Advanced Data Manipulation: Exercises

Exercise set 1

- 1. Display the data where the gene ontology biological process (the bp variable) is "leucine biosynthesis" (case-sensitive) and the limiting nutrient was Leucine. (Answer should return a 24-by-7 data frame 4 genes × 6 growth rates).
- 2. Which tene/rate combinations had high expression (in the top 1% of expressed genes)? *Hint:* see ?quantile and try quantile(ydat\$expression, probs=.99) to see the expression value which is higher than 99% of all the data, then filter() based on that. Try wrapping your answer with a View() function so you can see the whole thing. What does it look like those genes are doing? Answer should return a 1971-by-7 data frame.

Exercise set 2

- 1. First, re-run the command you used above to filter the data for genes involved in the "leucine biosynthesis" biological process and where the limiting nutrient is Leucine.
- 2. Wrap this entire filtered result with a call to arrange() where you'll arrange the result of #1 by the gene symbol.
- 3. Wrap this entire result in a View() statement so you can see the entire result.

Exercise set 3

Here's a warm-up round. Try the following.

1. Show the limiting nutrient and expression values for the gene ADH2 when the growth rate is restricted to 0.05. *Hint:* 2 pipes: filter and select.

```
## # A tibble: 6 X 2
##
      nutrient expression
##
         <chr>
                     <dbl>
## 1
       Glucose
                      6.28
## 2
       Ammonia
                      0.55
## 3 Phosphate
                     -4.60
       Sulfate
                     -1.18
## 5
                      4.15
       Leucine
## 6
        Uracil
```

2. What are the four most highly expressed genes when the growth rate is restricted to 0.05 by restricting glucose? Show only the symbol, expression value, and GO terms. *Hint:* 4 pipes: filter, arrange, head, and select.

```
## # A tibble: 4 X 4
##
     symbol expression
                                         bp
                                                                          mf
##
      <chr>
                 <dbl>
                                      <chr>>
## 1
       ADH2
                  6.28
                              fermentation* alcohol dehydrogenase activity
## 2 HSP26
                  5.86 response to stress*
                                                   unfolded protein binding
## 3
      MLS1
                  5.64
                           glyoxylate cycle
                                                  malate synthase activity
## 4
       HXT5
                  5.56
                           hexose transport glucose transporter activity*
```

3. When the growth rate is restricted to 0.05, what is the average expression level across all genes in the "response to stress" biological process, separately for each limiting nutrient? What about genes in the "protein biosynthesis" biological process? *Hint:* 3 pipes: filter, group_by, summarize.

```
## # A tibble: 6 X 2
##
      nutrient meanexp
##
         <chr>>
                  <dbl>
## 1
       Ammonia
                  0.943
## 2
       Glucose
                  0.743
## 3
       Leucine
                  0.811
## 4 Phosphate
                  0.981
## 5
                  0.743
       Sulfate
## 6
        Uracil
                  0.731
## # A tibble: 6 X 2
##
      nutrient meanexp
##
         <chr>>
                  <dbl>
## 1
       Ammonia
                -1.613
## 2
       Glucose
                -0.691
## 3
       Leucine
                -0.574
## 4 Phosphate
                -0.750
## 5
       Sulfate
                -0.913
## 6
        Uracil -0.880
```

Exercise set 4

That was easy, right? How about some tougher ones.

First, some review. How do we see the number of distinct values of a variable? Use n_distinct() within a summarize() call.

```
ydat %>% summarize(n_distinct(mf))

## # A tibble: 1 X 1

## `n_distinct(mf)`

## <int>
## 1 1086
```

1. Which 10 biological process annotations have the most genes associated with them? What about molecular functions? *Hint:* 4 pipes: group_by, summarize with n_distinct, arrange, head.

```
## # A tibble: 10 X 2
##
                                                                   bp
                                                                          n
##
                                                                <chr> <int>
## 1
                                                                        269
                                          biological process unknown
## 2
                                                protein biosynthesis
                                                                        182
## 3
                                protein amino acid phosphorylation*
                                                                         78
## 4
                                               protein biosynthesis*
                                                                         73
## 5
                             cell wall organization and biogenesis*
                                                                         64
## 6
      regulation of transcription from RNA polymerase II promoter*
                                                                         49
## 7
                             nuclear mRNA splicing, via spliceosome
                                                                         47
## 8
                                                         DNA repair*
                                                                         44
## 9
                                                aerobic respiration*
                                                                         42
## 10
                                                                         42
                                              ER to Golgi transport*
## # A tibble: 10 X 2
```

```
##
                                             mf
                                                    n
##
                                          <chr> <int>
## 1
                   molecular function unknown
                                                  886
##
  2
          structural constituent of ribosome
                                                  185
##
  3
                              protein binding
                                                  107
## 4
                                   RNA binding
                                                   63
## 5
                              protein binding*
                                                   53
## 6
                                  DNA binding*
                                                   44
##
                 structural molecule activity
                                                   43
## A
                              GTPase activity
                                                   40
## 9
      structural constituent of cytoskeleton
                                                   39
## 10
                transcription factor activity
                                                   38
```

