size of each chunk
- which functions to combine in one processing step

## Constraints

- number of cores available
- network bandwidth
- disk IO speed
- available memory

# More Applications

- Benford test on election campaign contribution data
- Forest greenness satellite imagery (Andrew Latimer)
- Simulating spread of disease (Nistara Randhawa)

# Steps in Code Analysis

- Parse code, gathering dependency information as explained in section ??.
- Use data description to determine if preprocessing of the data is required
- Identify expressions to parallelize
- May experimentally evaluate expressions
- Relate potentially parallel expressions to data description and timings
- Generate code that will run efficiently on the specific platform.

# Preserving language semantics can be challenging.

For example, R's dynamic lookups

```
f = function() 0
g = function() f() + 1
f = function() 10
g()                      # Returns 11!
```

The CodeDepends package provides tools for detecting this.

## Compiled R code provides even more efficiency.

- ① Parallelization will complement efforts to compile R
- ② Compiled code potentially allows the use of shared memory threads
- ③ May follow the OpenCL kernel model



## Last summer I worked on the Distributed Data Structures in R (DDR) project

- Relevant experience
- Idea: an abstraction layer for distributed and parallel data structures
- Created R lists and apply type functions to run on Spark

## How do we detect if a function in R is vectorized, and in which arguments?

`rnorm()` is vectorized in the last two arguments, but semantically different for a vector in the first argument.

```
> rnorm(5, mean = c(1, 2), sd = c(2, 10, 200))  
[1] 0.2134756 -4.1137221 256.0094734 0.4562226 -10.385537
```

It's all C Code.

```
> rnorm  
function (n, mean = 0, sd = 1)  
.Call(C_rnorm, n, mean, sd)
```

## R in a database

Suppose we want to call the vectorized function  $f()$ . If  $f()$  is available both in R and the database then we have two options:

Option 1: Run  $f$  inside database, returning result

```
dbGetQuery(con,"SELECT f(x) FROM mydata;")
```

Option 2: First fetch  $x$ , then call  $f$  within R

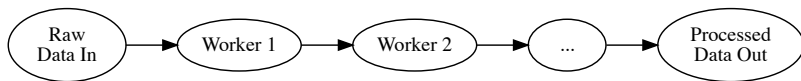
```
f(dbGetQuery(con,"SELECT x FROM mydata;"))
```

By “programming on the language” we can modify existing code.

```
lapply_to_mclapply = function(expr)
{
  # Changes lapply to parallel::mclapply
  lapply = quote(parallel::mclapply)
  expr = force(expr)
  # Following Wickham's Advanced R book
  call = substitute(substitute(expr))
  eval(call)
}

> e1 = quote(xmeans <- lapply(x, mean))
> lapply_to_mclapply(e1)
xmeans <- parallel::mclapply(x, mean)
```

## Pipeline parallelism is like a factory assembly line.



```
# Worker 1
```

```
x_chunk = rnorm(n_j)
```

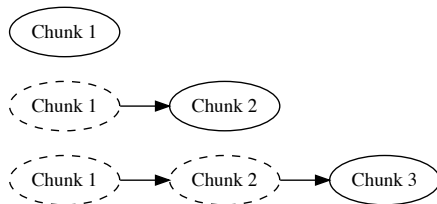
```
serialize(x_chunk, worker2)
```

```
# Worker 2
```

```
x_chunk = unserialize(worker1)
```

```
partial_means[i] = mean(x_chunk)
```

# An iterator produces data on demand



- Most flexible of the above options
- Natural in pipeline parallel model
- Operate well with high performance IO libraries
- Unfamiliar to R programmers
- Not ideal if you need the whole data set

Simple row based sampling misses the important areas of high density.

