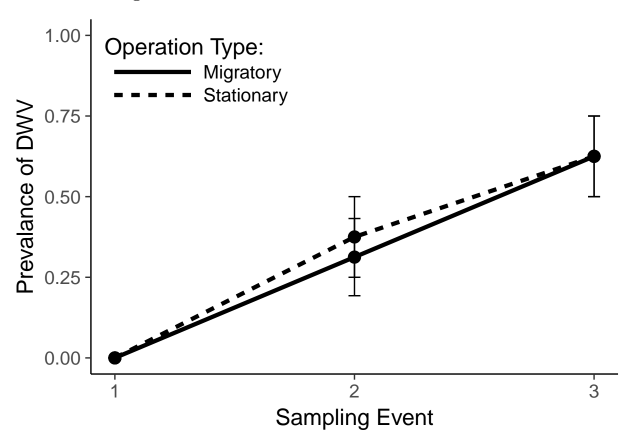
Migratory Stationary Technical Report

P. Alexander Burnham & Samantha A. Alger October 24, 2017

EXPERIMENT I - (MIGRATORY vs STATIONARY)

Deformed Wing Virus Prevalence:



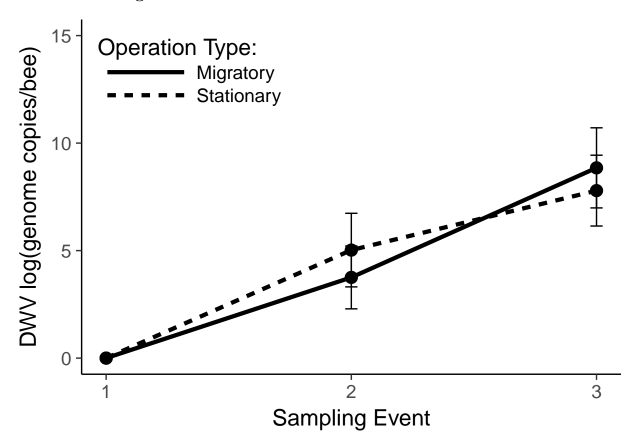
```
Treatment SamplingEvent n
                                   mean
     Migratory
                            1 16 0.0000 0.0000000 0.0000000
## 1
## 2
     Migratory
                            2 16 0.3125 0.4787136 0.1196784
## 3 Migratory
                            3 16 0.6250 0.5000000 0.1250000
                            1 16 0.0000 0.0000000 0.0000000
## 4 Stationary
                            2 16 0.3750 0.5000000 0.1250000
## 5 Stationary
## 6 Stationary
                            3 16 0.6250 0.5000000 0.1250000
```

Initial Time Step One Analaysis (Chi Square)

```
##
## Chi-squared test for given probabilities
##
## data: x
## X-squared = 0, df = 1, p-value = 1
```

Full repeated measures Analaysis (GLM)

Deformed Wing Virus Load:

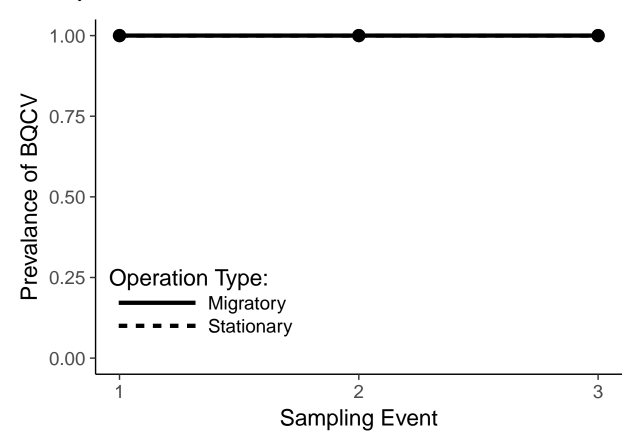


##		Treatment	SamplingEvent	n	mean	sd	se
##	1	Migratory	1	16	0.000000	0.000000	0.000000
##	2	Migratory	2	16	3.753490	5.849113	1.462278
##	3	Migratory	3	16	8.848642	7.451475	1.862869
##	4	Stationary	1	16	0.000000	0.000000	0.000000
##	5	Stationary	2	16	5.023841	6.845731	1.711433
##	6	Stationary	3	16	7.789912	6.587571	1.646893

Initial Time Step One Analaysis (ANOVA)

Full repeated measures Analaysis (Rep ANOVA)

Black Queen Cell Virus Prevalence:



```
##
      Treatment SamplingEvent n mean sd se
## 1
     Migratory
                             1 16
                                        0
                                     1
## 2
      Migratory
                             2 16
                                        0
                                           0
## 3 Migratory
                             3 16
## 4 Stationary
                             1 16
                                        0
                                           0
                             2 16
                                        0
                                           0
## 5 Stationary
                                     1
## 6 Stationary
                             3 16
                                        0
```

Initial Time Step One Analaysis (Chi Square)

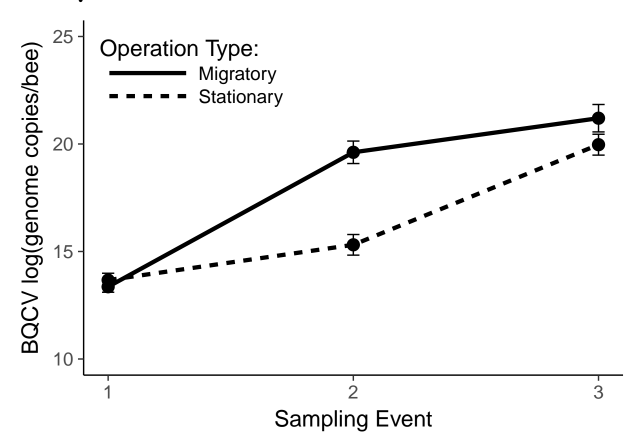
```
##
## Chi-squared test for given probabilities
##
```

```
## data: x
## X-squared = 0, df = 1, p-value = 1
```