2. How many distinct genes are there where we know what process the gene is involved in but we don't know what it does? *Hint:* 3 pipes; filter where bp!="biological process unknown" & mf=="molecular function unknown", and after selecting columns of interest, pipe the output to distinct(). The answer should be 737, and here are a few:

```
## # A tibble: 737 X 3
##
      symbol
                                                                             bp
##
       <chr>
                                                                          <chr>>
## 1
        SFR2
                                                         ER to Golgi transport
## 2
        EDC3
                                        deadenylylation-independent decapping
## 3
        PER1
                                                 response to unfolded protein*
       PEX25
##
                                      peroxisome organization and biogenesis*
## 5
        BNI5
                                                                   cytokinesis*
## 6
       CSN12 adaptation to pheromone during conjugation with cellular fusion
## 7
       SEC39
                                                             secretory pathway
## 8
        ABC1
                                                       ubiquinone biosynthesis
## 9
       PRP46
                                       nuclear mRNA splicing, via spliceosome
## 10
        EMAM
                                   mitochondrion organization and biogenesis*
## # ... with 727 more rows, and 1 more variables: mf <chr>
```

3. When the growth rate is restricted to 0.05 by limiting Glucose, which biological processes are the most upregulated? Show a sorted list with the most upregulated BPs on top, displaying the biological process and the average expression of all genes in that process rounded to two digits. *Hint:* 5 pipes: filter, group_by, summarize, mutate, arrange.

```
## # A tibble: 881 X 2
##
                                                    bp meanexp
##
                                                 <chr>
                                                         <dbl>
## 1
                                                          6.28
                                        fermentation*
## 2
                                                          5.29
                                     glyoxylate cycle
##
  3
      oxygen and reactive oxygen species metabolism
                                                          5.04
## 4
                                  fumarate transport*
                                                          5.03
## 5
                            acetyl-CoA biosynthesis*
                                                          4.32
## 6
                                      gluconeogenesis
                                                          3.64
## 7
                                                          3.57
                           fatty acid beta-oxidation
## 8
                                    lactate transport
                                                          3.48
## 9
                                 carnitine metabolism
                                                          3.30
## 10
                                  alcohol metabolism*
                                                          3.25
## # ... with 871 more rows
```

4. Group the data by limiting nutrient (primarily) then by biological process. Get the average expression for all genes annotated with each process, separately for each limiting nutrient, where the growth rate is restricted to 0.05. Arrange the result to show the most upregulated processes on top. The initial

result will look like the result below. Pipe this output to a View() statement. What's going on? Why didn't the arrange() work? *Hint:* 5 pipes: filter, group_by, summarize, arrange, View.

```
## Source: local data frame [5,257 x 3]
## Groups: nutrient [6]
##
##
       nutrient
                                                              bp meanexp
##
          <chr>
                                                            <chr>
                                                                    <dbl>
## 1
        Ammonia
                                           allantoate transport
                                                                     6.64
## 2
        Ammonia
                                          amino acid transport*
                                                                     6.64
## 3
      Phosphate
                                glycerophosphodiester transport
                                                                     6.64
        Glucose
## 4
                                                   fermentation*
                                                                     6.28
## 5
        Ammonia
                                             allantoin transport
                                                                     5.56
## 6
        Glucose
                                                                     5.29
                                                glyoxylate cycle
## 7
        Ammonia
                                            proline catabolism*
                                                                     5.14
## 8
                                                  urea transport
                                                                     5.14
        Ammonia
## 9
        Glucose oxygen and reactive oxygen species metabolism
                                                                     5.04
## 10
        Glucose
                                             fumarate transport*
                                                                     5.03
## # ... with 5,247 more rows
```

