Test for travel quarantine effect on SARS-CoV-2 imports

This script is to test whether the rate of new introductions from a country is effected by whether travellers from the country are under quarantine orders.

Date range for analysis is constrained to quarantine list data start/end dates. Phylogenetic import estimates are only for chains where travel context sequences clustered with them, and also only for one chain per polytomy. So these represent estimates for only a fraction of the total number of chains.

Q: why does it matters which transmission chain assumption we make if we take each polytomy as 1 import anyways? A: when polytomies are allowed to be Swiss, if the polytomy node is swiss then indeed, we cannot classify a location for the origin of the transmission chain This means should have more datapoints when s=F.

TODO: compare/contrast results when s=T and s=F

```
workdir <- "../grapevine/jan-dec_-01_max_sampling_-5_context-sf"</pre>
outdir <- paste(workdir, "output", sep = "/")</pre>
system(command = paste("mkdir -p", outdir))
db_connection <- open_database_connection(config_file = "../../../Repos/grapevine/workdir/input/config.)
## [1] "config_file location: ../../../Repos/grapevine/workdir/input/config.yml"
## [1] "config file exists TRUE"
Load data: independent variables
# indep variables: source country, whether travellers under quarantine order by day
travel_quarantine_by_country_day <- get_travel_quarantine_by_country_day(</pre>
 db_connection = db_connection
# indep variable: source country incidence per capita per day
country_incidence <- get_country_incidence(</pre>
 min_date = min(travel_quarantine_by_country_day$date),
 max_date = max(travel_quarantine_by_country_day$date),
 db_connection = db_connection
# indep variable: distance from country to Switzerland
country_distances <- get_country_distances(</pre>
 non_focal_countries = as.character(unique(travel_quarantine_by_country_day$iso_code)),
 focal_countries = "CHE".
 db_connection = db_connection
# indep variable: travel connection to Switzerland
country_arrivals <- read.delim(</pre>
 file = paste(
   workdir,
    "tmp/alignments/estimated_monthly_infectious_arrivals.txt",
    sep = "/"),
 stringsAsFactors = F) %>%
  select(date, iso_code, n_tourist_arrivals, n_commuter_permits)
```

```
# clean data
warn_missing_incidence_data(
 travel_quarantine_by_country_day = travel_quarantine_by_country_day,
 country_incidence = country_incidence
## Warning in warn_missing_incidence_data(travel_quarantine_by_country_day = travel_quarantine_by_count:
## iso_code
                        country_name
## 1
          ABW
                               Aruba
## 2
          GUM
                                Guam
## 3
                    French Polynesia
         PYF
## 4
          SXM
                        Sint Maarten
## 5
          VGB British Virgin Islands
travel_quarantine_by_country_day <- travel_quarantine_by_country_day %>%
 filter(iso_code %in% country_incidence$iso_code)
# merge data
indep_vars <- merge(</pre>
   x = travel_quarantine_by_country_day, y = country_incidence,
    by = c("iso_code", "date"),
   all = T)
indep_vars <- merge(</pre>
 x = indep_vars, y = country_distances,
 by = "iso_code",
 all.x = T)
indep_vars <- merge(</pre>
 x = indep_vars, y = country_arrivals,
by = c("iso_code", "date"),
all.x = T)
Load data: possible dependent variables
# dep variable: number of FOPH recorded exposures from each country each day
recorded_exposures <- get_exposures_per_country_day(</pre>
  min_date = min(travel_quarantine_by_country_day$date),
  max_date = max(travel_quarantine_by_country_day$date),
 db_connection = db_connection
)
## [1] "Getting number of cases by exposure country and date of case confirmation from BAG meldeformula:
## Warning in get_exposures_per_country_day(min_date = min(travel_quarantine_by_country_day$date), : Ren
## # A tibble: 2 x 4
## # Groups: date [2]
## english_name exp_land n_exposures date
                                <int> <date>
## <chr>
                 <chr>
## 1 <NA>
                  ANDERES
                                    1 2020-11-26
                  Namibia
## 2 <NA>
                                     1 2021-02-07
# dep variable: number of lineages departing a country each day
chains_asr <- load_chains_asr(s = s, workdir = workdir)</pre>
## Warning in scan(file = file, what = what, sep = sep, quote = quote, dec = dec, :
## EOF within quoted string
## Warning in scan(file = file, what = what, sep = sep, quote = quote, dec = dec, :
```

```
## EOF within quoted string
## Warning in read.table(file = file, header = header, sep = sep,
## quote = quote, : incomplete final line found by readTableHeader
## on '../grapevine/jan-dec_-01_max_sampling_-5_context-sf/tmp/asr/B.
## 1.1.74_m_3_p_1_s_F_tree_data_with_asr.txt'
## Warning in read.table(file = file, header = header, sep = sep,
## quote = quote, : incomplete final line found by readTableHeader
## on '../grapevine/jan-dec_-01_max_sampling_-5_context-sf/tmp/asr/B.
## 1.1.91_m_3_p_1_s_F_tree_data_with_asr.txt
chains_asr_representative <- pick_origin_representative_chains(chains_asr)</pre>
chains_asr_long <- pivot_chains_longer(chains_asr_representative)</pre>
asr_daily <- chains_asr_long %>%
 filter(foreign_tmrca >= min(travel_quarantine_by_country_day$date),
         foreign_tmrca <= max(travel_quarantine_by_country_day$date)) %>%
 group_by(foreign_tmrca, origin) %>%
 summarize(n_lineages = sum(asr_contribution, na.rm = T)) %>%
 ungroup() %>%
 mutate(iso_code = country_name_to_iso_code(country = origin)) %>%
 select(-origin) %>%
 rename("date" = "foreign_tmrca")
# merge data
dep_vars <- merge(</pre>
 x = recorded_exposures, y = asr_daily,
 by = c("iso_code", "date"),
all = T)
```