Full repeated measures Analaysis (GLM)

```
#BQCV prevalence using glmer
#Fullmod2 <- glmer(data=MigStatExp_1, formula = BQCVbinary~Treatment * SamplingEvent + (1/ID), family =
#Anova(Fullmod2)</pre>
```

Black Queen Cell Virus Load:



Treatment SamplingEvent n mean ## 1 Migratory 1 16 13.36015 1.021698 0.2554245 ## 2 Migratory 2 16 19.61332 2.090001 0.5225002 ## 3 Migratory 3 16 21.19942 2.549945 0.6374862 1 16 13.67099 1.269596 0.3173990 ## 4 Stationary ## 5 Stationary 2 16 15.31059 1.918286 0.4795715 3 16 19.96779 1.942679 0.4856697 ## 6 Stationary

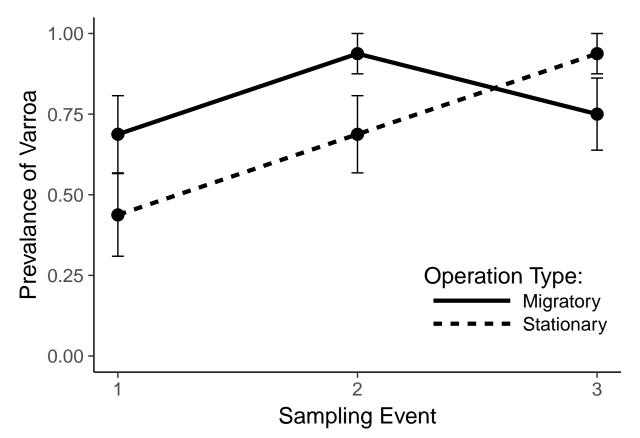
Initial Time Step One Analaysis (ANOVA)

```
## Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_1_T1$Treatment 1 0.77 0.773 0.582 0.451
## Residuals 30 39.84 1.328
```

Full repeated measures Analaysis (Rep ANOVA)

```
## Analysis of Deviance Table (Type II Wald chisquare tests)
##
## Response: logBQCV
##
                             Chisq Df Pr(>Chisq)
## Treatment
                           16.4884
                                   1 4.895e-05 ***
## SamplingEvent
                          187.2346
                                       < 2.2e-16 ***
## Treatment:SamplingEvent
                            2.2293
                                          0.1354
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Varroa Mite Prevalence:



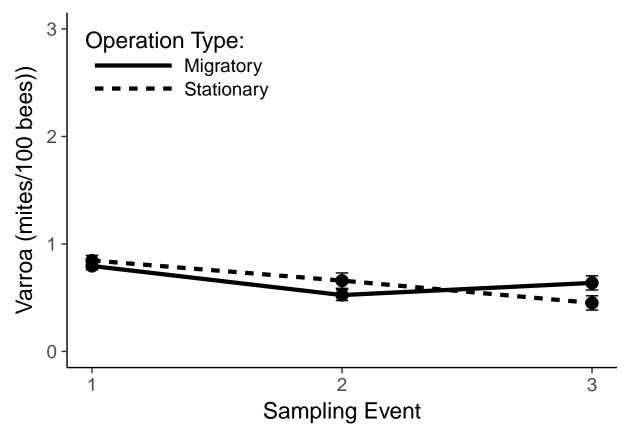
```
Treatment SamplingEvent
##
                                    {\tt mean}
                                                 sd
## 1
     Migratory
                             1 16 0.6875 0.4787136 0.1196784
## 2
     Migratory
                             2 16 0.9375 0.2500000 0.0625000
## 3 Migratory
                             3 16 0.7500 0.4472136 0.1118034
## 4 Stationary
                             1 16 0.4375 0.5123475 0.1280869
                             2 16 0.6875 0.4787136 0.1196784
## 5 Stationary
                             3 16 0.9375 0.2500000 0.0625000
## 6 Stationary
```

Initial Time Step One Analaysis (Chi Square)

```
##
## Pearson's Chi-squared test with Yates' continuity correction
##
```

```
## data: x
## X-squared = 1.1429, df = 1, p-value = 0.285
Full repeated measures Analaysis (GLM)
## Analysis of Deviance Table (Type II Wald chisquare tests)
##
## Response: VarroaBinary
##
                           Chisq Df Pr(>Chisq)
                          1.2899
                                       0.25607
## Treatment
                                  1
## SamplingEvent
                          4.8963
                                  1
                                       0.02691 *
## Treatment:SamplingEvent 3.2100
                                  1
                                       0.07319 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Varroa Mite Load:



```
##
      Treatment SamplingEvent
                                      mean
                                                  sd
                               n
                            1 16 0.7949475 0.1438689 0.03596722
## 1
     Migratory
## 2
     Migratory
                            2 16 0.5245116 0.2034428 0.05086071
## 3 Migratory
                            3 16 0.6375294 0.2638807 0.06597017
## 4 Stationary
                            1 16 0.8461480 0.1903694 0.04759234
## 5 Stationary
                            2 16 0.6589632 0.2858586 0.07146466
## 6 Stationary
                            3 16 0.4513384 0.2653305 0.06633262
```

Initial Time Step One Analaysis (ANOVA)

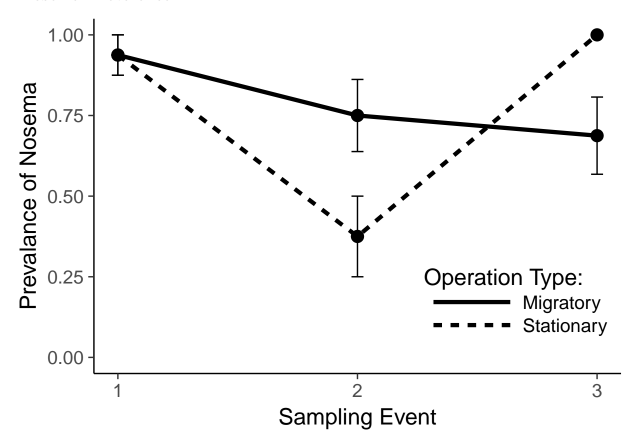
```
## Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_1_T1$Treatment 1 0.0210 0.02097 0.737 0.398
## Residuals 30 0.8541 0.02847

Full repeated measures Analaysis (Rep ANOVA)
## Analysis of Deviance Table (Type II Wald chisquare tests)
```

