Running ISA in parallel with the snow package

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1 Running ISA in parallel

In this document we show a little example on how to speed up ISA analysis [?, ?, ?, ?] by running the ISA iterations in parallel, on a computer cluster, or a multiprocessor machine.

Since a typical ISA analysis consists of using a range of row/column thresholds and these runs are independent of each other; it is trivial to parallelize the task by performing the iterations for different threshold parameters on different processors or computers. Here we show an example on how to do this easily with the snow [?] and the Rmpi [?] packages:

- > library(isa2)
- > library(snow)
- > library(Rmpi)

We generate some simple in-silico data.

```
> pdata <- isa.in.silico()</pre>
```

Next, we create the MPI cluster, with eight working nodes. You need to have a working MPI installation for this. See more in the documentation of the Rmpi and snow packages.

```
> clu <- makeMPIcluster(8)
```

8 slaves are spawned successfully. 0 failed.

Now we load the isa2 package on all nodes and distribute the input data to them.

```
> invisible(clusterEvalQ(clu, library(isa2)))
> clusterExport(clu, "pdata")
```

Next, we create a big matrix in which each row is a combination of the threshold parameters. This will be needed for the parallel run.

```
> thr <- seq(1,3,by=0.2)
> thr.list <- expand.grid(thr, thr)</pre>
```

First we run the ISA on a single processor only, and measure the running time.

```
> system.time(modules <- isa(pdata[[1]], thr.row=thr, thr.col=thr))
         system elapsed
   user
 24.114
          1.221 25.839
  Let us now do a parallel run, again, with measuring the running time. If
you are really running this on multiple CPUs, then it is much faster.
> system.time(modules.par <- parApply(clu, thr.list, 1, function(x) {
   isa(pdata[[1]], thr.row=x[1], thr.col=x[2])
 }))
   user
         system elapsed
  0.331
          0.208 10.537
  Finally, stop the cluster.
> stopCluster(clu)
[1] 1
```

2 Session information

The version number of R and packages loaded for generating this vignette were:

- R version 3.2.0 beta (2015-04-02 r68137), x86_64-apple-darwin10.8.0
- Locale: en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
- Base packages: base, datasets, graphics, grDevices, methods, stats, utils
- \bullet Other packages: isa
20.3.4,Rmpi0.6-5,snow0.3-13
- Loaded via a namespace (and not attached): grid 3.2.0, lattice 0.20-31, tools 3.2.0

References

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- [Ihmels, 2002] Ihmels, J., Friedlander, G., Bergmann, S., Sarig, O., Ziv, Y., Barkai, N. (2002). Revealing modular organization in the yeast transcriptional network. *Nat Genet*, page 370–7.
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