Merge independent and dependent variables, aggregate timechunks

```
model_data_daily <- merge(</pre>
  x = indep_vars, y = dep_vars,
 by = c("iso_code", "date"),
 all = T) \%
  tidyr::replace_na(list(
    "quarantine_order" = F, # not on the quarantine list = no quarantine order
"n_exposures" = 0, # no exposures in bag_meldeformular = 0 exposures
    "n_lineages" = 0, # no foreign attachment points in the month have travel context descendents from
   "n_tourist_arrivals" = 0,
   "n_commuter_permits" = 0)) %>% # no tourist arrivals to hotels recorded or commuter permits = 0 ar
  group_by(iso_code) %>%
 mutate(quarantine_y_and_n = length(unique(quarantine_order)) == 2) %>%
 filter(quarantine_y_and_n) %>% # only countries on & off quarantine list at some point
 select(-quarantine_y_and_n) %>%
 ungroup()
model_data_weekly <- model_data_daily %>%
 mutate(date = format(date, "%y-%U")) %>%
  group_by(date, iso_code) %>%
 summarize(
   n_exposures = sum(n_exposures),
   n_lineages = sum(n_lineages),
   dist_to_CHE = dist_to_CHE[1],
  new_cases_per_million = mean(new_cases_per_million),
```

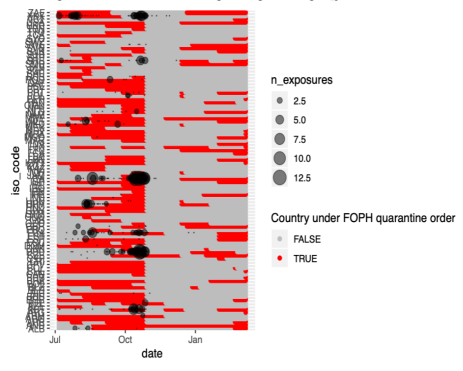
```
quarantine_order = sum(quarantine_order) / n()) # indep var frac of days on list

model_data_monthly <- model_data_daily %>%
    mutate(date = format(date, "%y-%m-01")) %>%
    group_by(date, iso_code) %>%
    summarize(
    n_exposures = sum(n_exposures),
    n_lineages = sum(n_lineages),
    dist_to_CHE = dist_to_CHE[1],
    new_cases_per_million = mean(new_cases_per_million),
    quarantine_order = sum(quarantine_order) / n(), # indep var frac of days on list
    n_tourist_arrivals = sum(n_tourist_arrivals, na.rm = T), # a monthly variable only
    n_commuter_permits = sum(n_commuter_permits, na.rm = T)) # a monthly variable only
```

Plot data summary

```
plot_dep_var_by_quarantine_status(
  model_data = model_data_daily,
  outdir = outdir,
  dep_varname = "n_exposures")
```