Analysis of Deviance Table (Type II Wald chisquare tests)
##
Response: Varroa
Chisq Df Pr(>Chisq)
Treatment 0.0000 1 0.99741
SamplingEvent 25.2426 1 5.055e-07 ***
Treatment:SamplingEvent 4.6647 1 0.03079 *

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

Nosema Prevalence:



Treatment SamplingEvent n mean sd se ## 1 Migratory 1 16 0.9375 0.2500000 0.0625000 ## 2 Migratory 2 16 0.7500 0.4472136 0.1118034 ## 3 Migratory 3 16 0.6875 0.4787136 0.1196784 ## 4 Stationary 1 16 0.9375 0.2500000 0.0625000 ## 5 Stationary 2 16 0.3750 0.5000000 0.1250000 ## 6 Stationary 3 16 1.0000 0.0000000 0.0000000

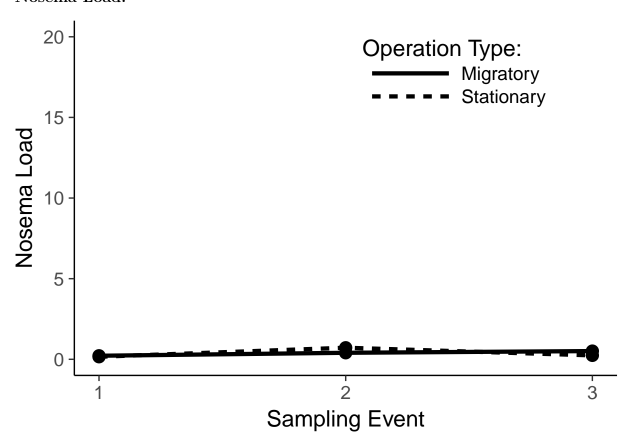
Initial Time Step One Analaysis (Chi Square)

```
##
## Pearson's Chi-squared test
##
## data: x
## X-squared = 0, df = 1, p-value = 1
```

Full repeated measures Analaysis (GLM)

```
## Analysis of Deviance Table (Type II Wald chisquare tests)
##
## Response: NosemaBinary
## Chisq Df Pr(>Chisq)
## Treatment 0.0367 1 0.8481
## SamplingEvent 0.6236 1 0.4297
## Treatment:SamplingEvent 2.3758 1 0.1232
```

Nosema Load:



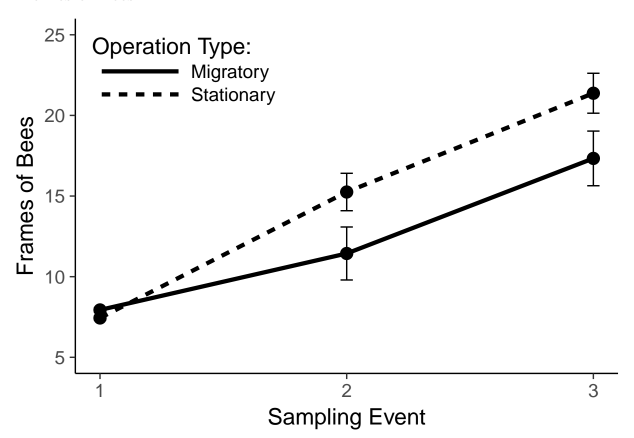
Treatment SamplingEvent n mean sd Migratory 1 16 0.2202589 0.2608239 0.06520598 Migratory 2 16 0.4034428 0.3788187 0.09470467 3 16 0.5099130 0.3585029 0.08962572 ## 3 Migratory ## 4 Stationary 1 16 0.1575894 0.2394642 0.05986605 2 16 0.7177525 0.3956736 0.09891840 ## 5 Stationary ## 6 Stationary 3 16 0.2422378 0.1773289 0.04433223

Initial Time Step One Analaysis (ANOVA)

```
## Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_1_T1$Treatment 1 0.0314 0.03142 0.501 0.484
## Residuals 30 1.8806 0.06269
```

Full repeated measures Analaysis (Rep ANOVA)

Frames of Bees:

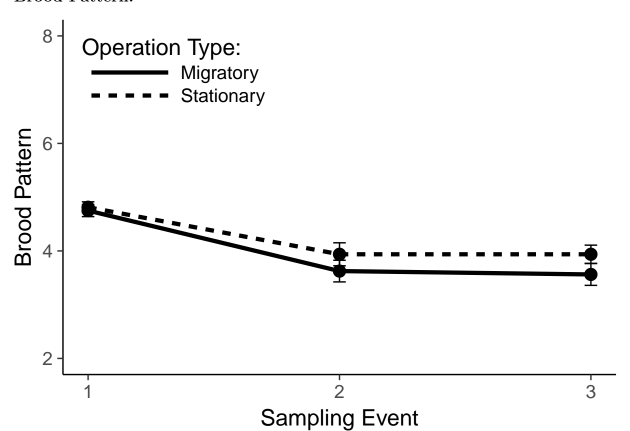


```
##
     Treatment SamplingEvent n
                                     mean
                                                 sd
                            1 16 7.93750 0.5737305 0.1434326
## 1 Migratory
     Migratory
                            2 16 11.43750 6.5723537 1.6430884
## 3 Migratory
                            3 16 17.33333 6.7788186 1.6947046
## 4 Stationary
                            1 16 7.43750 0.5123475 0.1280869
                            2 16 15.25000 4.6404023 1.1601006
## 5 Stationary
## 6 Stationary
                           3 16 21.37500 4.9514308 1.2378577
```

