

# The Eval Monad and Strategies

Stephen A. Edwards

Columbia University

Fall 2023



The Eval Monad: rpar and rseq

Speeding up a Sudoku Solver

- Static Partitioning

- Dynamic Partitioning

- Amdahl's Law

Control.DeepSeq: Reducing to Normal Form

- rnf for a Custom Data Type

Evaluation Strategies

- Basic Strategies

- Composing Strategies

- Evaluating Lists in Parallel

Example: The K-Means Problem

- Generating a Data Set

- Parallelizing K-Means

- Performance of Parallel K-Means

Garbage Collected Sparks

Parallelizing Lazy Streams

## The Eval Monad: An Alternative to *seq* and *par*

When in doubt, create a Monad

```
module Control.Parallel.Strategies where

data Eval a = ...

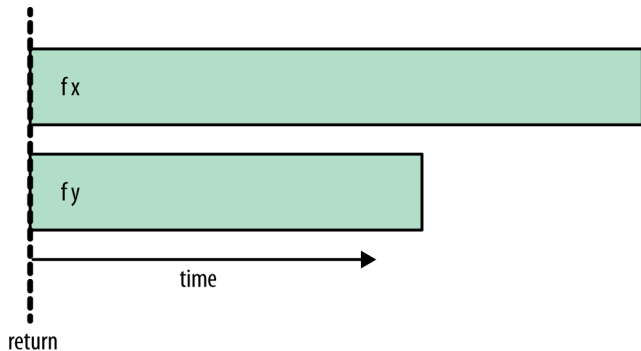
instance Monad Eval where ...

runEval :: Eval a -> a    -- Get the result

rpar  :: a -> Eval a     -- Spark evaluation
rseq  :: a -> Eval a     -- Wait for evaluation to WHNF
```

## rpar/rpar

```
runEval $ do
  a <- rpar (f x)
  b <- rpar (f y)
  return (a, b)
```

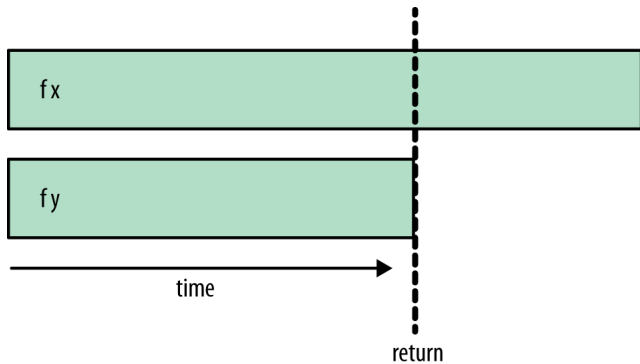


Start parallel evaluation of f x and f y

Return immediately

## rpar/rseq

```
runEval $ do
  a <- rpar (f x)
  b <- rseq (f y)
  return (a, b)
```



Start parallel evaluation of `f x` and `f y`

Wait for `f y` to finish, then return

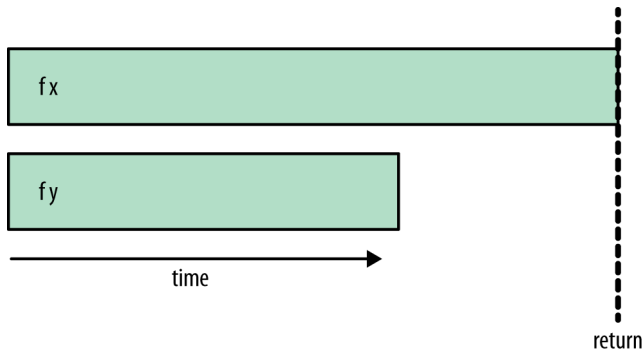
*Do we really know `f y` is faster?*

## rpar/rpar/rseq/rseq

```
runEval $ do
  a <- rpar (f x)
  b <- rseq (f y)
  rseq a
  return (a, b)
```

Equivalent and symmetrical:

```
runEval $ do
  a <- rpar (f x)
  b <- rpar (f y)
  rseq a
  rseq b
  return (a, b)
```



Start parallel evaluation of f x and f y

Wait for both to finish, then return

## Marlow's Sudoku Solver Example (single-threaded)

```
$ wget https://github.com/simonmar/parconc-examples/archive/master.tar.gz
$ tar --strip-components=1 -zxf master.tar.gz \
  parconc-examples-master/{Sudoku.hs,sudoku17.1000.txt}
```

```
import Sudoku ; import Control.Exception
import System.Environment ; import Data.Maybe

main :: IO ()
main = do [f] <- getArgs
          file <- readFile f
          let puzzles    = lines file
              solutions = map solve puzzles
          print (length (filter isJust solutions))
```

```
$ stack ghc -- -O2 sudoku1.hs -rtsopts
$ ./sudoku1 sudoku17.1000.txt +RTS -s
```

