Merging regions

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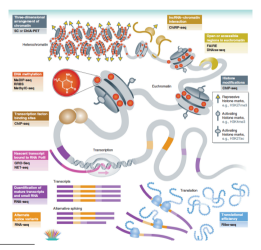
- Problem setting
 - Background
 - Question

Proposed method

Second Example
Second Example

Results

High-Throughput Genomics Panorama¹



¹Wendy Weijia Soon, Manoj Hariharan, and Michael P. Snyder. "High-throughput sequencing for biology and medicine". In: Molecular Systems Biology 9.1 (). UR

What is common?

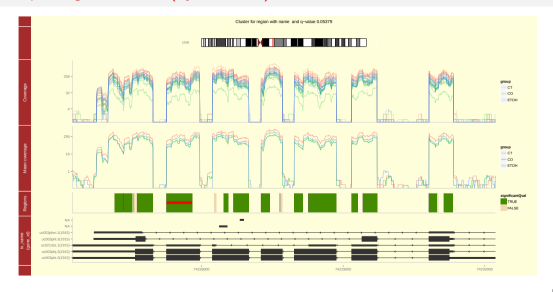
- Measurements along the genome (sometimes summarized)
- Two or more groups of samples
- Typical question: are there differences between the groups?
 - ightharpoonup ightharpoonup Find the candidate regions.

Issue: regions might be highly fragmented.

Why?

- Biological reasons: regions correspond to two exons (intron is the cause).
- Measurement not ideal: coverage dips.

Example region cluster (by distance)



Question

Are two adjacent regions similar?

- Can we *link* them?
- Are regions overlapping the same exon more frequently linked?

Translating framework

- What is measured?
 - ► Coverage =: *Y*
 - ▶ Transformed: $log_2(Y + 32)$
- Individual (cluster of measurements) → sample
- Repeated visits → individual base pairs (from a given chromosome)
 - Note that the data is correlated!

Consider a region pair:

- region1: first region
- 2 regionM: middle part
- region2: second region

Proposed method

Model for sample *i*:

$$\log_2(Y_{ijk} + 32) = \alpha + \beta_1 \text{sampleDepth}_i + \beta_2 \text{group}_j + \beta_3 \text{region}_k + \epsilon$$

Using region1 as the reference, we want to test β_3 (region₂) = 0.

Data sets used

- derHippo: RNA-seq brain hippocampus study
- 25 samples
- 3 groups: CO, ETOH and CT

```
chr6 890 pairs
```

300 (\sim 33.7%) with regions 1 & 2 having a width greater than 1, region M < 250

chr22 573 pairs

187 (\sim 32.6%) passing the filtering

Example:

chr 6, chose the largest cluster, then the pair starting with the largest region from the cluster.

Example region pair



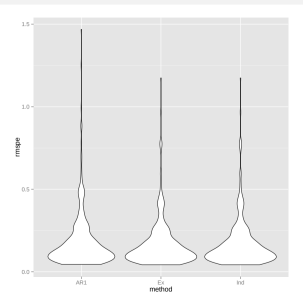
Data

```
pairs[i2,]
##
        start1 end1 startM endM start2 end2 cluster
## 552 74227546 74227657 74227658 74227752 74227753 74227934
                                                         168
##
      width1 widthM width2 widthNoM
## 552 112 95 182
                            294
dim(covdata[[i2]])
## [1] 9725 6
head(covdata[[i2]], n = 1)
##
    base region sample coverage sampleDepth group
       1 region1
                  CO1
                        7.401 28.25
## 1
```

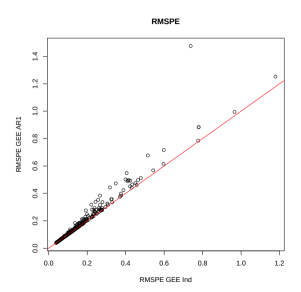
GEE - AR1

```
##
## Call:
## geeglm(formula = coverage ~ sampleDepth + group + region, family = gaussian,
      data = covdata[[i2]], id = sample, corstr = "ar1")
##
   Coefficients:
                Estimate Std.err Wald Pr(>|W|)
## (Intercept) -11.02816
                         3.69521
                                   8.91 0.0028 **
## sampleDepth 0.66137
                          0.12836
                                   26.55 2.6e-07 ***
## groupCO -0.76057 0.10828 49.34 2.2e-12 ***
## groupETOH -0.33737 0.10847
                                    9.67 0.0019 **
## regionregionM -1.79501 0.10960 268.25 < 2e-16 ***
## regionregion2 -0.11747 0.00614 366.09 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Estimated Scale Parameters:
             Estimate Std.err
## (Intercept) 0.458 0.0224
##
## Correlation: Structure = ar1 Link = identity
## Estimated Correlation Parameters:
        Estimate Std.err
## alpha 0.971 0.00691
## Number of clusters: 25 Maximum cluster size: 389
```

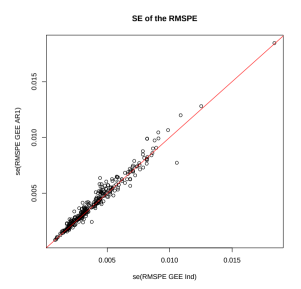
Root mean squared prediction error (RMSPE) chr6



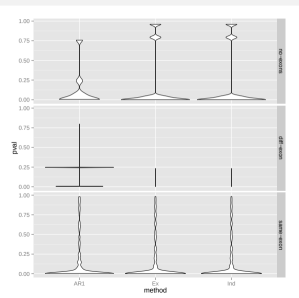
RMSPE chr6: GEE AR1 vs GEE Ind



SE RMSPE chr6: GEE AR1 vs GEE Ind



P-values by exon status: chr6



Test result by exon status

ullet Adjust for multiple testing by using q-value < 0.10

BothAccept	Ar1Accept	IndAccept	BothReject	chr	ExonStatus
0	1	2	7	chr6	no-exons
0	1	0	13	chr6	diff-exon
19	24	23	210	chr6	same-exon
0	4	1	11	chr22	no-exons
0	0	0	2	chr22	diff-exon
5	15	14	135	chr22	same-exon

Conclusions

- With longer region pairs, fitting GEE takes a significant amount of time.
- GEE with Independence working correlation had lower RMSPE.
- For pairs sharing an exon, 11-20% were linked.
- Method works, but is restrictive.

References

- Project code and results: https://github.com/lcolladotor/756final_code
- A. Frazee, S. Sabunciyan, K. D. Hansen, R. A. Irizarry, and J. T. Leek (2013).
 Differential expression analysis of rna-seq data at single base resolution, Biostatistics, recently accepted.
- L. Collado-Torres, A. Frazee, M. Love, R. A. Irizarry, A. E. Jaffe, J. T. Leek (2013). derfinder: Software for annotation-agnostic RNA-seq differential expression analysis. Manuscript in preparation.
- derfinder package https://github.com/lcolladotor/derfinder

Thank you!