Initial Time Step One Analaysis (ANOVA)

```
##
                            Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_1_T1$Treatment 1 2.000 2.0000 6.761 0.0143 *
## Residuals
                            30 8.875 0.2958
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
\hbox{\tt \#\# Warning in chisq.test(x): Chi-squared approximation may be incorrect}
##
## Pearson's Chi-squared test
##
## data: x
## X-squared = 5.8889, df = 2, p-value = 0.05263
Full repeated measures Analaysis (Rep ANOVA)
## Analysis of Deviance Table (Type II Wald chisquare tests)
## Response: FOB
                             Chisq Df Pr(>Chisq)
##
## Treatment
                            3.5968 1
                                        0.05789 .
                                         < 2e-16 ***
## SamplingEvent
                          152.8381 1
## Treatment:SamplingEvent 5.6507 1
                                        0.01745 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Brood Pattern:



```
## Treatment SamplingEvent n mean sd se

## 1 Migratory 1 16 4.7500 0.4472136 0.1118034

## 2 Migratory 2 16 3.6250 0.8062258 0.2015564

## 3 Migratory 3 16 3.5625 0.8139410 0.2034853

## 4 Stationary 1 16 4.8125 0.4031129 0.1007782

## 5 Stationary 2 16 3.9375 0.8539126 0.2134781

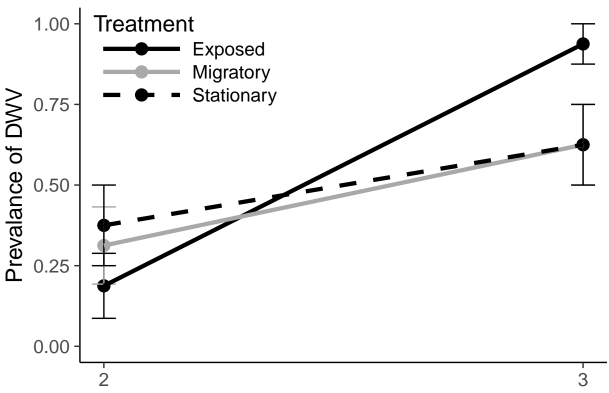
## 6 Stationary 3 16 3.9375 0.6800735 0.1700184
```

Initial Time Step One Analaysis (ANOVA)

```
## Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_1_T1$Treatment 1 0.031 0.03125 0.172 0.681
## Residuals 30 5.437 0.18125
```

EXPERIMENT II - (EXPOSED vs STATIONARY)

Deformed Wing Virus Prevalence:



Sampling Event

```
##
     Treatment SamplingEvent n
                                   mean
## 1
       Exposed
                            2 16 0.1875 0.4031129 0.1007782
## 2
       Exposed
                            3 16 0.9375 0.2500000 0.0625000
## 3 Migratory
                            2 16 0.3125 0.4787136 0.1196784
## 4 Migratory
                            3 16 0.6250 0.5000000 0.1250000
## 5 Stationary
                            2 16 0.3750 0.5000000 0.1250000
                            3 16 0.6250 0.5000000 0.1250000
## 6 Stationary
```

Initial Time Step One Analaysis (Chi Square)

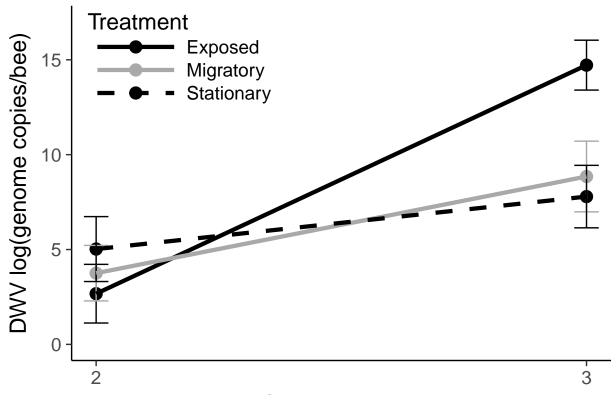
```
## Warning in chisq.test(x): Chi-squared approximation may be incorrect
##
##
   Pearson's Chi-squared test with Yates' continuity correction
##
## X-squared = 0.61836, df = 1, p-value = 0.4317
```

Full repeated measures Analaysis (GLM)

```
## Analysis of Deviance Table (Type II Wald chisquare tests)
##
```

Response: DWVbinary

Deformed Wing Virus Load:



Sampling Event

```
##
      Treatment SamplingEvent n
                                      mean
                                                 sd
## 1
       Exposed
                                  2.671716 6.184479 1.546120
## 2
                            3 16 14.718737 5.260002 1.315001
       Exposed
## 3 Migratory
                                 3.753490 5.849113 1.462278
## 4 Migratory
                            3 16 8.848642 7.451475 1.862869
                            2 16
## 5 Stationary
                                 5.023841 6.845731 1.711433
## 6 Stationary
                            3 16 7.789912 6.587571 1.646893
```

Initial Time Step One Analaysis (ANOVA)

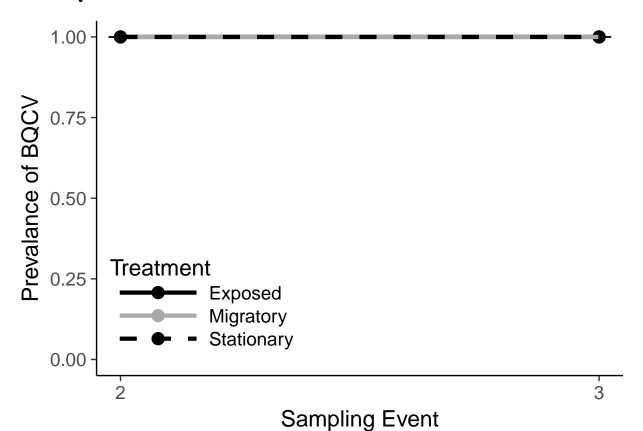
```
## Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_2_analysis_T2$Treatment 1 44.3 44.26 1.04 0.316
## Residuals 30 1276.7 42.56
```

Full repeated measures Analaysis (Rep ANOVA)

##