## Static Partitioning

```
import Sudoku
import Data.Maybe ; import System.Environment
import Control.Parallel.Strategies ; import Control.DeepSeq
main :: IO ()
main = do puzzles <- getArgs >>= \[f] -> lines <$> readFile f

    let (as,bs) = splitAt (length puzzles `div` 2) puzzles

        solutions = runEval $ do
            as' <- rpar (force (map solve as)) -- To normal
            bs' <- rpar (force (map solve bs)) -- form
            _ <- rseq as'
            _ <- rseq bs'
            return (as' ++ bs')

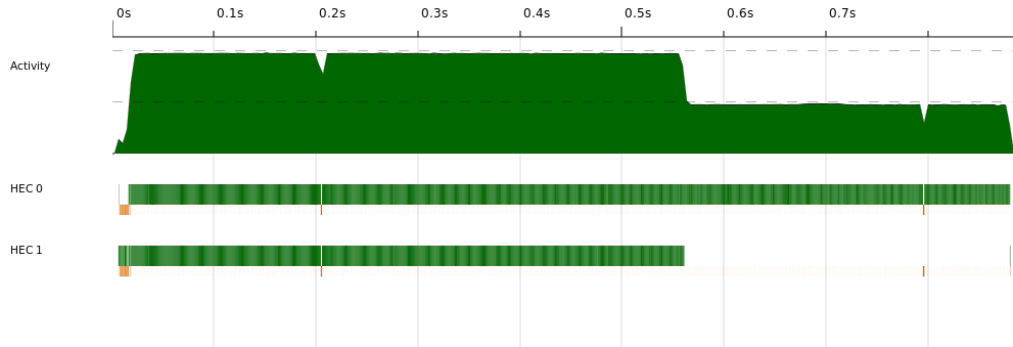
    print (length (filter isJust solutions))
```



# The Problem with Static Partitioning

```
$ stack install parallel
$ stack ghc -- -O2 -Wall sudoku2.hs -threaded -rtsopts -eventlog
$ ./sudoku2 sudoku17.1000.txt +RTS -N2 -s -ls
$ threadscope sudoku2.eventlog
```

Speedup on two cores:  $1.37/0.891 = 1.53\times$



## Better Balancing using parMap

```
import Sudoku
import System.Environment ; import Data.Maybe
import Control.Parallel.Strategies hiding (parMap)

parMap :: (a -> b) -> [a] -> Eval [b]
parMap _ []      = return []
parMap f (a:as) = do b  <- rpar  (f a) -- Spark evaluation of f a
                    bs <- parMap f as -- Recurse
                    return (b:bs)

main :: IO ()
main = do puzzles <- getArgs >>= \[f] -> lines <$> readFile f

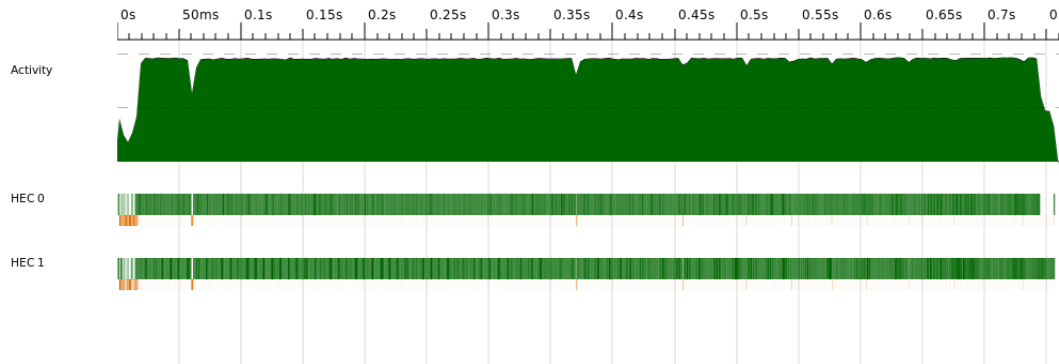
        let solutions = runEval (parMap solve puzzles)

        print (length (filter isJust solutions))
```

# The Advantage of Dynamic Partitioning

```
$ stack ghc -- -O2 -Wall sudoku3.hs -threaded -rtsopts -eventlog  
$ ./sudoku3 sudoku17.1000.txt +RTS -N2 -s -ls  
$ threadscope sudoku3.eventlog
```

Speedup on two cores:  $1.37/0.751 = 1.82\times$  versus  $1.53\times$  with static partitioning



# Amdahl's Law: Why Parallelism is Difficult

Validity of the single processor approach to achieving large scale computing capabilities

by DR. GENE M. AMDAHL  
*International Business Machines Corporation*  
Sunnyvale, California

AFIPS Joint Computer Conference, 1967

$P$  is the parallel fraction of the task

$N$  is the degree of parallelism

$S$  is the speedup

$$S = \frac{1}{(1 - P) + P/N}$$

$\lim_{N \rightarrow \infty} S$  is the depressing part:

If  $P = 0.5$ ,  $S \rightarrow 2$  as  $N \rightarrow \infty$

If  $P = 0.95$ ,  $S \rightarrow 20$  as  $N \rightarrow \infty$

## DeepSeq and Normal Form

Weak Head Normal Form = Top is data constructor or lambda, not application

Normal Form = Data constructors or lambdas all the way down

In Control.DeepSeq,

```
class NFData a where  
  rnf :: a -> ()      -- "Reduce to Normal Form"  
  rnf a = a `seq` ()  -- Default, e.g., for numbers  
  
deepseq :: NFData a => a -> b -> b  
deepseq a b = rnf a `seq` b -- Evaluate a to NF; return b  
  
force   :: NFData a => a -> a  
force a = a `deepseq` a    -- Evaluate a to NF; return it  
  
($!!)   :: NFData a => (a -> b) -> a -> b  
f $!! x = x `deepseq` f x -- evaluate x then f x
```

