05 - Anomaly Detection

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05-anomaly.pdf

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1 Anomaly Detection Intro

1.1 Required R Packages

We will be using the R packages of:

- tidyverse for data manipulation and visualization
- mclust for model-based clustering

```
library(mclust) # install.packages("mclust")
library(tidyverse)
library(readxl)
```

1.2 Anomaly Detection

Anomaly Detection: The identification of unusual observations. Statistically, this means finding observations that come from a different distribution that the *normal* or *usual* observations.

1. Goodness of Fit (GOF)

- Tests if data conform to a given distribution (or distributional family)
- Use case: Failure of the first digits in a financial statement to conform to Benford's distribution may indicate fraud.

2. Two-Sample Tests (A/B Testing)

- Tests if two datasets come from the same distribution
- Often simplified to test if one group has a larger mean than the other
- Use case: Determine if a new surgical technique leads to faster recovery times.

3. Outlier Detection

- Tests if a single observation or small set of observations come from the same distribution as the rest of the data
- Use case: Detect and correct data entry errors.

4. Hotspot Detection

- Identification of regions that have unusually high density
- Use case: add additional police patrols to regions and times that are experiencing an unusually high crime rate

5. Outbreak Detection

- A sequential method that repeatedly tests for a change in an event (i.e., point process) distribution
- Focus on determining if a change occurred, and then determining when it occurred
- For outbreak detection, the changes of interest are those that conform to an expected *outbreak pattern*
- Use case: quickly detect the presence of West Nile Virus from the *chief complaints* field of health records and initiate a rapid mosquito control spraying

1.3 Example #1: Benford's Distribution

State/Territory	Real or Faked Area (km ²)		
Afghanistan	645,807	796,467	
Albania	28,748	9,943	
Algeria	2,381,741	3,168,262	
American Samoa	197	301	
Andorra	464	577	
Anguilla	96	82	
Antigua and Barbuda	442	949	
Argentina	2,777,409	4,021,545	
Armenia	29,743	54,159	
Aruba	193	367	
Australia	7,682,557	6,563,132	
Austria	83,858	64,154	
Azerbaijan	86,530	71,661	
Bahamas	13,962	9,125	
Bahrain	694	755	
Bangladesh	142,615	347,722	
Barbados	431	818	
Belgium	30,518	47,123	
Belize	22,965	20,648	
Benin	112,620	97,768	

Table from Fewster (2009) A Simple Explanation of Benford's Law, *The American Statistician*, 63, 1, pp 26–32

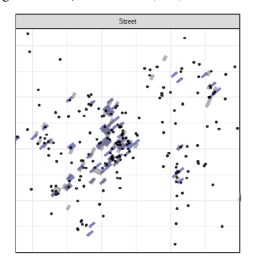
- Someone that fakes numbers, say on a financial statement, may be tempted to use a random number generator
 - But watch out for Benford's Law
- Note on terminology:
 - Law = probability distribution
 - Anomalous Numbers = Random numbers (no known relationship)
- Benford's PMF:

$$\Pr(\text{first digit} = x) = \log_{10} \left(1 + \frac{1}{x} \right) \quad \text{for } x = 1, 2, \dots, 9$$

1.4 Example #2: Crime Hotspots

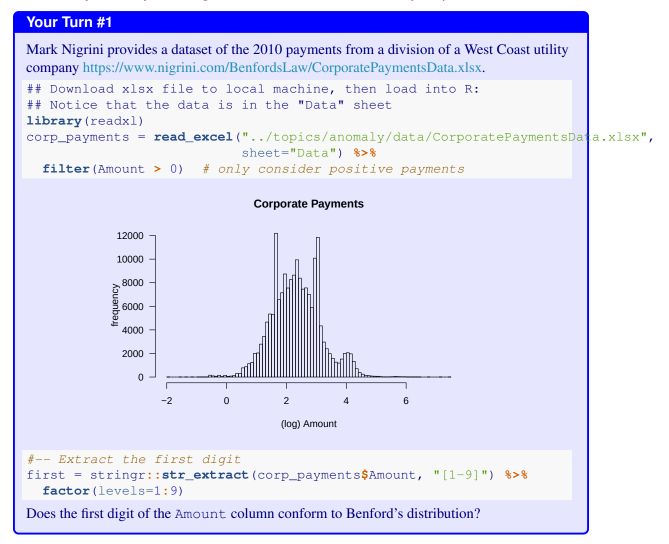
In 2017, the National Institute of Justice (NIJ) held a Crime Forecasting Challenge