```
## Error: ID
##
            Df Sum Sq Mean Sq F value Pr(>F)
                 83.8
## Treatment 1
                       83.78
                                2.056 0.162
## Residuals 30 1222.6
                        40.75
## Error: Within
                          Df Sum Sq Mean Sq F value
                                     877.7 23.510 3.57e-05 ***
## SamplingEvent
                           1 877.7
## Treatment:SamplingEvent 1 344.5
                                      344.5
                                             9.229
                                                     0.0049 **
## Residuals
                          30 1120.0
                                      37.3
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Analysis of Deviance Table (Type II Wald chisquare tests)
## Response: logDWV
                            Chisq Df Pr(>Chisq)
## Treatment
                           2.0559 1
                                      0.151622
## SamplingEvent
                          23.5099 1
                                     1.243e-06 ***
## Treatment:SamplingEvent 9.2288 1
                                      0.002382 **
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Black Queen Cell Virus Prevalence:



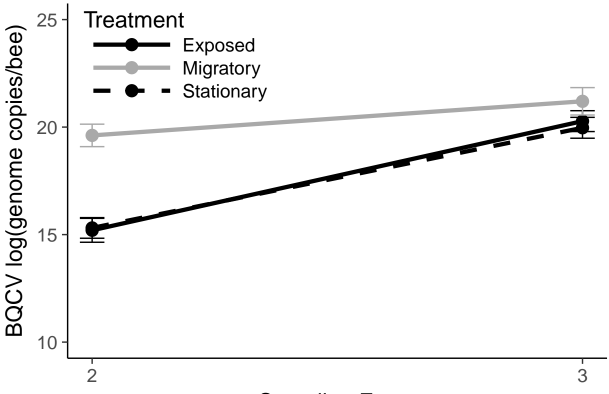
```
## 2 Exposed 3 16 1 0 0 ## 3 Migratory 2 16 1 0 0 ## 4 Migratory 3 16 1 0 0 ## 5 Stationary 2 16 1 0 0 ## 6 Stationary 3 16 1 0 0
```

Initial Time Step One Analaysis (Chi Square)

```
##
## Chi-squared test for given probabilities
##
## data: x
## X-squared = 0, df = 1, p-value = 1
```

Full repeated measures Analaysis (GLM)

Black Queen Cell Virus Load:



Sampling Event

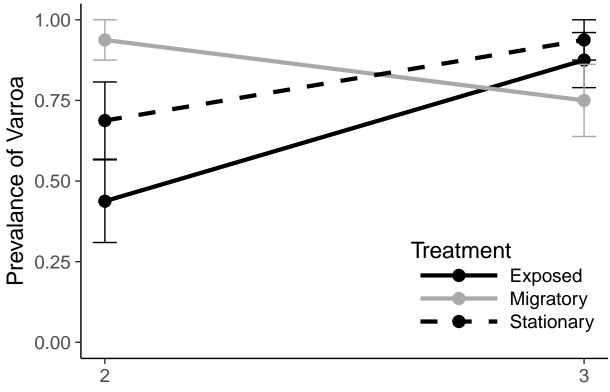
##		Treatment	SamplingEvent	n	mean	sd	se
##	1	Exposed	2	16	15.19960	2.222482	0.5556204
##	2	Exposed	3	16	20.27747	1.940819	0.4852047
##	3	Migratory	2	16	19.61332	2.090001	0.5225002
##	4	Migratory	3	16	21.19942	2.549945	0.6374862
##	5	Stationary	2	16	15.31059	1.918286	0.4795715
##	6	Stationary	3	16	19.96779	1.942679	0.4856697

Initial Time Step One Analaysis (ANOVA)

```
## MigStatExp_2_analysis_T2$Treatment 1 0.1 0.099 0.023 0.881 ## Residuals 30 129.3 4.310
```

Full repeated measures Analaysis (Rep ANOVA)

Varroa Mite Prevalence:



Sampling Event

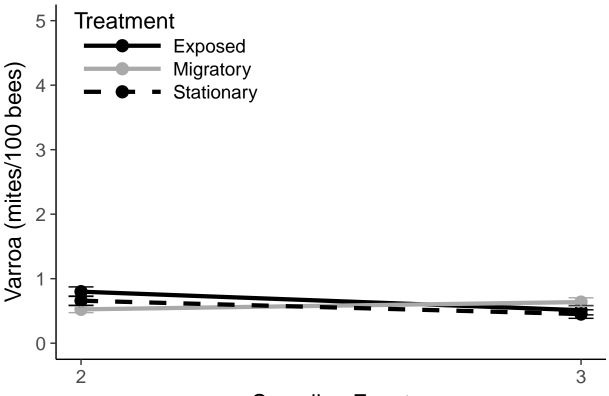
##		Treatment	SamplingEvent	n	mean	sd	se
##	1	Exposed	2	16	0.4375	0.5123475	0.12808688
##	2	Exposed	3	16	0.8750	0.3415650	0.08539126
##	3	Migratory	2	16	0.9375	0.2500000	0.06250000
##	4	Migratory	3	16	0.7500	0.4472136	0.11180340
##	5	Stationary	2	16	0.6875	0.4787136	0.11967839
##	6	Stationary	3	16	0.9375	0.2500000	0.06250000

Initial Time Step One Analaysis (Chi Square)

```
##
## Pearson's Chi-squared test with Yates' continuity correction
##
## data: x
## X-squared = 1.1429, df = 1, p-value = 0.285
```

Full repeated measures Analaysis (GLM)

Varroa Mite Load:



Sampling Event

##		Treatment	SamplingEvent	n	mean	sd	se
##	1	Exposed	2	16	0.7986416	0.2953717	0.07384293
##	2	Exposed	3	16	0.5112203	0.2887780	0.07219449
##	3	Migratory	2	16	0.5245116	0.2034428	0.05086071
##	4	Migratory	3	16	0.6375294	0.2638807	0.06597017