## Normal Form vs. Weak Head Normal Form

```
Prelude> import Control.DeepSeq
Prelude Control.DeepSeq> let x = [1..10] :: [Int]
Prelude Control.DeepSeq> :sprint x
x = _
Prelude Control.DeepSeq> :sprint x
x = _
Prelude Control.DeepSeq> x `seq` ()
()
Prelude Control.DeepSeq> :sprint x
x = 1 : _
Prelude Control.DeepSeq> (last $ take 3 x) `seq` ()
()
Prelude Control.DeepSeq> :sprint x
x = 1 : 2 : 3 : _
Prelude Control.DeepSeq> x `deepseq` ()
()
Prelude Control.DeepSeq> :sprint x
x = [1,2,3,4,5,6,7,8,9,10]
```

## Roll-your-own NFData

```
import Control.DeepSeq
data Tree a = Empty | Branch (Tree a) a (Tree a)
instance NFData a => NFData (Tree a) where
  rnf Empty = ()
  rnf (Branch l a r) = rnf l `seq` rnf a `seq` rnf r
```

```
*Main> let singleton x = Branch Empty x Empty
*Main> let x = Branch (singleton 'a') 'b' (singleton 'c')
*Main> :sprint x
x = Branch _ 'b' _
*Main> x `seq` ()
()
*Main> :sprint x
x = Branch _ 'b' _
*Main> rnf x
()
*Main> :sprint x
x = Branch (Branch Empty 'a' Empty) 'b' (Branch Empty 'c' Empty)
```

## Evaluation Strategies

“Strategies are a means for modularizing parallel code by **separating the algorithm from the parallelism**. Sometimes they require you to rewrite your algorithm, but once you do so, you will be able to **parallelize it in different ways** just by substituting a new Strategy.” —Simon Marlow

1. Build a lazy data structure representing the computation
2. Apply a Strategy that traverses the computation applying *rpar* and *rseq*

A Strategy: an identity function in the *Eval* monad (`Control.Parallel.Strategies`)

```
type Strategy = a -> Eval a
```



## An Example: A Parallel Strategy for Pairs

```
rpar :: Strategy a           -- Spark evaluation
runEval :: Eval a -> a     -- Evaluate; return value

parPair :: Strategy (a,b)   -- Simple parallel strategy for pairs
parPair (a,b) = do
  a' <- rpar a               -- Spark parallel evaluation of
  b' <- rpar b               -- a and b to WHNF
  return (a',b')
```

```
runEval (parPair (fib 35, fib 36))
```

More elegantly,

```
using :: a -> Strategy a -> a -- In Control.Parallel.Strategies
x `using` s = runEval (s x)    -- Apply s, return x
```

```
(fib 35, fib 36) `using` parPair
```

## Basic Strategies

In Control.Parallel.Strategies:

```
r0      :: Strategy a
r0 x    = return x      -- Do not evaluate x

rseq    :: Strategy a  -- Evaluate to WHNF; wait for completion

rdeepseq :: NFData a => Strategy a -- Fully evaluate then proceed
rdeepseq x = rseq (rnf x) >> return x

rpar    :: Strategy a  -- Spark evaluation (in parallel) to WHNF

rparWith :: Strategy a -> Strategy a
rparWith s x          -- Spark evaluation of x 'using' s
```

## Building Strategies from Strategies

A skeleton for expressing strategies for evaluating tuples:

```
-- In Control.Parallel.Strategies,  
evalTuple2 :: Strategy a -> Strategy b -> Strategy (a, b)  
evalTuple2 sa sb (a, b) = do  a' <- sa a  
                               b' <- sb b  
                               return (a', b')
```

```
parPair :: Strategy (a, b)  
parPair = evalTuple2 rpar rpar -- Spark elements' evaluation to WHNF
```

What if we wanted to fully evaluate the two elements in parallel?

```
parPair :: Strategy a -> Strategy b -> Strategy (a, b)  
parPair sa sb = evalTuple2 (rparWith sa) (rparWith sb)
```

```
parPair rdeepseq rdeepseq (fib 25, fib 26)
```

## parPair rdeepseq rdeepseq (a, b)

“Spark two parallel threads that fully evaluate a and b to normal form”

A cartoon of how this works:

```
parPair rdeepseq rdeepseq (a, b)
  = evalTuple2 (rparWith rdeepseq) (rparWith rdeepseq) (a, b)

  = do a' <- (rparWith rdeepseq) a
        b' <- (rparWith rdeepseq) b
        return (a', b')

  = do a' <- rpar (a `using` \x -> rseq (rnf x) >> return x)
        b' <- rpar (b `using` \x -> rseq (rnf x) >> return x)
        return (a', b')
```

## Evaluating a List in Parallel

In `Control.Parallel.Strategies`,

```
evalList :: Strategy a -> Strategy [a] -- Apply a strategy
evalList _ [] = return []             -- to each list element
evalList s (x:xs) = do x' <- s x
                      xs' <- evalList s xs
                      return (x':xs')
```

```
parList :: Strategy a -> Strategy [a] -- Evaluate each list element
parList s = evalList (rparWith s)     -- in parallel with strategy
```

Combining these to evaluate all list elements to WHNF in parallel:

```
parMap :: (a -> b) -> [a] -> [b]
parMap f xs = map f xs `using` parList rseq
```