• Predict the crime hotspots for 4 crime types (burglary, street crime, motor vehicle theft, all types) for 5 forecasting windows (1 week ahead, ..., 3 months ahead)



2 Goodness of Fit Testing

Tests if data conform to a given distribution (or distributional family)



2.1 GOF Hypothesis Test

- Let $D = (X_1, X_2, \dots, X_n)$ be the observed random variables (i.e., the data).
- Let \mathcal{H}_0 be the *null* hypothesis
- Choose a test statistic $T = T(X_1, \dots, X_n)$ that is a function of the observed data
 - T is a random variable; it has a distribution.
 - Let $t = T(x_1, \dots, x_n)$ be the *observed* value of the test statistic
 - Common to adjust the test statistic so that *extreme* means large values of T
- The p-value is the probability that chance alone would produce a test statistic as extreme as the observed test statistic if the null hypothesis is true
 - E.g., p-value = $Pr(T \ge t | \mathcal{H}_0)$
- Think of T or the p-value as the evidence against the null hypothesis
 - Its common to set a threshold (e.g., p-value ≤ .05) and reject the null hypothesis when this threshold is crossed.
 - This is a form of *outlier detection*. Reject null if $t_{\rm obs}$ is an *outlier*; that is $t_{\rm obs}$ is from a different distribution that what is specified in \mathcal{H}_0 .

- To calculate a p-value, we need to know/estimate the distribution of $T|\mathcal{H}_0!$
 - Even if we don't know the distribution of T under the null, we can often approximate it using simulation (Monte Carlo)

2.1.1 Example: one sample t-test

- In 2012, the Obama administration issued new rules on the fuel efficiency requirements for new cars and trucks by 2025.
 - The fleetwise fuel efficiency requirement is 54.5 mpg
- Suppose a car maker in 2025 designed a car to get an average fuel efficiency of 54.5 mpg.
 - Also, they think the fuel efficiency will be Normally distributed (Gaussian)
- The government officials randomly tested n=9 cars. They got a sample mean of $\bar{x}=53.0$ and sample standard deviation of s=2.5.

Your Turn #2

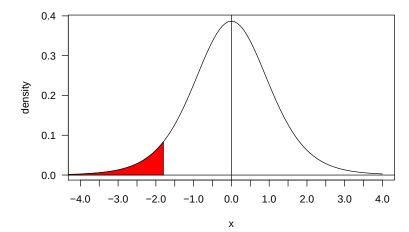
Is this enough evidence to conclude the car manufacturer failed to meet the requirements? Or can the results be attributed to chance fluctuation?

- Null Hypothesis:
 - the mpg come from a Normal distribution
 - mean of $\mu_0 = 54.5$
 - independent
 - standard deviation, σ is unknown
 - $X|\mathcal{H}_0 \stackrel{\text{iid}}{\sim} \mathcal{N}(\mu = 54.5, \sigma)$
- Choose the test statistic:

Test Statistic

- Under the null, T has a t-distribution with df = n 1
 - Shoutout to William Sealy Gosset, a.k.a Student

Distribution of sample statistic: T ~ t(df=8)



