A "big data" strategy relevant to transcriptional regulatory networks

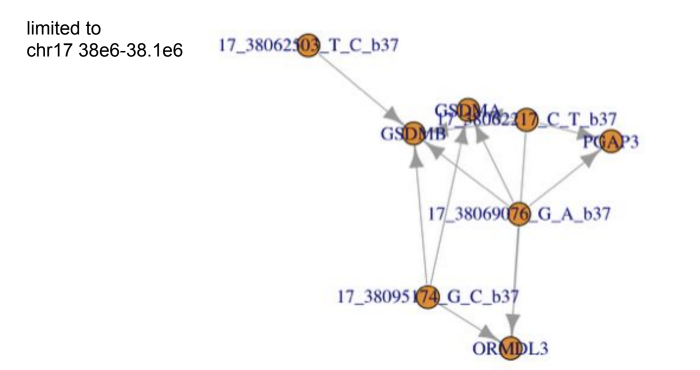
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Road map

- A view of the task
- mongolite package to resolve queries
- TxRegInfra package to organize interfaces

Organizing TFs, TFBS, tissue-specific eQTL results (SNP:gene associations declared in GTEx v6)

SNP:Gene assoc in GTEx lung within STAT1 binding sites by FIMO



this is fairly primitive! want to know more about edges, variant characteristics, etc.

how to scratch the surface even to this extent is not clear/shared ... or ... neo4j? graphQL?

Some guiding principles

- Objective is to smooth the path from large genomic reference resources or experimental results to exploratory and confirmatory analysis activities -- leading to a new resource or interface
- Principle 1: record provenance and bind it to the resource
- Principle 2: minimize modification to original resource
- Principle 3: consider how the X[G, S] idiom can be supported
 - G refers to 'features', S refers to 'samples'
 - The underlying structure need not be 'rectangular'
- Principle 4: produce a useful 'colData'
- Principle 5: support subsetByOverlaps

Principles 2+4 for eQTL+FP+GWAS

Show 10 centries

Search:

basetypemidSpleen_allpairs_Y_eQTLSpleeneQTLallpairs_v7Stomach_allpairs_Y_eQTLStomacheQTLallpairs_v7Testis_allpairs_Y_eQTLTestiseQTLallpairs_v7Thyroid_allpairs_Y_eQTLThyroideQTLallpairs_v7Verus_allpairs_Y_eQTLVeruseQTLallpairs_v7Vagina_allpairs_Y_eQTLVaginaeQTLallpairs_v7Vhole_Blood_allpairs_Y_eQTLWholeeQTLBlood_allpairs_v7CD14_DS17215_ng19_FPCD14FPDS17215_ng19CD34_DS12274_ng19_FPCD34FPDS12274_ng19	
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CD19_DS17186_hg19_FP CD19 FP DS17186_hg19	
CD34_DS12274_hg19_FP CD34 FP DS12274_hg19	
Showing 41 to 50 of 2,640 entries Previous 1 4 5 6 264	Next

searchability is valuable

Show 10 centries					Search: lu	ng	
	base	\$	type		mid	I	÷
Lung_allpairs_v7_eQTL	Lung	eQTL		all	allpairs_v7		
fLung_DS14724_hg19_FP	fLung	FP		DS	14724_hg19		
Showing 1 to 2 of 2 entries (filtered from 2,640 total entries)					Previous	1	Next

Classes are useful -- this is not quite mature, 'RaggedExperiment' extension ... accommodate irregular 'assays'

```
> rme
class: RaggedMongoExpt
dim: 2640 2640
assays(0):
rownames: NULL
colnames(2640): Adipose_Subcutaneous_allpairs_v7_eQTL
  Adipose_Visceral_Omentum_allpairs_v7_eQTL ...
  iPS_19_11_DS15153_hg19_FP vHMEC_DS18406_hg19_FP
colData names(6): base type ... type mid
. .....
```