## Simpler Parallel Sudoku

```
import Sudoku(solve)
import System.Environment(getArgs)
import Data.Maybe(isJust)
import Control.Parallel.Strategies(using, parList, rseq)

main :: IO ()
main = do [fname] <- getArgs
          puzzles <- lines <$> readFile fname

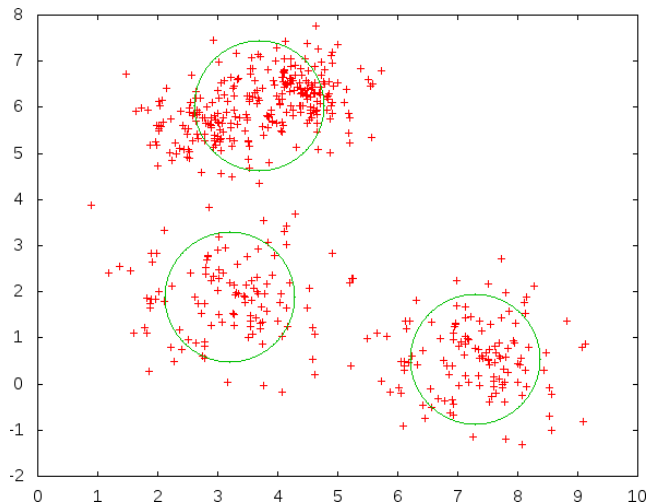
          let solutions = map solve puzzles `using` parList rseq

          print $ length $ filter isJust solutions
```

Note that *rseq* only evaluates to WHNF, but that suffices for Sudoku

About the same performance as the “parMap” version presented earlier

## Example: The K-Means Problem



Lloyd's (approximation) algorithm

Give a number of clusters  $k$ ,

1. Guess a center for each cluster
2. Group points by closest centerpoint
3. Calculate the centroid (average) of each group
4. Repeat steps 3–4 until satisfied

## Example: The K-Means Problem

```
$ wget https://github.com/simonmar/parconc-examples/archive/master.tar.gz
$ tar --strip-components=1 -zxf master.tar.gz \
  parconc-examples-master/kmeans
```

2D points (to simplify visualization) and clusters, in KMeansCore.hs,

```
data Point = Point !Double !Double -- ! disables laziness
```

```
zeroPoint :: Point
```

```
zeroPoint = Point 0 0
```

```
sqDistance :: Point -> Point -> Double -- Distance squared for speed
```

```
sqDistance (Point x1 y1) (Point x2 y2) = ((x1-x2)^2) + ((y1-y2)^2)
```

```
data Cluster = Cluster { clId    :: Int    -- number of this cluster
                        , clCent :: Point  -- centroid of this cluster
                        }
```



## Example: The K-Means Problem

For computing the centroids (average of all points in a cluster), in `kmeans.hs`,

```
data PointSum = PointSum !Int !Double !Double

addToPointSum :: PointSum -> Point -> PointSum
addToPointSum (PointSum count xs ys) (Point x y)
    = PointSum (count+1) (xs + x) (ys + y)

pointSumToCluster :: Int -> PointSum -> Cluster
pointSumToCluster i (PointSum count xs ys) = Cluster {
    clId      = i
    , clCent  = Point (xs / fromIntegral count)
                  (ys / fromIntegral count)
}
```

## 1. Accumulate Points in PointSums for Nearest Centroid

```
assign :: Int -> [Cluster] -> [Point] -> Vector PointSum
assign nclusters clusters points = Vector.create $ do
  vec <- MVector.replicate nclusters (PointSum 0 0 0)

  let addpoint p = do
        let c = nearest p ; cid = clId c
            ps <- MVector.read vec cid
            MVector.write vec cid $! addToPointSum ps p

      mapM_ addpoint points
  return vec
where
  nearest p = fst $ minimumBy (compare `on` snd)
    [ (c, sqDistance (clCent c) p) | c <- clusters ]
```

Vectors are Haskell's fixed-length, random-access arrays that are "mutable" in the right monad. See `Data.Vector` and `Data.Vector.Mutable`

## 2. Create New Clusters from PointSums

```
makeNewClusters :: Vector PointSum -> [Cluster]
makeNewClusters vec =
  [ pointSumToCluster i ps
  | (i,ps@(PointSum count _ _)) <- zip [0..] (Vector.toList vec)
  , count > 0
  ]
```

One step of the algorithm: group by nearest centroid; calculate new centroids

```
step :: Int -> [Cluster] -> [Point] -> [Cluster]
step nclusters clusters points
  = makeNewClusters (assign nclusters clusters points)
```

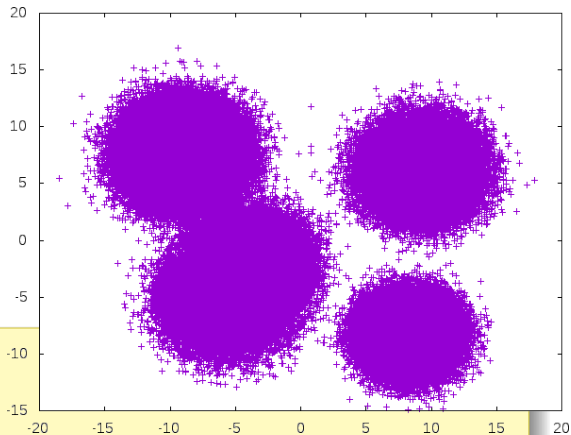
## The Sequential Loop: step until converged or give up

```
kmeans_seq :: Int -> [Point] -> [Cluster] -> IO [Cluster]
kmeans_seq nclusters points clusters =
  let loop :: Int -> [Cluster] -> IO [Cluster]
      loop n clusters | n > tooMany = do
        putStrLn "giving up."
        return clusters
      loop n clusters = do
        printf "iteration %d\n" n
        putStrLn (unlines (map show clusters))
        let clusters' = step nclusters clusters points
            if clusters' == clusters
            then return clusters
            else loop (n+1) clusters'
  in loop 0 clusters
```

```
tooMany = 80
```