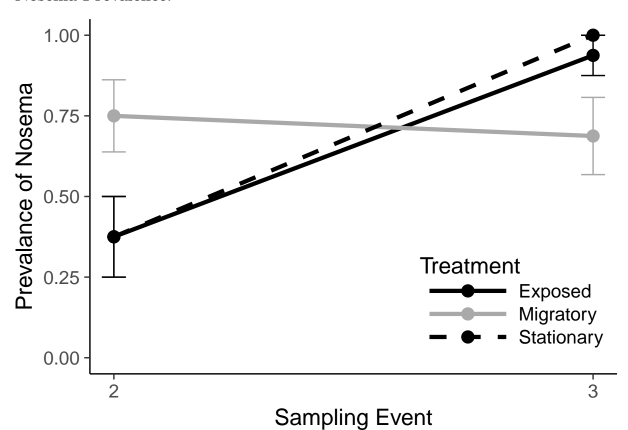
```
## 5 Stationary 2 16 0.6589632 0.2858586 0.07146466
## 6 Stationary 3 16 0.4513384 0.2653305 0.06633262
```

Initial Time Step One Analaysis (ANOVA)

```
## Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_2_analysis_T2$Treatment 1 0.1561 0.15608 1.848 0.184
## Residuals 30 2.5344 0.08448
```

Full repeated measures Analaysis (Rep ANOVA)

Nosema Prevalence:



Treatment SamplingEvent n mean sd se ## 1 Exposed 2 16 0.3750 0.5000000 0.1250000 ## 2 Exposed 3 16 0.9375 0.2500000 0.0625000

```
## 4 Migratory
                           3 16 0.6875 0.4787136 0.1196784
                          2 16 0.3750 0.5000000 0.1250000
## 5 Stationary
## 6 Stationary
                           3 16 1.0000 0.0000000 0.0000000
Initial Time Step One Analaysis (Chi Square)
##
   Pearson's Chi-squared test
##
##
## data: x
## X-squared = 0, df = 1, p-value = 1
Full repeated measures Analaysis (GLM)
## Warning in checkConv(attr(opt, "derivs"), opt$par, ctrl = control$checkConv, : Model is nearly unide
## - Rescale variables?
## Analysis of Deviance Table (Type II Wald chisquare tests)
```

2 16 0.7500 0.4472136 0.1118034

Chisq Df Pr(>Chisq)

0.999946

0.949451

0.005309 **

0.0000 1

7.7711 1

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

3 Migratory

Response: NosemaBinary

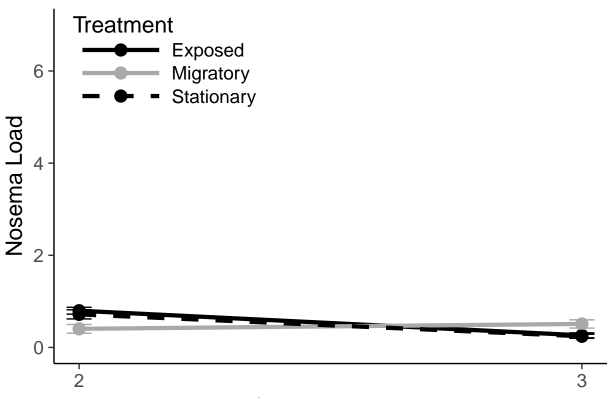
Treatment:SamplingEvent 0.0040 1

##

Treatment

SamplingEvent

Nosema Load:



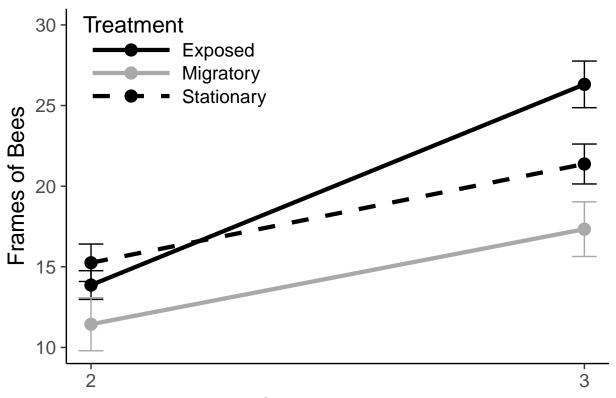
Sampling Event

```
Treatment SamplingEvent n
##
## 1
       Exposed
                            2 16 0.7965278 0.2966401 0.07416003
## 2
       Exposed
                            3 16 0.2621866 0.2162779 0.05406948
## 3 Migratory
                            2 16 0.4034428 0.3788187 0.09470467
## 4 Migratory
                            3 16 0.5099130 0.3585029 0.08962572
                            2 16 0.7177525 0.3956736 0.09891840
## 5 Stationary
## 6 Stationary
                            3 16 0.2422378 0.1773289 0.04433223
```

Initial Time Step One Analaysis (ANOVA)

```
## Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_2_analysis_T2$Treatment 1 0.050 0.04964 0.406 0.529
## Residuals 30 3.668 0.12228
```

Frames of Bees:



Sampling Event

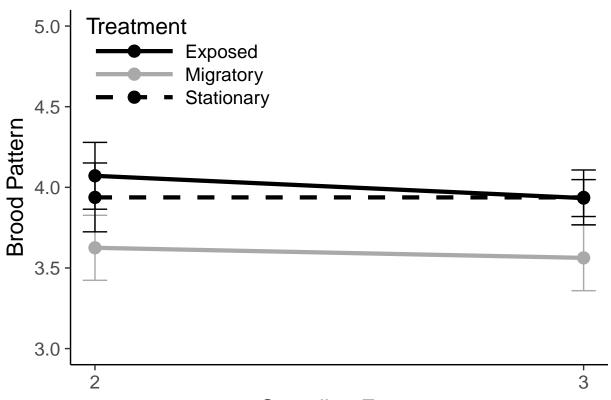
```
Treatment SamplingEvent n
##
                                     mean
## 1
       Exposed
                            2 16 13.86667 3.563038 0.8907594
## 2
       Exposed
                            3 16 26.31250 5.781796 1.4454490
## 3 Migratory
                            2 16 11.43750 6.572354 1.6430884
## 4 Migratory
                            3 16 17.33333 6.778819 1.6947046
                            2 16 15.25000 4.640402 1.1601006
## 5 Stationary
## 6 Stationary
                            3 16 21.37500 4.951431 1.2378577
```

