Project II: Cuckoo Eggs Dimension

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Introduction

The Common Cuckoo (*Cuculus canorus*) is a bird species known for its distinctive behaviours and characteristics, particularly its unusual reproductive strategy. As brood parasites, cuckoos lay their eggs in the nests of other, often smaller, bird species. The host bird then incubates the cuckoo egg and unknowingly raises the chick as its own (Davies, 2011; Lahti, 2005). According to Kruger and Davies (2003), Davies (2011), and Lahti (2005), the survival of the cuckoo species hinges on their remarkable adaptation strategies, which include evading host defences through deception and matching their egg and chick development with the host's incubation and provisioning patterns.

Research conducted in Russia has provided evidence of evolutionary adaptation in cuckoos, particularly in their ability to vary egg size to match that of their hosts. This adaptation reduces the likelihood of egg rejection, as host birds are more inclined to reject eggs that are significantly different or larger than their own clutch (Meshcheryagina et al., 2018). The study found that while the breadth of cuckoo eggs closely matched that of the host species, the length did not, suggesting a trade-off that limits the cuckoo's ability to maintain larger egg volumes. This alignment in egg breadth is likely an evolved response to avoid host rejection (Meshcheryagina et al., 2018).

This project aims to investigate the egg dimensions—length and breadth—of six different cuckoo bird species, analysing 120 eggs. Various statistical analyses will be performed to determine differences in these dimensions both between and within the species.

Importing Data

For data manipulation, visualisation and analysis and to effectively read and access the relevant cuckoo data, the "tidyverse" was installed and activated. The read.csv () function was used to import and read the data that is in CSV format. The data was assigned a new name termed cuckoo data.

```
library(tidyverse)
Warning: package 'tidyverse' was built under R version 4.3.3
Warning: package 'ggplot2' was built under R version 4.3.3
Warning: package 'readr' was built under R version 4.3.3
-- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
v dplyr
           1.1.2
                    v readr
                               2.1.5
v forcats
           1.0.0
                    v stringr
                               1.5.0
          3.5.1
                    v tibble
                               3.2.1
v ggplot2
v lubridate 1.9.2
                    v tidyr
                               1.3.0
v purrr
           1.0.1
-- Conflicts ----- tidyverse_conflicts() --
x dplyr::filter() masks stats::filter()
x dplyr::lag()
                masks stats::lag()
```

cuckoo_data <- read.csv("https://raw.githubusercontent.com/xrander/bootcamp-test/master/data,</pre>

i Use the conflicted package (http://conflicted.r-lib.org/) to force all conflicts to become

To view and understand the contents of the data names() function was used to return the names of the columns in the cuckoo_data, this also ensures that the data has been imported correctly. The names of the the cuckoo_data are "length" "breadth" "species"

```
names(cuckoo_data)
```

```
[1] "length" "breadth" "species"
```

Table 1: Data Preview

$head(cuckoo_data, n = 10)$

```
length breadth
                       species
             16.1 Meadow Pipit
1
     21.7
2
     22.6
            17.0 Meadow Pipit
    20.9
3
           16.2 Meadow Pipit
4
    21.6
          16.2 Meadow Pipit
5
    22.2
          16.9 Meadow Pipit
6
    22.5
           16.9 Meadow Pipit
7
    22.2
           17.3 Meadow Pipit
    24.3
8
            16.8 Meadow Pipit
9
     22.3
             16.8 Meadow Pipit
     22.6
10
            17.0 Meadow Pipit
```

$tail(cuckoo_data, n = 10)$

	length	${\tt breadth}$	species
111	21.0	15.5	Wren
112	22.3	16.0	Wren
113	21.0	15.9	Wren
114	20.3	15.5	Wren
115	20.9	15.9	Wren
116	22.0	16.0	Wren
117	20.0	15.7	Wren
118	20.8	15.9	Wren
119	21.2	16.0	Wren
120	21.0	16.0	Wren

The names of the variables in the data is given above. The head() function was implemented to preview the first 10 rows of the data, while the tail() function was run to preview the last 10 rows of the dataset, thus making it easier to observe patterns in the data.

A preview of the data is presented in Table 1.