## Generating a Data Set

```
$ cabal install normaldistribution  
$ ghc -O2 GenSamples.hs  
$ ./GenSamples 5 50000 100000 1010  
$ ls -l points.bin  
-rw-rw-r-- 1 sedwards sedwards 16M Nov 23 14:58 points.bin  
$ gnuplot -e 'set terminal png;set nokey;plot "points" > points.png
```

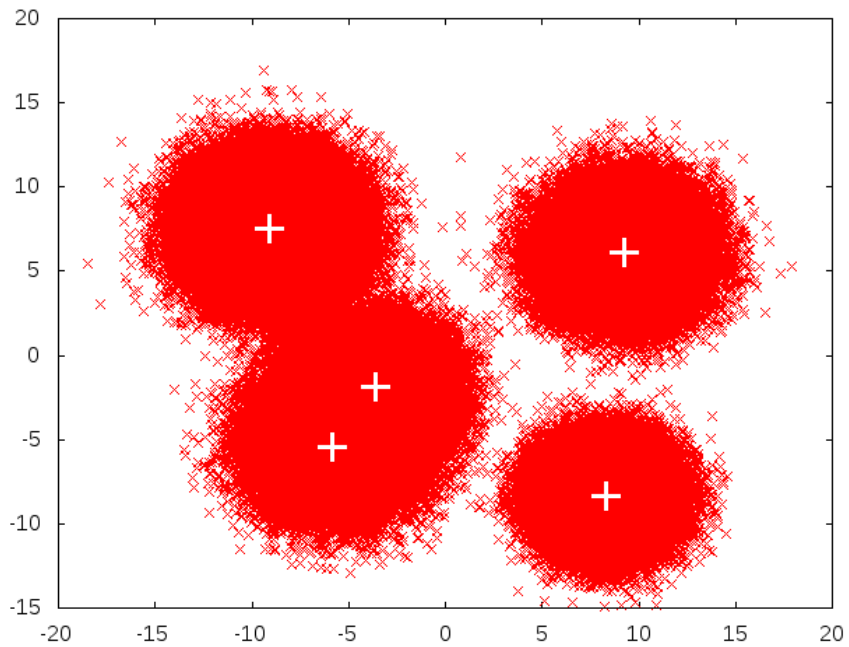


## Compiling and Running K-Means

```
$ stack install monad-par
$ cd kmeans
$ stack ghc -- -O2 -threaded -rtsopts -eventlog kmeans.hs
```

Run it in sequential mode:

```
$ ./kmeans seq
...
iteration 20
Cluster {clId = 0, clCent = Point -5.84359465 -5.46502314}
Cluster {clId = 1, clCent = Point 8.316354592 -8.33043084}
Cluster {clId = 2, clCent = Point -9.06455081 7.561852464}
Cluster {clId = 3, clCent = Point 9.243597731 6.138576051}
Cluster {clId = 4, clCent = Point -3.62170911 -1.82458124}
Total time: 0.73
```



## Parallelizing K-Means

Computing nearest center for each point is the main operation to parallelize. This is a fold with an associative accumulation function *addToPointSum*.

Too many points and not enough work per point for per-point parallelism; overhead would dominate. Better to split work into coarser chunks.

```
split :: Int -> [a] -> [[a]]           -- Divide into numChunks chunks
split numChunks xs = chunk (length xs `quot` numChunks) xs

chunk :: Int -> [a] -> [[a]]          -- Split into n-point chunks
chunk _ [] = []
chunk n xs = let (as,bs) = splitAt n xs in  as : chunk n bs

addPointSums :: PointSum -> PointSum -> PointSum -- Accumulate PointSums
addPointSums (PointSum c1 x1 y1) (PointSum c2 x2 y2)
  = PointSum (c1+c2) (x1+x2) (y1+y2)

combine :: Vector PointSum -> Vector PointSum -> Vector PointSum
combine = Vector.zipWith addPointSums           -- Accumulate vectors
```



## Code for a Parallel *step*

Analyze the chunks in parallel; merge; and make new clusters:

```
parSteps_strat :: Int -> [Cluster] -> [[Point]] -> [Cluster]
parSteps_strat nclusters clusters pointss
  = makeNewClusters $
    foldr1 combine $           -- Merge the results from each chunk
      (map (assign nclusters clusters) pointss) -- Analyze chunks
      `using` parList rseq)    -- in parallel
```