Initial Time Step One Analaysis (ANOVA)

```
## Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_2_analysis_T2$Treatment 1 14.8 14.81 0.858 0.362
## Residuals 29 500.7 17.27
## 1 observation deleted due to missingness
```

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

Brood Pattern:



Sampling Event

```
##
     Treatment SamplingEvent n
                                     mean
                                                 sd
## 1
       Exposed
                            2 16 4.071429 0.8287419 0.2071855
## 2
        Exposed
                            3 16 3.933333 0.4577377 0.1144344
## 3 Migratory
                            2 16 3.625000 0.8062258 0.2015564
## 4 Migratory
                            3 16 3.562500 0.8139410 0.2034853
## 5 Stationary
                            2 16 3.937500 0.8539126 0.2134781
## 6 Stationary
                            3 16 3.937500 0.6800735 0.1700184
```

Initial Time Step One Analaysis (ANOVA)

```
## Df Sum Sq Mean Sq F value Pr(>F)
## MigStatExp_2_analysis_T2$Treatment 1 0.134 0.1339 0.189 0.667
## Residuals 28 19.866 0.7095
## 2 observations deleted due to missingness
```

```
## Analysis of Deviance Table (Type II Wald chisquare tests)
##
## Response: BroodPattern
## Chisq Df Pr(>Chisq)
## Treatment 0.1331 1 0.7152
```

```
## SamplingEvent 0.1815 1 0.6701
## Treatment:SamplingEvent 0.2156 1 0.6424
```

OVER-ARCHING STUDY-WIDE ANALYSES

Tempurature and Rainfall Date between CA and NC

Avg Temp. between NC and CA

```
## Df Sum Sq Mean Sq F value Pr(>F)
## State 1 2.7 2.67 0.059 0.809
## Residuals 52 2350.7 45.21
```

Avg Windspeed between NC and CA

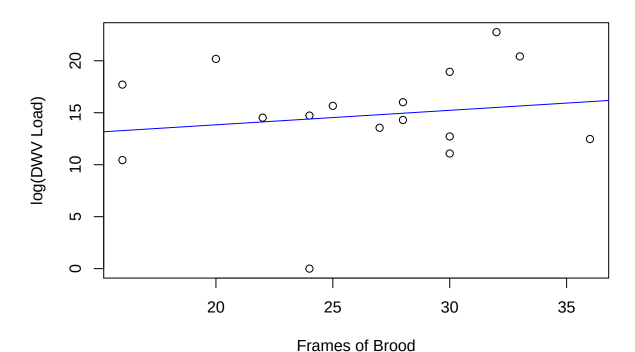
Rainfall between NC and CA

```
## Df Sum Sq Mean Sq F value Pr(>F)
## State 1 0.040 0.04000 1.323 0.256
## Residuals 49 1.482 0.03024
## 3 observations deleted due to missingness
```

Linear regression between DWV load and Frames of bees

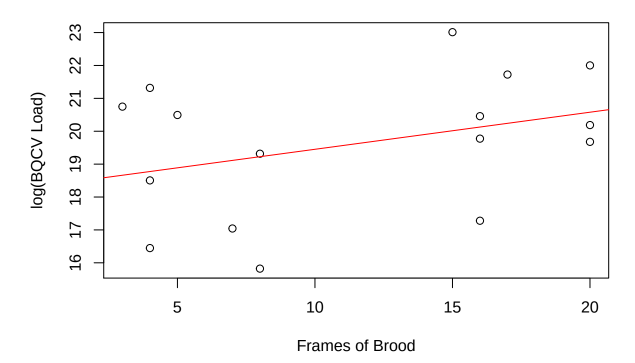
Regression between DWV and FOB

T3 Exposed



```
##
## Call:
## lm(formula = MigStat_3_Exp$logDWV ~ MigStat_3_Exp$FOB)
##
## Residuals:
##
        Min
                  1Q
                       Median
                                    3Q
                                            Max
   -14.3965 -2.5956
                       0.3709
                                3.8857
##
                                         7.2414
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                      11.0517
                                  6.4637
                                            1.71
                                                    0.109
                                            0.58
## MigStat_3_Exp$FOB
                       0.1394
                                  0.2403
                                                    0.571
## Residual standard error: 5.38 on 14 degrees of freedom
## Multiple R-squared: 0.02347,
                                    Adjusted R-squared: -0.04628
## F-statistic: 0.3364 on 1 and 14 DF, p-value: 0.5711
```

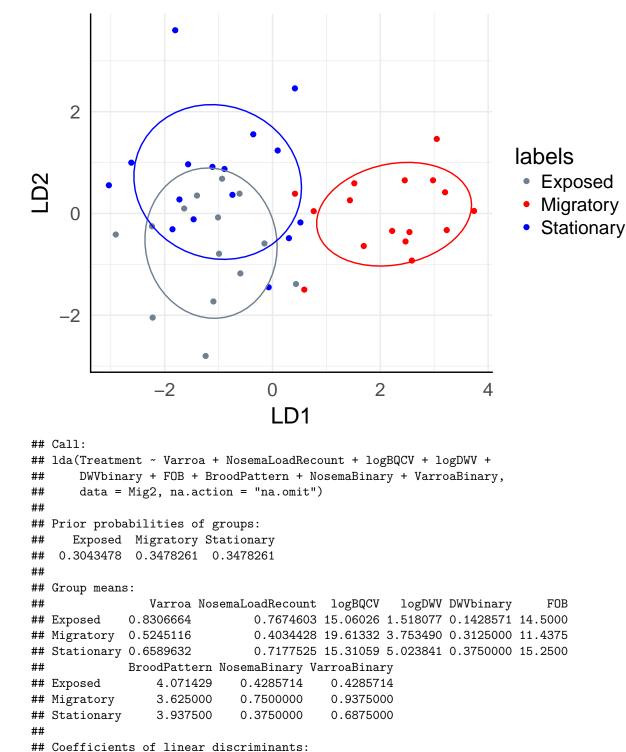
T2 Migratory



```
##
## Call:
## lm(formula = MigStat_1_Mig$logBQCV ~ MigStat_1_Mig$FOB)
##
## Residuals:
##
       Min
                1Q Median
                                3Q
                                       Max
   -3.4030 -1.1930 -0.0894
##
                           1.5144
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                     18.32414
                                 1.04016
                                         17.617 5.96e-11 ***
## MigStat_1_Mig$FOB
                     0.11272
                                 0.07947
                                           1.418
                                                    0.178
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 2.023 on 14 degrees of freedom
## Multiple R-squared: 0.1256, Adjusted R-squared: 0.06318
## F-statistic: 2.012 on 1 and 14 DF, p-value: 0.178
```