MongoDB+mongolite for each 'file' as 'document' rme = f(URL, db, colData)... URL here is to AWS

```
> rme@con
<Mongo collection> 'test'
$aggregate(pipeline = "{}", options = "{\"allowDiskUse\":true}", handler = NULL, pagesize = 1000, it
erate = FALSE)
$count(query = "{}")
$disconnect(gc = TRUE)
$distinct(key, guery = "{}")
$drop()
$export(con = stdout(), bson = FALSE, query = "{}", fields = "{}", sort = "{\"_id\":1}")
$find(query = "{}", fields = "{\"_id\":0}", sort = "{}", skip = 0, limit = 0, handler = NULL, pagesi
ze = 1000)
$import(con, bson = FALSE)
$index(add = NULL, remove = NULL)
$info()
$insert(data, pagesize = 1000, stop on error = TRUE, ...)
$iterate(query = "{}", fields = "{\"_id\":0}", sort = "{}", skip = 0, limit = 0)
$mapreduce(map, reduce, query = "{}", sort = "{}", limit = 0, out = NULL, scope = NULL)
$remove(query, just_one = FALSE)
$rename(name, db = NULL)
$replace(query, update = "{}", upsert = FALSE)
$run(command = "{\"ping\": 1}", simplify = TRUE)
$update(query, update = "{\"$set\":{}}", filters = NULL, upsert = FALSE, multiple = FALSE)
~
```

Task: use the resource to learn about eQTLs in two tissues

- Interested in subcutaneous adipose and lung
- We'll look in a region on chr17

```
> tiss = c("Adipose_Subcutaneous_allpairs_v7_eQTL",
   "Lung_allpairs_v7_eQTL")
+
> grange = GRanges("17", IRanges(38e6,width=1e6))
> ans = lapply(tiss, function(x) sbov(rme[,x], grange))
...Warning messages:
1: In sbov(rme[, x], grange) : genome is not set for for guery GRanges
2: In sbov(rme[, x], grange) : genome is not set for for guery GRanges
> sapply(ans,length)
[1] 2644 2049
> names(ans) = tiss
> sapply(ans,length)
Adipose_Subcutaneous_allpairs_v7_eQTL
                                                       Lung_allpairs_v7_eQTL
                                 2644
                                                                        2049
```

> he	ad(ans[[1]])							
GRang	es object	with 6 ran	nges and	16 met	adata c	olumns	:		
	seqnames	ranges	strand	1	g	ene_id			variant_id
	<rle></rle>	<iranges></iranges>	<rle></rle>	1	<f< td=""><td>actor></td><td></td><td></td><td><factor></factor></td></f<>	actor>			<factor></factor>
[1]	17	38001559	*	ENSG0	0000073	605.14	1	7_38001	559_A_G_b37
[2]	17	38001571	*	ENSG	0000014	1744.3	1	7_38001	571_G_A_b37
[3]	17	38001659	*	ENSG0	0000073	605.14	17	_380016	59_T_TC_b37
[4]	17	38004929	*	ENSG0	0000073	605.14	17_38	8004929_	GATTT_G_b37
[5]	17	38004929	*						GATTT_G_b37
[6]	17	38004929	*	ENSG					GATTT_G_b37
	tss_dista	ance ma_sam		a_count		maf pva	al_nom	ninal	slope
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[1]		3344	50	51	0.0662	338 (0.0014	7706 -0	.271482
[2]		7337	16		0.0221	and the state of the	00027	0805 0	.359381
[3]		3244	50		0.0662			7706 -0	
[4]		9974	260		0.428				.122441
[5]		8925	260	330	0.428				.106083
[6]	-7	9066	260	330	0.428	571 1.	45895	ie-06 0	.248438
	slope_s			alue	chr		pos	A1	A2
	<numeric:< td=""><td></td><td></td><td></td><td></td><td></td><td></td><td>factor></td><td><factor></factor></td></numeric:<>							factor>	<factor></factor>
	0.084649		22269301		17			A	G
	0.097578		54545107	10 10 10 10 10 10 10 10 10 10 10 10 10 1	17		100 C	G	A
	0.084649		22269301		17			т	TC
	0.037228		03818591		17			GATTT	G
	0.023989				17			GATTT	G
[6]		7 0.000236	49124524	8636	17		+929	GATTT	G
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	 [4] b37 Adipose_Subcutaneous_allpairs_v7_eQTL [5] b37 Adipose_Subcutaneous_allpairs_v7_eQTL 								
[5]									
[6]	b37	Adipose_S	ubcutane	ous_all	pairs_v	7_eQTL			