## The Parallel Loop: Divide into chunks; apply parSteps\_strat

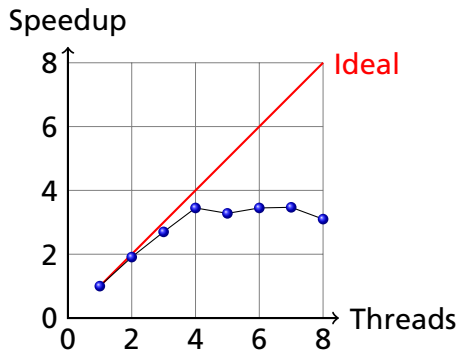
```
kmeans_strat :: Int -> Int -> [Point] -> [Cluster] -> IO [Cluster]
kmeans_strat numChunks nclusters points clusters =
  let chunks = split numChunks points          -- One big change

      loop :: Int -> [Cluster] -> IO [Cluster]
      loop n clusters | n > tooMany = do
        printf "giving up."
        return clusters
      loop n clusters = do
        printf "iteration %d\n" n
        putStr (unlines (map show clusters))
        let clusters' = parSteps_strat nclusters clusters chunks
            if clusters' == clusters
              then return clusters
              else loop (n+1) clusters'
  in loop 0 clusters
```

## Performance of kmeans\_strat on 1–8 Cores

```
./kmeans strat 64 +RTS -N1  
./kmeans strat 64 +RTS -N8
```

Cores	Time (s)	Speedup
1	0.77	1.00
2	0.40	1.91
3	0.29	2.70
4	0.22	3.45
5	0.23	3.28
6	0.22	3.45
7	0.22	3.47
8	0.25	3.10



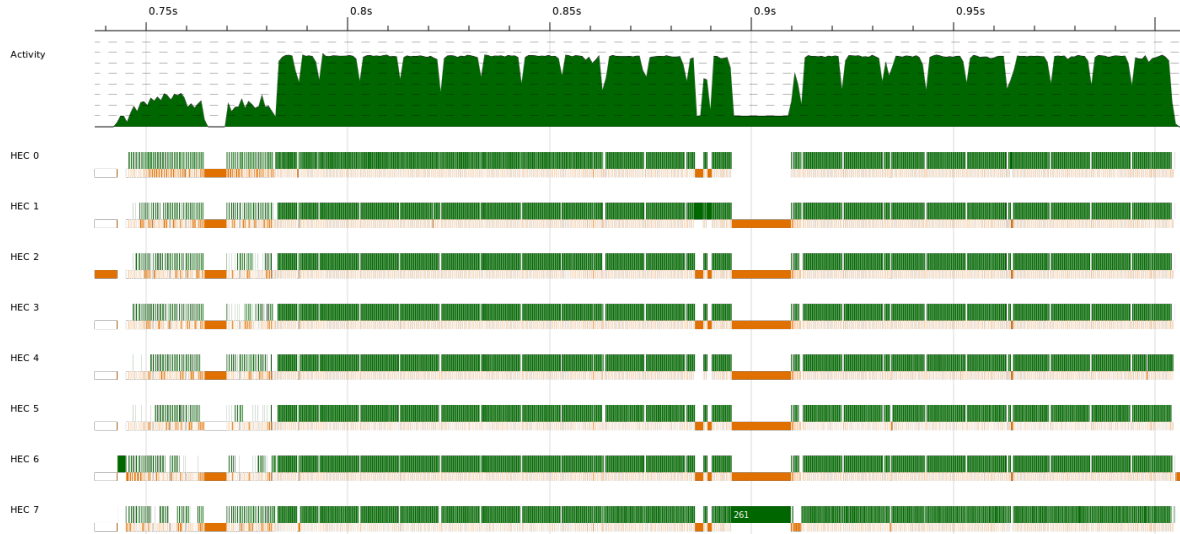
Using "Total time" reported by the program; ignores reading point data

# Threadscope on kmeans\_strat: Overall



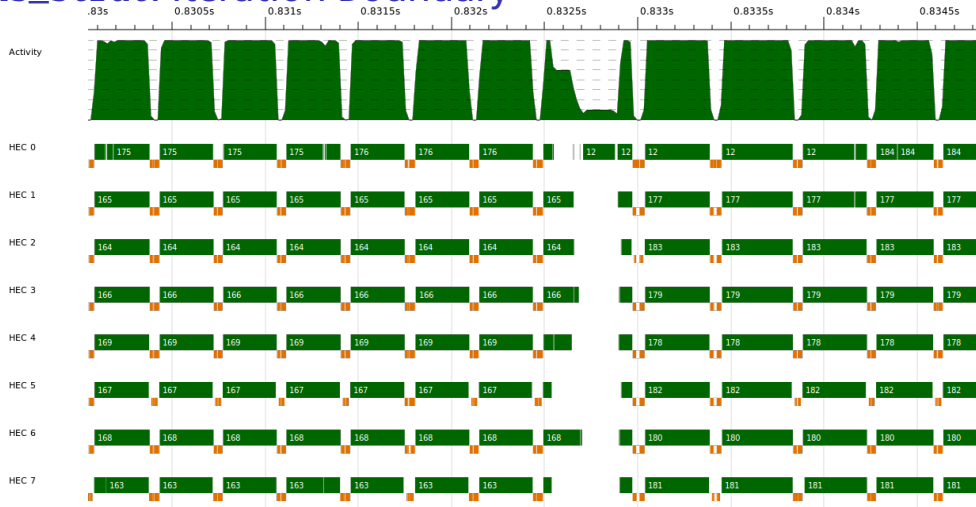
Lots of sequential file processing start: not being counted against speedup