• $Pr(T \le -1.8 | \mathcal{H}_0) = pt((53.0-54.5)/(2.5/3), df=8) = 0.055$

2.1.2 Alternative Hypotheses

- The choice of test statistic depends on the expected deviations from the null
 - That is, we can come up with *better* test statistic if we know what sort of deviations from \mathcal{H}_0 are expected.
 - better meaning more power to correctly reject the null

2.2 Test Statistics

• Going back to the original question about the corporate payments conforming to Benford's distribution, we can state the *null hypothesis* formally:

$$\mathcal{H}_0: X \stackrel{\text{iid}}{\sim} Benf$$

- *X* is the *first* digit(s)
- Benf stands for Benford's distribution for the first digit(s). This has pmf:

$$f(x) = \log_{10} \left(1 + \frac{1}{x} \right)$$

- Note: there are no parameters to estimate!
- A generic alternative hypothesis is:

$$\mathcal{H}_1: X \not\sim Benf$$

• R code for a Benford pmf

```
#-- pmf for Benford's distribution
dbenford <- function(x) log10(1 + 1/x)

#-- first digit
dbenford(1:9)
#> [1] 0.30103 0.17609 0.12494 0.09691 0.07918 0.06695 0.05799 0.05115 0.04576

#-- first two digits
expand.grid(first=1:9, second=0:9) %>%
    mutate(two = paste0(first, second) %>% as.integer) %>%
    mutate(f = dbenford(two)) %>%
    select(first, second, f) %>%
    spread(second, f) %>%
    knitr::kable(digits=3)
```

first	0	1	2	3	4	5	6	7	8	9
1	0.041	0.038	0.035	0.032	0.030	0.028	0.026	0.025	0.023	0.022
2	0.021	0.020	0.019	0.018	0.018	0.017	0.016	0.016	0.015	0.015
3	0.014	0.014	0.013	0.013	0.013	0.012	0.012	0.012	0.011	0.011
4	0.011	0.010	0.010	0.010	0.010	0.010	0.009	0.009	0.009	0.009
5	0.009	0.008	0.008	0.008	0.008	0.008	0.008	0.008	0.007	0.007
6	0.007	0.007	0.007	0.007	0.007	0.007	0.007	0.006	0.006	0.006
7	0.006	0.006	0.006	0.006	0.006	0.006	0.006	0.006	0.006	0.005
8	0.005	0.005	0.005	0.005	0.005	0.005	0.005	0.005	0.005	0.005
9	0.005	0.005	0.005	0.005	0.005	0.005	0.005	0.004	0.004	0.004

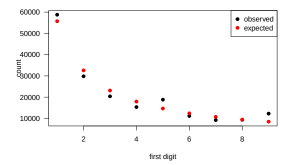
2.2.1 χ^2 Test Statistic

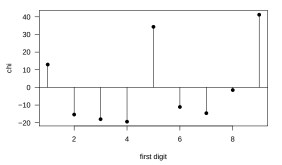
- The Pearson's χ^2 (chi-squared) test statistic is commonly used in goodness-of-fit testing.
- It requires the data to be discrete or categorical
 - Continuous data can be binned
- Test Statistic:

$$\chi^2 = \sum_{j=1}^{J} \frac{(y_j - E_j)^2}{E_j} = \sum_{j=1}^{J} \frac{(y_j - np_j)^2}{np_j}$$

- *J*: number of categories or possible values
- y_j : observed count in category j
- E_j : expected count (i.e., under \mathcal{H}_0) in category j
- Let $n = \sum_{j=1}^{J} y_j$ be the total number of observations
- p_j : the proportion of events under the null (i.e., $\Pr(X=j|\mathcal{H}_0)$)
- Asymptotically, χ^2 converges to a chi-squared distribution with J-1 degrees of freedom
- R code for corporate payments data

```
#-- Get counts
Y = table(first) %>% as.integer # ensure first is factor with properly ordered
n = length(first) # number of observations
#-- chi-squared
n = length(first) # number of observations
E = n*dbenford(1:9) # expected count vector
chi = (Y-E)/sqrt(E) # vector of deviations
(chisq = sum(chi^2)) # chi-squared test statistic
#> [1] 4317
```





• Note: there is a build-in R function chisq.test() which does these calculations

```
chisq.test(Y, p=dbenford(1:9))$statistic
#> X-squared
#> 4317
```

2.2.2 Likelihood Ratio Test Statistic

• When an *alternative hypothesis* can be specified, the log-likelihood ratio test statistic is commonly used in goodness-of-fit testing.

