## **Explore data**

This analysis is very similar to what I did last May for the #TidyTuesday data set on cocktail recipes, so take a look at both to see what is the same and what is different for the two different data sets. Our modeling goal is to use unsupervised algorithms for dimensionality reduction with United Nations voting data to understand which countries are similar.

```
library(tidyverse)
unvotes <- read_csv("https://raw.githubusercontent.com/rfordatascience/tidytuesday/
master/data/2021/2021-03-23/unvotes.csv")
issues <- read_csv("https://raw.githubusercontent.com/rfordatascience/tidytuesday/
master/data/2021/2021-03-23/issues.csv")</pre>
```

Let's create a wide version of this data set via pivot wider () to use for modeling.

```
unvotes_df <- unvotes %>%
  select(country, rcid, vote) %>%
  mutate(
    vote = factor(vote, levels = c("no", "abstain", "yes")),
    vote = as.numeric(vote),
    rcid = paste0("rcid_", rcid)
    ) %>%
    pivot_wider(names_from = "rcid", values_from = "vote", values_fill = 2)
```

## Principal component analysis

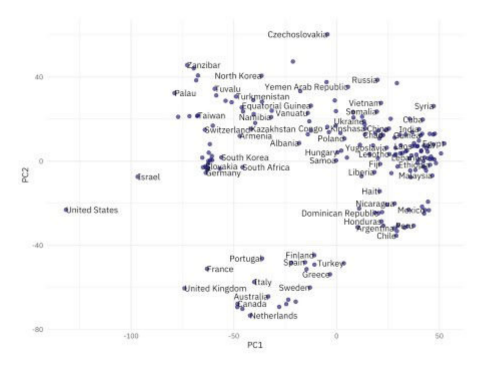
This analysis only uses the recipes package, the tidymodels package for data preprocessing and feature engineering that contains functions for unsupervised methods. There are lots of options available, like step\_ica() and step\_kpca(), but let's implement a basic principal component analysis.

```
library(recipes)
pca_rec <- recipe(~., data = unvotes_df) %>%
 update role(country, new role = "id") %>%
 step normalize(all predictors()) %>%
 step pca(all predictors(), num comp = 5)
pca_prep <- prep(pca_rec)</pre>
pca prep
## Data Recipe
##
## Inputs:
##
        role #variables
##
         id
                  6202
## predictor
```

```
##
## Training data contained 200 data points and no missing data.
##
## Operations:
##
## Centering and scaling for rcid_3, rcid_4, rcid_5, rcid_6, rcid_7,
... [trained]
## PCA extraction with rcid_3, rcid_4, rcid_5, rcid_6, rcid_7, ...
[trained]
```

We can look at where the countries are in the principal component space by **baking** the prepped recipe.

```
bake(pca_prep, new_data = NULL) %>%
  ggplot(aes(PC1, PC2, label = country)) +
  geom_point(color = "midnightblue", alpha = 0.7, size = 2) +
  geom_text(check_overlap = TRUE, hjust = "inward", family =
"IBMPlexSans") +
  labs(color = NULL)
```



We can look at which votes contribute to the component by **tidying** the prepped recipe. Let's join the roll call votes up with the topics to see which topics contribute to the top principal components.

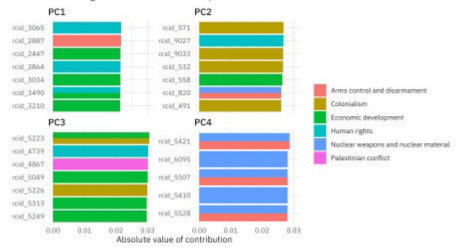
```
pca_comps <- tidy(pca_prep, 2) %>%
  filter(component %in% paste0("PC", 1:4)) %>%
  left_join(issues %>% mutate(terms = paste0("rcid_", rcid))) %>%
  filter(!is.na(issue)) %>%
  group_by(component) %>%
  top_n(8, abs(value)) %>%
  ungroup()

pca_comps %>%
  mutate(value = abs(value)) %>%
  ggplot(aes(value, fct_reorder(terms, value), fill = issue)) +
```

```
geom_col(position = "dodge") +
facet_wrap(~component, scales = "free_y") +
labs(
    x = "Absolute value of contribution",
    y = NULL, fill = NULL,
    title = "What issues are most important in UN voting country
differences?",
    subtitle = "Human rights and economic development votes account for
the most variation"
)
```

## What issues are most important in UN voting country differences?

Human rights and economic development votes account for the most variation.



The PCA implementation did not know about the topics of the votes, but notice how the first principal component is mostly about human rights and economic development, the second principal component is mostly about colonialsim, and so on.

## **UMAP**

To switch out for a different dimensionality reduction approach, we just need to change to a different recipe <code>step\_()</code>. Let's try out UMAP, a different algorithm for dimensionality reduction based on ideas from topological data analysis, which is available in the <code>embed</code> package.

```
library(embed)

umap_rec <- recipe(~., data = unvotes_df) %>%
    update_role(country, new_role = "id") %>%
    step_normalize(all_predictors()) %>%
    step_umap(all_predictors())

umap_prep <- prep(umap_rec)

umap_prep

## Data Recipe
##
## Inputs:
##
## role #variables</pre>
```

```
## id 1
## predictor 6202
##
## Training data contained 200 data points and no missing data.
##
## Operations:
##
## Centering and scaling for rcid_3, rcid_4, rcid_5, rcid_6, rcid_7,
... [trained]
## UMAP embedding for rcid_3, rcid_4, rcid_5, rcid_6, rcid_7, ...
[trained]
```

When we visualize where countries are in the space created by this dimensionality reduction approach, it looks very different!

```
bake(umap_prep, new_data = NULL) %>%
  ggplot(aes(umap_1, umap_2, label = country)) +
  geom_point(color = "midnightblue", alpha = 0.7, size = 2) +
  geom_text(check_overlap = TRUE, hjust = "inward", family =
"IBMPlexSans") +
  labs(color = NULL)
```