Linear Classification Anlaysis for all Vars on Grouping into Treatments (3 time points):

LDA for Time Point 2:



LD2

LD1

NosemaLoadRecount 0.08804005 -4.3858680

0.97951917 -2.4086816

0.53052423 -0.1380832

-0.13171807 0.4506278

1.50597482 -5.0280473

-0.11430496 0.1567110

##

Varroa

logBQCV

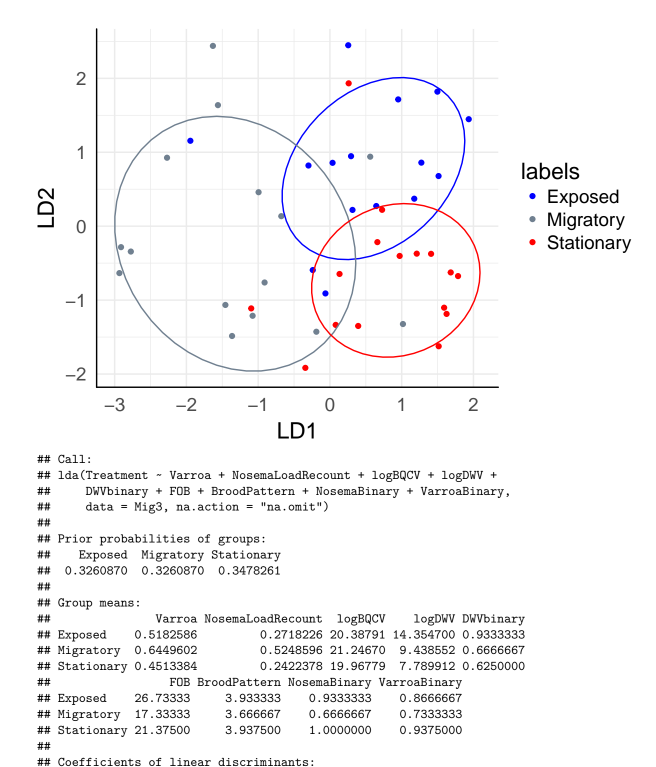
DWVbinary

logDWV

FOB

```
## BroodPattern
                      0.38830031 -0.9453298
## NosemaBinary
                      0.46686771 -3.2848585
## VarroaBinary
                      1.95470288 -0.1136285
##
## Proportion of trace:
##
     LD1
            LD2
## 0.8836 0.1164
## Warning in cbind(Mig2$Treatment, CA1$class): number of rows of result is
## not a multiple of vector length (arg 2)
##
               predicted
                Exposed Migratory Stationary
## actual
##
                      6
     Exposed
                                3
                      2
                                           3
##
     Migratory
                               11
##
     Stationary
                      7
                                3
                                           6
##
               predicted
## actual
                Exposed Migratory Stationary
##
    Exposed
                 0.3750
                           0.1875
                                      0.4375
##
    Migratory
                 0.1250
                           0.6875
                                      0.1875
##
    Stationary 0.4375
                           0.1875
                                      0.3750
##
## Call:
## adonis(formula = envdist1 ~ Treatment, data = Mig2x)
##
## Permutation: free
## Number of permutations: 999
## Terms added sequentially (first to last)
##
##
             Df SumsOfSqs MeanSqs F.Model
                                                R2 Pr(>F)
## Treatment 2
                  0.57735 0.288674
                                     5.077 0.19103 0.001 ***
## Residuals 43
                  2.44495 0.056859
                                           0.80897
                  3.02230
                                           1.00000
## Total
            45
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

LDA for Time Point 3:



LD2

1.45159756

LD1

-1.76281514 1.80931348

-0.12388210 0.03298634

-0.19425568 0.22967117

2.01214636 -2.04228774

0.12074871 0.09144119

##

Varroa

logBQCV

DWVbinary

logDWV

FOB

NosemaLoadRecount -2.31126434

```
## BroodPattern
                      0.15738416 -0.28505404
## NosemaBinary
                     -0.01490752 0.52084598
## VarroaBinary
                      0.74061573 0.43930995
##
## Proportion of trace:
##
     LD1
            LD2
## 0.6873 0.3127
## Warning in cbind(Mig3$Treatment, CA2$class): number of rows of result is
## not a multiple of vector length (arg 2)
##
              predicted
                Exposed Migratory Stationary
## actual
##
                      9
     Exposed
                                3
                      5
                                7
                                           4
##
     Migratory
##
     Stationary
                      3
                                2
                                          11
##
              predicted
## actual
                Exposed Migratory Stationary
    Exposed
                 0.5625
                           0.1875
##
    Migratory
                0.3125
                           0.4375
                                      0.2500
##
    Stationary 0.1875
                           0.1250
                                      0.6875
##
## Call:
## adonis(formula = envdist ~ Treatment, data = Mig3x)
##
## Permutation: free
## Number of permutations: 999
## Terms added sequentially (first to last)
##
             Df SumsOfSqs MeanSqs F.Model
##
                                                R2 Pr(>F)
## Treatment 2
                  0.40211 0.201054 4.7793 0.18187 0.002 **
## Residuals 43
                 1.80891 0.042068
                                           0.81813
                                           1.00000
## Total
            45
                  2.21102
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```