- The general binary hypothesis formulation is:
 - $\mathcal{H}_0: X \sim f_0(X)$ (null hypothesis)
 - $\mathcal{H}_1: X \sim f_1(X)$ (alternative hypothesis)
- The likelihood ratio is:

$$LR = \frac{f_1(X_1, \dots, X_n)}{f_0(X_1, \dots, X_n)}$$
$$= \prod_{i=1}^n \frac{f_1(X_i)}{f_0(X_i)} \quad \text{if } X\text{'s are iid}$$

• The log-likelihood ratio, when the observations are iid, becomes:

$$llr = \sum_{i=1}^{n} \log \frac{f_1(X_i)}{f_0(X_i)}$$
$$= \sum_{i=1}^{n} \log f_1(X_i) - \sum_{i=1}^{n} \log f_0(X_i)$$

- The hypotheses for the corporate payments data:
 - $\mathcal{H}_0: X \stackrel{\text{iid}}{\sim} Benf$
 - $\mathcal{H}_1: X \stackrel{\text{iid}}{\sim} Cat(p_1, p_2, ..., p_9)$ where $\{p_k\}$ do *not* match Benford's probabilities.
- There are many reasonable choices for setting \mathcal{H}_1 parameters (p_1,\ldots,p_9)
 - Discrete Uniform: $p_1 = ... = p_9 = 1/9$
 - MLE: $\hat{p}_k = y_i/n$

Your Turn #3

Write out the log-likelihood for the two alternatives.

Using MLE, the llr = 2031.83.

• Note: $2 \times llr$ has an asymptotic chi-squared distribution (same as the chi-squared test statistic).

2.3 Testing

- The two-test statistics, χ^2 and llr, provide evidence against \mathcal{H}_0 .
- But how do we know if these values are *unusually* large? Perhaps by chance alone (i.e. the data sample we observed) the values are as large as they are.
- We can answer this with a solid probabilistic statement if we knew the *distribution of the test* statistic under the null hypothesis

- We don't often know this exactly, but there are often good approximations that hold as the sample size grows (asymptotically).
- There are two primary options:
 - 1. Use an asymptotic distribution (e.g., the chi-squared distribution)
 - 2. Use Monte Carlo simulation

2.3.1 Monte Carlo Simulation

- If we can sample from the null hypothesis, then it becomes straightforward to estimate the distribution of *any* test statistic and consequently, *p*-values.
- Monte Carlo based GOF Test
 - 1. Calculate the test statistic for the original observation, t
 - 2. Generate M samples from the null distribution

```
- \{Y_1, \ldots, Y_M\}
```

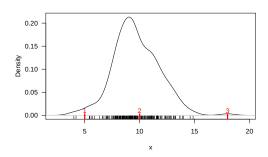
3. For each sample, calculate the test statistic T^*

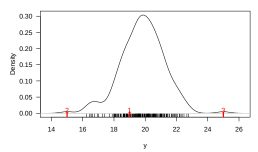
```
-\{T_1^*,\ldots,T_M^*\}
```

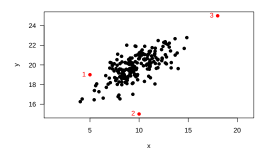
4. p-value = $\frac{1 + \text{number of } T^*\text{'s greater than or equal to } t}{M + 1}$

3 Outlier Detection

- Outlier Detection tests if a single observation or small set of observations come from the same distribution as the rest of the data
- Are any of the *red* points outliers?