To further understand and observe the cuckoo_data the car::some() function was run, first setting a seed, to randomly select and generate a sample of rows from the cuckoo_data dataset. The set. seed() allows the same results every time the code is run.

```
set.seed(3891)
car::some(cuckoo_data)
```

	length	${\tt breadth}$	specie			
21	22.0	17.0	${\tt Meadow}$	Pipit		
31	22.5	17.1	${\tt Meadow}$	Pipit		
34	22.0	17.2	${\tt Meadow}$	Pipit		
58	23.2	16.4	Tree	Pipit		
80	23.0	16.1		${\tt Robin}$		
90	23.0	17.0		${\tt Robin}$		
99	24.9	16.8	Pied Wa	agtail		
109	20.9	15.7		Wren		
112	22.3	16.0		Wren		
117	20.0	15.7		Wren		

Exploratory Data Analysis

To investigate the main characteristics and understand the structure of the dataset, various analyses were conducted. The summary() function gave statistics of the three variables in the data frame; length, breadth, and species below:

A quick descriptive statistics of the data is given below:

summary(cuckoo_data)

len	gth	breadth		spec	cies
Min.	:19.60	Min.	:15.00	Length	n:120
1st Qu.	:21.90	1st Qu.	:16.20	Class	:character
Median	:22.35	Median	:16.60	Mode	:character
Mean	:22.45	Mean	:16.55		
3rd Qu.	:23.23	3rd Qu.	:17.00		
Max.	:25.00	Max.	:17.50		

The str() function was run to show the structure of the data along with the type and length of variables in the dataset. For cuckoo_data the str() function showed that the data has 120 observations, 3 variables, and the variables length and breadth are numerical, while species variable is character, as seen below:

```
str(cuckoo_data)
```

```
'data.frame': 120 obs. of 3 variables:
$ length: num 21.7 22.6 20.9 21.6 22.2 22.5 22.2 24.3 22.3 22.6 ...
$ breadth: num 16.1 17 16.2 16.2 16.9 16.9 17.3 16.8 16.8 17 ...
$ species: chr "Meadow Pipit" "Meadow Pipit" "Meadow Pipit" "Meadow Pipit" ...
```

To identify distinct species present in the data frame, the unique() function was used and it generated six species names in the data: Meadow Pipit, Tree Pipit, Hedge Sparrow, Robin, Pied Wagtail and Wren.

```
unique(cuckoo_data$species)
```

```
[1] "Meadow Pipit" "Tree Pipit" "Hedge Sparrow" "Robin" [5] "Pied Wagtail" "Wren"
```

The skim() function was run to obtain an extended structure of the data and gave statistical values of the data such as mean, median, and standard deviation along with missing values and histograms of the data.

```
skimr::skim(cuckoo data)
```

Table 2: Data summary

Name Number of rows	cuckoo_data 120
Number of columns	3
Column type frequency:	
character	1
numeric	2
Group variables	None

Variable type: character

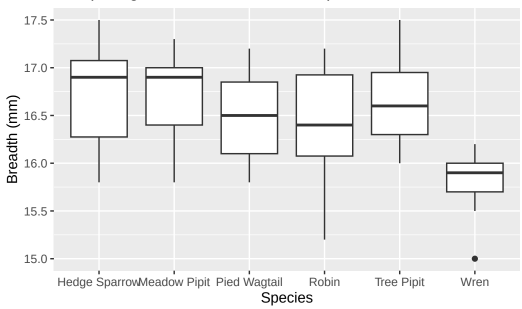
skim_variable	n_missing	$complete_rate$	min	max	empty	n_unique	whitespace
species	0	1	4	13	0	6	0

Variable type: numeric

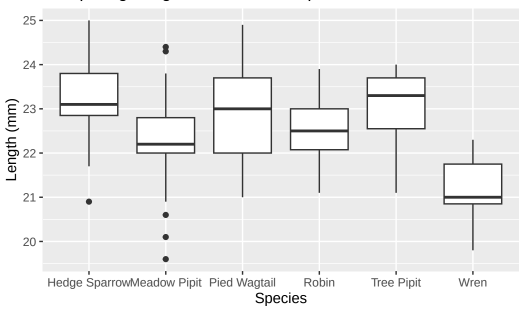
skim_variablen	_missing com	plete_ra	temean	sd	p0	p25	p50	p75	p100	hist
length	0	1	22.45	1.07	19.6	21.9	22.35	23.22	25.0	
breadth	0	1	16.55	0.52	15.0	16.2	16.60	17.00	17.5	