# kmeans\_strat: Parallel Section Only



Off to a difficult start; iterations are periodic hiccups; big garbage collect

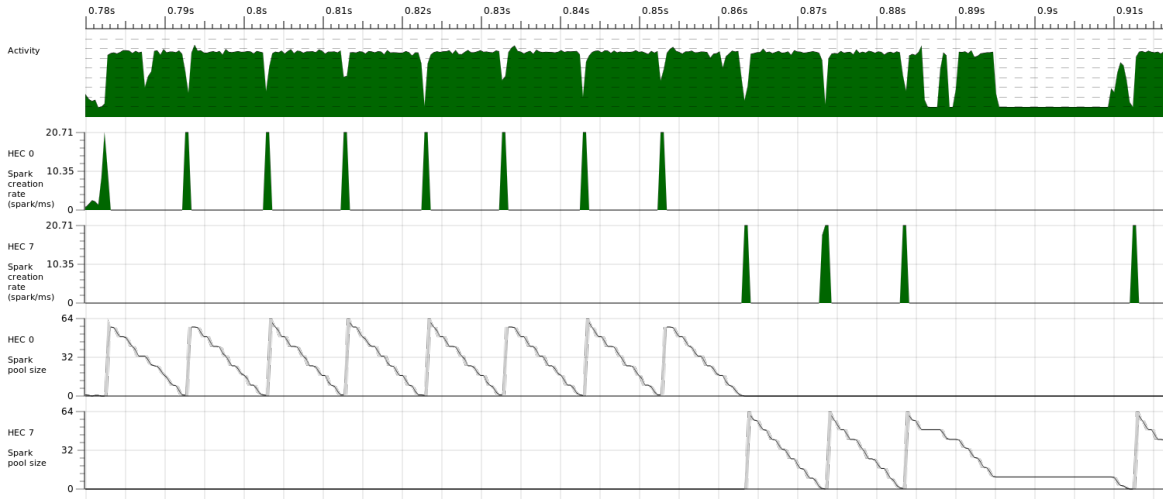
# kmeans\_strat: Iteration Boundary



Program suddenly turns completely sequential; darn Amdahl.

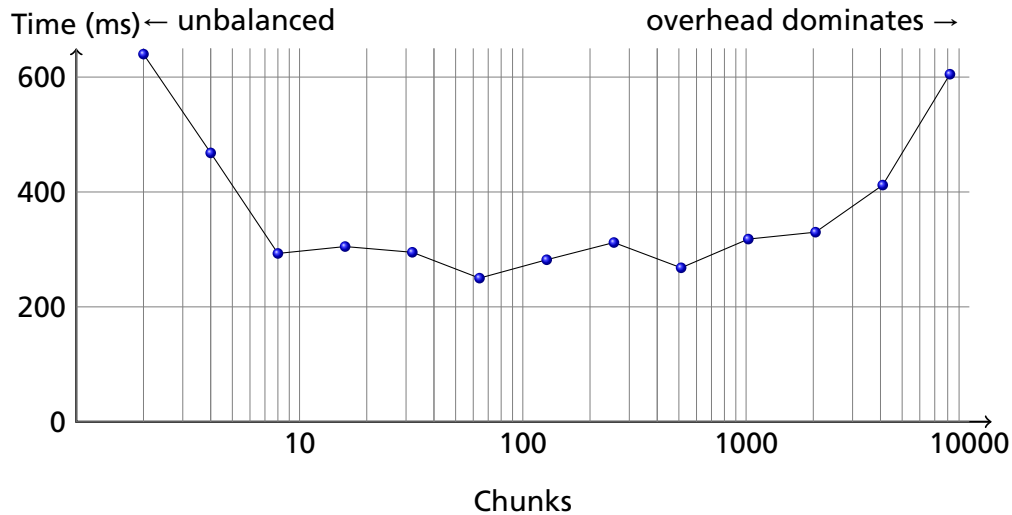
Marlow found printing was a major culprit, but removing it didn't matter on

# kmeans\_strat: Spark creation and conversion



Iteration: sudden spark creation activity in single HEC pool, then slow conversion. Main thread migrated after 8 iterations. (Under the "Traces" tab)

# The Effects of Granularity (N=8)





## GC'ed Sparks and Speculative Parallelism [Marlow p. 48]

parList/evalList creates a new list, which seems wasteful

```
parList :: Strategy a
        -> Strategy [a]
parList s =
    evalList (rparWith s)

evalList :: Strategy a
         -> Strategy [a]
evalList _ [] = return []
evalList s (x:xs) =
    do x' <- s x
       xs' <- evalList s xs
       return (x':xs') -- Cons
```