- ## Saving 6.5×10 in image
- ## Warning: Removed 24292 rows containing missing values (geom_point).
- ## Warning: Removed 24292 rows containing missing values (geom_point).



```
plot_dep_var_by_quarantine_status(
 model_data = model_data_daily,
 outdir = outdir,
 dep_varname = "n_lineages")
## Saving 6.5 \times 10 in image
## Warning: Removed 24372 rows containing missing values (geom_point).
## Warning: Removed 24372 rows containing missing values (geom_point).
                                                   n_lineages
                                                    1
iso_code
                                                   Country under FOPH quarantine order
                                                    FALSE
                                                    TRUE
                     Oct
                                   Jan
       Jul
                         date
```

Define model

```
fit_model <- function(
  model_data, return_fitted_model = F, dep_varname, indep_varnames)
) {
  my_formula <- paste(dep_varname, " ~ ", pasteO(indep_varnames, collapse = " + "), sep = "")
  print(paste("Fitted model:", my_formula))
  fitted_model <- lm(
    data = model_data,
    formula = as.formula(my_formula))
  print(summary(fitted_model))
  if (return_fitted_model) {
    return(fitted_model)
  }
}</pre>
```

Fit model(s) Assumption: travel is constant year-round

```
fit_model(
 model_data = model_data_monthly,
  dep_varname = "n_exposures",
 indep_varnames = c("new_cases_per_million", "quarantine_order", "dist_to_CHE",
                    "n_tourist_arrivals", "n_commuter_permits"))
## [1] "Fitted model: n_exposures ~ new_cases_per_million + quarantine_order + dist_to_CHE + n_tourist_;
##
## Call:
## lm(formula = as.formula(my_formula), data = model_data)
## Residuals:
## Min
               10 Median
                              30
## -20.139 -0.590 -0.232 0.143 98.602
##
## Coefficients:
                         Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                        7.992e-01 3.358e-01 2.380 0.0175 *
## new_cases_per_million -1.754e-03 9.124e-04 -1.923 0.0548 .
## quarantine_order
                        8.914e-01 3.800e-01 2.346
                                                       0.0192 *
## dist_to_CHE
                        -1.160e-04 4.802e-05 -2.416
                                                      0.0159 *
                         9.958e-05 2.213e-05 4.500 7.72e-06 ***
## n_tourist_arrivals
                         8.294e-05 1.170e-05 7.091 2.72e-12 ***
## n_commuter_permits
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 4.564 on 885 degrees of freedom
## (9 observations deleted due to missingness)
## Multiple R-squared: 0.1672, Adjusted R-squared: 0.1625
## F-statistic: 35.54 on 5 and 885 DF, p-value: < 2.2e-16
fit_model(
 model_data = model_data_monthly,
  dep_varname = "n_lineages",
 indep_varnames = c("new_cases_per_million", "quarantine_order", "dist_to_CHE",
                    "n_tourist_arrivals", "n_commuter_permits"))
## [1] "Fitted model: n_lineages ~ new_cases_per_million + quarantine_order + dist_to_CHE + n_tourist_a:
## Call:
## lm(formula = as.formula(my_formula), data = model_data)
##
## Residuals:
               1Q Median
     Min
## -9.7390 -0.0805 -0.0222 0.0274 17.2399
##
## Coefficients:
                         Estimate Std. Error t value Pr(>|t|)
##
                         1.298e-01 6.729e-02 1.929 0.0540 .
## new_cases_per_million -4.375e-04 1.828e-04 -2.393
                                                      0.0169 *
                        1.000e-01 7.614e-02
## quarantine_order
                                              1.314
                        -1.536e-05 9.621e-06 -1.596
## dist_to_CHE
                                                      0.1108
## n_tourist_arrivals
                         3.828e-05 4.434e-06 8.632 <2e-16 ***
## n_commuter_permits
                         5.304e-05 2.343e-06 22.636 <2e-16 ***
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.9145 on 885 degrees of freedom
    (9 observations deleted due to missingness)
## Multiple R-squared: 0.5781, Adjusted R-squared: 0.5757
## F-statistic: 242.5 on 5 and 885 DF, p-value: < 2.2e-16
model_data_monthly_gathered <- model_data_monthly %>%
 tidyr::gather(
   key = "variable", value = "value",
   dist_to_CHE, new_cases_per_million, quarantine_order, n_tourist_arrivals,
   n_commuter_permits)
ggplot(data = model_data_monthly_gathered,
      aes(x = value, y = n_lineages)) +
  geom_point() +
 facet_wrap(~variable, scales = "free")
```

Warning: Removed 9 rows containing missing values (geom_point).