Visualisation

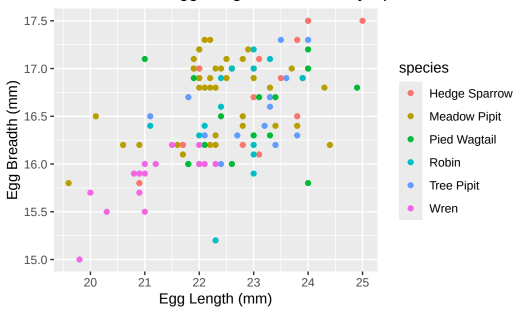
Comparing Breadth size between Species



Comparing Length size bewteen Species

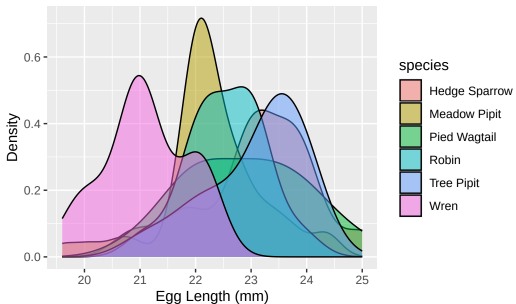


Scatter Plot of Egg Length vs Breadth by Species



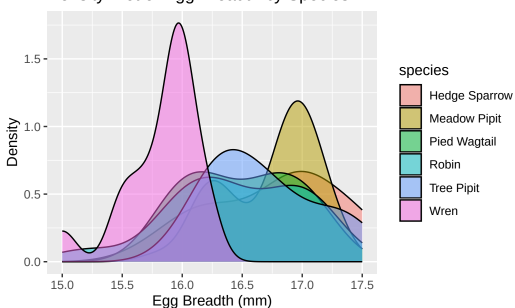
```
ggplot(cuckoo_data, aes(x = length, fill = species)) +
  geom_density(alpha = 0.5) +
  labs(title = "Density Plot of Egg Length by Species", x = "Egg Length (mm)", y = "Density"
```

Density Plot of Egg Length by Species



```
ggplot(cuckoo_data, aes(x = breadth, fill = species)) +
  geom_density(alpha = 0.5) +
  labs(title = "Density Plot of Egg Breadth by Species", x = "Egg Breadth (mm)", y = "Density")
```

Density Plot of Egg Breadth by Species



Inferential Statistical Analysis

Two-way ANOVA to test how the species (dependent variable) influences the length (independent variable) of eggs from different bird species (independent variable).

```
breadth_anova <- aov(breadth ~ species, data = cuckoo_data) |>
    anova()
breadth_anova
```

Analysis of Variance Table

```
Response: breadth

Df Sum Sq Mean Sq F value Pr(>F)

species 5 10.373 2.07457 10.885 1.412e-08 ***

Residuals 114 21.727 0.19059

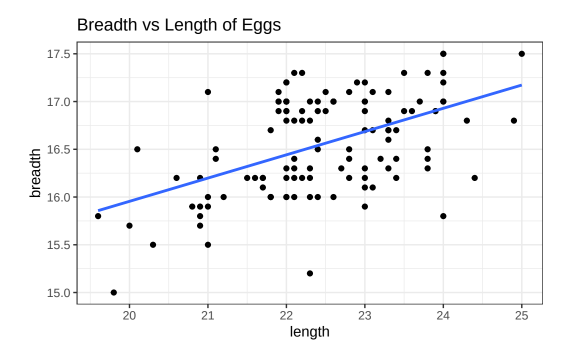
---

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Two-way ANOVA test to test how bird species independently influence breadth.

```
length_anova <- aov(length ~ species, data = cuckoo_data)|>
  anova()
length_anova
Analysis of Variance Table
Response: length
          Df Sum Sq Mean Sq F value Pr(>F)
          5 42.81 8.5620 10.449 2.852e-08 ***
Residuals 114 93.41 0.8194
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
library(ggplot2)
lm_model <- lm(breadth ~ length + species, data = cuckoo_data) |>
  summary()
ggplot(cuckoo_data, aes(x = length , y = breadth)) +
  geom_point() +
  geom_smooth(se = FALSE, method = "lm") +
  ggtitle("Breadth vs Length of Eggs") +
 theme_bw()
```

```
`geom_smooth()` using formula = 'y ~ x'
```



Discussion and Conclusion