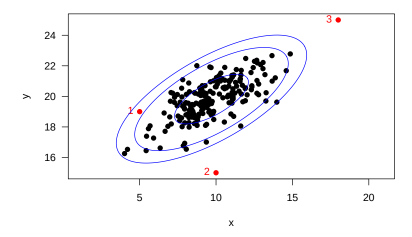
3.1 Distance based approach

- One approach, with strong connections to clustering, is to calculate the *distance* from an observation to the centroid
 - This assumes the "normal" observations are from a unimodal distribution
 - To allow for an ellipse shape (orientation) and different spreads in each dimension, use the squared *Mahalanobis Distance*

$$D_i^2 = (\mathbf{x_i} - \bar{\mathbf{x}})^\mathsf{T} \hat{\Sigma}^{-1} (\mathbf{x_i} - \bar{\mathbf{x}})$$

- Note that $\bar{\mathbf{x}}$ and $\hat{\Sigma}$ are estimated from all of the data.
- Estimated Parameters:
 - $-\bar{x} = (9.70, 19.82)$
 - $\hat{\Sigma} = (4.2171, 1.9482, 1.9482, 1.906)$

obs	X	у	Dsq
1	5	19	7.056
2	10	15	24.471
3	18	25	18.123



3.2 Likelihood Based Approach

- From the plots, it appears the a 2D Gaussian/Normal model could be a decent approximation to the distribution of the non-outlier observations
- We can use this to calculate the log-likelihood of observation i using the estimated parameters

Gaussian Log-Likelihood

$$\log L_i = \mathcal{N}(\mathbf{x}_i; \mu = \bar{x}, \Sigma = \hat{\Sigma})$$
=

- Notice that this is a function of the squured Mahalanobis distance!
- Robust estimation:
 - If indeed we have outliers, then these will be affecting out estimated parameters \bar{x} and $\hat{\Sigma}$.
 - Robust estimation techniques can help limit the damage caused by the outliers
 - Another, more structured approach, is mixture models!

3.3 Mixture Model Approach

- We can go back to our mixture model formulation and propose that the outliers come from a different distribution than the normal observations
- In mixture formulation

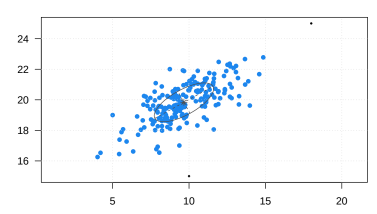
$$f(x) = \pi h(x) + (1 - \pi)g(x)$$

- h(x) is the pdf for the *outliers*
- -g(x) is the pdf for the normal observations
- π is the prior probability that an observation will be an outlier
- $E[\text{number of outliers}] = n\pi$

- There are several options for the outlier distribution h(x)
 - The uniform distribution on the minimum bounding box is one simple approach
 - * h(x) = 1/V, where V is the volume of the bounding box
 - * $V = \prod_{j=1}^{p} (max(x_j) min(x_j))$
- The Mclust() function in the R package mclust permits this formulation using the initialization=list(noise=TRUE) argument

```
#-- Fit mixture model with uniform noise
library (mclust)
mc = Mclust(X, initialization=list(noise=TRUE))
summary (mc)
#> Gaussian finite mixture model fitted by EM algorithm
#>
#> Mclust EVE (ellipsoidal, equal volume and orientation) model with 1 component
#> and a noise term:
#>
   log.likelihood
                   n df BIC
           -713.4 203 7 -1464 -1473
#>
#>
#> Clustering table:
   1 0
#>
#> 201 2
plot (mc, what="classification", asp=1, las=1)
grid()
```

Classification



4 Two-Sample Testing (A/B Testing)

4.1 Example (A/B Testing): Clinical Trials

A placebo-controlled randomized trial proposes to assess the effectiveness (i.e., cure rate) of Drug A in curing infants suffering from sepsis. A clinical trial of n=600 infants using Drug A found that 40% were cured of sepsis while 36% of the n=1200 infants on a placebo were cured.