seqinfo: 1 sequence from an unspecified genome; no seqlengths

annotation from GTEx

Are there tissue-specific SNP-associated genes?

```
> names(ans) = c("subcut", "lung")
> setdiff(ans$subcut$gene_id, ans$lung$gene_id)
 [1] "ENSG00000126368.5" "ENSG00000108306.7"
                                             "ENSG00000214546.3"
 [4] "ENSG00000173991.5" "ENSG00000229028.2" "ENSG00000131746.8"
 [7] "ENSG00000221852.4" "ENSG0000126337.9" "ENSG00000131759.13"
[10] "ENSG00000186832.4" "ENSG00000128422.11" "ENSG00000173812.6"
> setdiff(ans$lung$gene id, ans$subcut$gene id)
[1] "ENSG00000108298.5" "ENSG00000266753.2" "ENSG00000265799.1"
[4] "ENSG00000204889.6" "ENSG00000270145.1" "ENSG00000196859.3"
[7] "ENSG00000265666.1" "ENSG0000186847.5" "ENSG00000171346.9"
```

```
> ulung = setdiff(ans$lung$gene_id, ans$subcut$gene_id)
> uluna
[1] "ENSG00000108298.5" "ENSG0000266753.2" "ENSG00000265799.1"
[4] "ENSG00000204889.6" "ENSG00000270145.1" "ENSG00000196859.3"
[7] "ENSG00000265666.1" "ENSG00000186847.5" "ENSG00000171346.9"
> ul = gsub("\..*", "", ulung)
> ul
[1] "ENSG00000108298" "ENSG00000266753" "ENSG00000265799" "ENSG0000020
[5] "ENSG00000270145" "ENSG00000196859" "ENSG00000265666" "ENSG0000018
[9] "ENSG00000171346"
> na.omit(mapIds(org.Hs.eg.db, keys=ul, keytype="ENSEMBL", column="GO"
'select()' returned 1:many mapping between keys and columns
ENSG00000108298 ENSG00000204889 ENSG00000196859 ENSG00000186847 ENSG00
   "GO:0000184"
                   "GO:0005198"
                                   "GO:0005198"
                                                    "GO:0005200"
                                                                    "GO
attr(, "na.action")
ENSG00000266753 ENSG00000265799 ENSG00000270145 ENSG00000265666
              2
                              3
                                               5
                                                               7
attr(, "class")
[1] "omit"
> mapIds(GO.db, keys=.Last.value, keytype="GOID", column="TERM")
'select()' returned 1:1 mapping between keys and columns
                                                            GO:0000184
"nuclear-transcribed mRNA catabolic process, nonsense-mediated decay"
                                                            GO:0005198
                                        "structural molecule activity"
                                                            GO:0005198
                                        "structural molecule activity"
                                                            GO:0005200
                             "structural constituent of cytoskeleton"
                                                            GO:0005200
                             "structural constituent of cytoskeleton"
```

```
primitive approach to
acquiring GO
mappings for eGenes
unique to lung in this
region
```

For TFBS in this interval

See the TFutils package fimo_granges()

If time permits

indexedGF for GWAS summaries

vjcitn.shinyapps.io/ca43k