5. Let's try to further process that result to get only the top three most upregulated biolgocal processes for each limiting nutrient. Google search "dplyr first result within group." You'll need a filter(row_number()....) in there somewhere. Hint: 5 pipes: filter, group_by, summarize, arrange, filter(row_number().... Note: dplyr's pipe syntax used to be %.% before it changed to %>%. So when looking around, you might still see some people use the old syntax. Now if you try to use the old syntax, you'll get a deprecation warning.

```
## Source: local data frame [18 x 3]
  Groups: nutrient [6]
##
##
       nutrient
                                                              bp meanexp
##
          <chr>
                                                            <chr>
                                                                    <dbl>
## 1
        Ammonia
                                           allantoate transport
                                                                     6.64
## 2
        Ammonia
                                          amino acid transport*
                                                                     6.64
## 3
      Phosphate
                               glycerophosphodiester transport
                                                                     6.64
## 4
        Glucose
                                                                     6.28
                                                   fermentation*
## 5
        Ammonia
                                             allantoin transport
                                                                     5.56
## 6
        Glucose
                                                                     5.29
                                                glyoxylate cycle
## 7
        Glucose oxygen and reactive oxygen species metabolism
                                                                     5.04
## 8
         Uracil
                                             fumarate transport*
                                                                     4.32
## 9
      Phosphate
                                 vacuole fusion, non-autophagic
                                                                     4.20
## 10
        Leucine
                                                   fermentation*
                                                                     4.15
## 11 Phosphate
                         regulation of cell redox homeostasis*
                                                                     4.03
## 12
        Leucine
                                             fumarate transport*
                                                                     3.72
## 13
        Leucine
                                                glyoxylate cycle
                                                                     3.65
## 14
        Sulfate
                                         protein ubiquitination
                                                                     3.40
## 15
        Sulfate
                                            fumarate transport*
                                                                     3.27
## 16
         Uracil
                                          pyridoxine metabolism
                                                                     3.11
## 17
                                         asparagine catabolism*
         Uracil
                                                                     3.06
                                  sulfur amino acid metabolism*
## 18
                                                                     2.69
        Sulfate
```

6. There's a slight problem with the examples above. We're getting the average expression of all the biological processes separately by each nutrient. But some of these biological processes only have a single gene in them! If we tried to do the same thing to get the correlation between rate and expression, the calculation would work, but we'd get a warning about a standard deviation being zero. The correlation coefficient value that results is NA, i.e., missing. While we're summarizing the correlation between rate

and expression, let's also show the number of distinct genes within each grouping.

```
ydat %>%
  group_by(nutrient, bp) %>%
  summarize(r=cor(rate, expression), ngenes=n_distinct(symbol))
## Warning in cor(c(0.05, 0.05, 0.05, 0.05, 0.05, 0.05, 0.05, 0.05, 0.05,
## 0.05, : the standard deviation is zero
## Source: local data frame [5,286 x 4]
## Groups: nutrient [?]
##
##
      nutrient
                                                      bp
                                                               r ngenes
##
         <chr>>
                                                   <chr>
                                                           <dbl>
                                                                  <int>
## 1
       Ammonia
                            'de novo' IMP biosynthesis*
                                                         0.3125
                                                                      8
## 2
       Ammonia
                 'de novo' pyrimidine base biosynthesis -0.0482
                                                                       3
               'de novo' pyrimidine base biosynthesis*
## 3
       Ammonia
                                                         0.1670
                                                                       4
       Ammonia
## 4
                     35S primary transcript processing
                                                                     13
                                                         0.5080
## 5
       Ammonia
                    35S primary transcript processing*
                                                          0.4240
                                                                     30
## 6
       Ammonia
                                   acetate biosynthesis 0.4677
                                                                       1
## 7
       Ammonia
                                     acetate metabolism 0.9291
                                                                       1
## 8
       Ammonia
                                    acetate metabolism* -0.6855
                                                                      1
## 9
       Ammonia
                                acetyl-CoA biosynthesis -0.8512
                                                                       1
## 10
                 acetyl-CoA biosynthesis from pyruvate 0.0951
       Ammonia
                                                                       1
## # ... with 5,276 more rows
```

7. Take the above code and continue to process the result to show only results where the process has at least 5 genes. Add a column corresponding to the absolute value of the correlation coefficient, and show for each nutrient the singular process with the highest correlation between rate and expression, regardless of direction. *Hint:* 4 more pipes: filter, mutate, arrange, and filter again with row_number()==1. Ignore the warning.

```
## Source: local data frame [6 x 5]
## Groups: nutrient [6]
##
##
      nutrient
                                                          bp
                                                                 r ngenes
                                                                           absr
##
         <chr>>
                                                       <chr> <dbl>
                                                                    <int>
                                                                          <dbl>
## 1
       Glucose telomerase-independent telomere maintenance -0.95
                                                                           0.95
## 2
       Ammonia telomerase-independent telomere maintenance -0.91
                                                                        7
                                                                           0.91
##
  3
       Leucine telomerase-independent telomere maintenance -0.90
                                                                        7
                                                                           0.90
## 4 Phosphate telomerase-independent telomere maintenance -0.90
                                                                           0.90
        Uracil telomerase-independent telomere maintenance -0.81
                                                                        7
                                                                           0.81
## 5
## 6
       Sulfate
                                  translational elongation* 0.79
                                                                           0.79
```