Your Turn #4

- 1. Is Drug A better than the placebo?
- 2. How much better?

Let $p_1 = \Pr(\text{ cure } | \text{ Drug A}) \text{ and } p_2 = \Pr(\text{ cure } | \text{ Placebo})$

- $\mathcal{H}_0: p_1 = p_2 \text{ or } p_1 p_2 = 0$
- $\mathcal{H}_a: p_1 > p_2 \text{ or } p_1 p_2 > 0$

4.1.1 Simulation Based Testing

The 1800 patients were randomly assigned to the treatment (Drug A) or placebo group. It turned out that:

- of the $n_1 = 600$ given Drug A, $n_1\bar{p}_1 = 600(0.4) = 240$ were cured
- of the $n_2 = 1200$ given Drug A, $n_1\bar{p}_2 = 1200(0.36) = 432$ were cured
- of the $n_1 + n_2 = 1800$ patients, a total of 672 (37.3%) were cured

Under the null hypothesis, \mathcal{H}_0 : $p_1 = p_2$, there is no real difference in the cure rate between treatment and placebo. The observed difference is due **only** to the random assignment.

4.1.2 Permutation Test

We can see what the outcomes would have been if we used a different assignment into treatment and placebo.

- Regroup all patients
- Draw n_1 samples, at random, and calculate \bar{p}_1^*
- Use the remaining n_2 to calculate \bar{p}_2^*
- Calculate the test statistic $Z^* = (\bar{p}_1^* \bar{p}_2^*)$

This is a possible outcome if the null hypothesis was actually true.

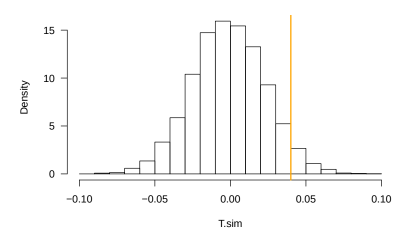
- If we repeat this procedure for all possible re-groupings, then we get the exact¹ distribution of the test statistic, if the null hypothesis was true.
- But, there are $\binom{1800}{600}$ (huge number) possible regroupings (permutations)
- Monte Carlo simulation can be used to approximate this distribution
- Just repeat the re-grouping procedure many times (say 1000 or 10000)
 - gives a set of observed values under the null model
- The estimated p-value is the proportion of simulated test statistic values that are more extreme than the **observed** test statistic

¹conditional on the data and study design

4.1.3 R Code

```
#- observed data
n1 = 600
p1 = 0.40
n2 = 1200
p2 = 0.36
p0 = (n1*p1 + n2*p2)/(n1+n2) # average cure rate
T.obs = p1 - p2
                     # Test Statistic: observed difference
#- Simulation Data
n = n1 + n2
                       # number of patients
x = n1*p1 + n2*p2
                       # total number cured
#- Run Simulation
set.seed(100)
                                         # set seed for replication
nsim = 10000
                                         # of simulations
x1.sim = rhyper(nsim, m=x, n=n-x, k=n1) # simulated # cured in pop 1
x2.sim = x - x1.sim
                                         # simulated # cured in pop 2
T.sim = x1.sim/n1 - x2.sim/n2
                                         # simulated test statistics
#- plots
hist(T.sim, breaks=seq(-.1,.1,by=.01), freq=FALSE, las=1) # histogram
abline(v=T.obs,col="orange",lwd=2) # add observed test statistic
```

Histogram of T.sim



```
#- p-value
(sum(T.sim >= T.obs) + 1) / (nsim +1)  # non-parametric p-value
#> [1] 0.05349
```

- 4.2 A/B Testing
- 4.3 Two-Sample Goodness of Fit
- 5 Hotspot Detection
- 5.1 Mixture Model Formulation
- 6 Outbreak Detection