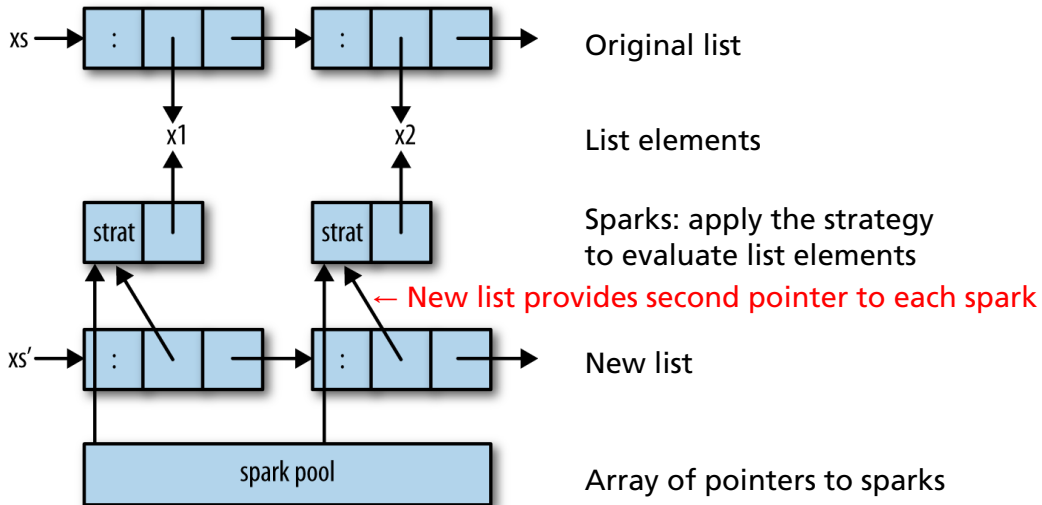
Consider this walk-the-list alternative that “touches” elements with *rparWith*

```
parList :: Strategy a
        -> Strategy [a]
parList strat xs = do go xs
                    return xs

where
    go [] = return ()
    go (x:xs) = do rparWith strat x
                  go xs
```

Doesn't work: each spark created by *rparWith* is garbage-collected because it is never used. **Critical that the result of *rpar/rparWith* be returned.**

# Heap Layout of Working parList: New List Inhibits Spark GC



Marlow, fig. 3-8

## Parallelizing Lazy Streams: RSA Encoder/Decoder from Marlow

```
$ stack ghc -- -O2 -Wall -rtsopts rsa
$ ./rsa encrypt /usr/share/dict/words > /dev/null +RTS -s
  5,089,757,232 bytes allocated in the heap
    3,043,360 bytes copied during GC
      107,888 bytes maximum residency (3 sample(s))
        27,968 bytes maximum slop
          0 MB total memory in use

Total    time    5.740s ( 5.767s elapsed)

$ ls -sh /usr/share/dict/american-english
920K /usr/share/dict/american-english
```

Dictionary file is about 1 MB, but runtime only uses 107,888 bytes maximum because of `Data.ByteString.Lazy.Char8`

# Parallelizing RSA

Sequential implementation:

```
encrypt :: Integer -> Integer -> ByteString -> ByteString
encrypt n e = B.unlines                                     -- Join results
               . map (B.pack . show . power e n . code)   -- Encrypt
               . chunk (size n)                            -- Split
```

First try (rsa1.hs): use *parList rdeepseq*

```
encrypt n e = B.unlines
               . withStrategy (parList rdeepseq)
               . map (B.pack . show . power e n . code)
               . chunk (size n)
```

`withStrategy s e = e `using` s`

## Speedup using *parList rdeepseq*

```
$ stack ghc -- -O2 -Wall -threaded -rtsopts rsa1
$ ./rsa1 encrypt /usr/share/dict/words > /dev/null +RTS -N8 -s
  5,319,033,432 bytes allocated in the heap
  18,619,728 bytes copied during GC
  3,029,464 bytes maximum residency (10 sample(s))
   570,920 bytes maximum slop
     2 MB total memory in use

SPARKS: 9988
      (8254 converted, 1734 overflowed, 0 dud, 0 GC'd, 0 fizzled)

Total    time    14.403s ( 2.991s elapsed)
```

Speedup of 1.92 over sequential (rsa.hs) (4.8× itself)

Maximum memory use now 3 MB (cf. 107 KB): *parList* traverses the whole list.

## Control.Parallel.Strategies.parBuffer: Regulate number of outstanding sparks

*parBuffer 100* creates 100 outstanding sparks; sparks more once consumed

```
parBuffer :: Int -> Strategy a -> Strategy [a]
```

```
encrypt n e = B.unlines    -- rsa2.hs  
    . withStrategy (parBuffer 100 rdeepseq) -- 100 max  
    . map (B.pack . show . power e n . code)  
    . chunk (size n)
```

```
$ ./rsa2 encrypt /usr/share/dict/words > /dev/null +RTS -N8 -s  
506,640 bytes maximum residency (18 sample(s))  
SPARKS: 9988  
        (9987 converted, 0 overflowed, 0 dud, 0 GC'd, 1 fizzled)  
Total   time   12.160s ( 1.641s elapsed)
```

Down to 500 KB residency, 3.5× over sequential, excellent 7.4× self-speedup

# Spark Creation and Pool with parBuffer



HEC5 got the spark creation process  
Spark pool remains around 100  
Hiccups primarily garbage collection

No creation or pools on other HECs  
Gray on graph denotes variance  
About 11% overhead

## RSA Strategies (parList, parBuffer) Compared

Technique	Memory (K)	Sparks					Time (s)	Speedup
		Converted	Overflowed	Dud	GC'ed	Fizzled		
Sequential	105						5.77	1
parList	2958	8254	1734	0	0	0	2.99	1.92
parBuffer	495	9987	0	0	0	1	1.64	3.52

Both generate the same number of sparks

*parList* forces the entire file to be loaded (memory consumption) and generates all the sparks at the beginning (spark pool overflow).