Supporting Materials for 'The Integration of Norse-Derived Terms in English'

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1 Introduction

The analysis covers the following sources:

- FCPC Peterborough, 1154 (12th C)
- Ormulum South Lincolnshire, ca. 1180 (12th ${\rm C})$
- Havelok, composed in Lincolnshire, but manuscript from West Norfolk 13th century (13th C)
- Cursor mundi, c. 1300 (14th C)
- Gawain-poet Cheshire / Staffordshire, ca. 1380 (14th C)
- St Erkenwald, written late fourteenth or the early fifteenth century, manuscript from 1477 (15th C)
- Mannyng South Lincolnshire, ca. 1450 (15th C)
- Wars of Alexander northern England, ca. 1450 (15th C)
- \bullet Texts from Lincolnshire, Norfolk and Nottinghamshire from the Corpus of Middle English (1399-1525), (15th C)
- Texts by Richard Rolle (before 1465), (15thC)

The data analysed here is the product of several processing steps. The file data/SharedIntegrationOfCognatesData.xls has the original transcribed data. This is cleaned and processed by the script analysis/analyseTextDistances.py, which draws some code from analysis/CLTSFeatureBasedAlignment.py which is mainly contributed by Johann Mattis List (see https://calc.hypotheses.org/1962 and https://gist.github.com/LinguList/7fac44813572f65259c872ef89fa64ad). The script calculates the distances between pairs of Norse and English forms according to three measures:

- Simple distance from Keller (2020).
- A feature-based distance.
- A historical distance that uses the likelihood of one segment historically replacing another.

Each row in the data represents a comparison between a Norse form and an English form within a given cognate set, including the three measures of distance and frequency of occurance in each source.

1.1 Variables

The variables in the data (and some that are calculated in the script below) are described as follows:

- "Set": the cognate set the pair of comparisons belong to.
- "Class": the word class the set can appear as (and some binary variables representing the same information).
- "NorseLexeme", "EngLexeme", "NorseForm", "EngForm": The lexemes and full forms for Norse and English terms.
- "NorseFormDiagnostic", "EngFormDiagnostic": the relevant parts of the form that are diagnostic of the etymology.
- "Alignment", "Feature Alignment", "Historical Alignment": multiple sequence alignment of the forms according to the three measures.
- "RawDistance", "NormDistance", "RawFeatureDistance", "NormFeatureDistance", "RawHistoricalDistance", "NormHistoricalDistance": Raw and normed distances between
- "NFreq...", "EFreq...": Frequencies of the Norse and English forms in each source.
- "Alliteration": only true if the source uses alliterative verse and the Norse form and the English form do not start with segments in the same alliterative category. That is, it is true if part of the decision about which form to use might be influenced by the need to alliterate. The alliterative texts include: Gawain, St Erkenwold, Wars of Alexander.
- "totalNFreq": The total Norse frequency across all sources.
- "total EFreq": The total English frequency across all sources.
- "NorseProp": The proportion of total Norse frequency compared to total Norse Frequency + total English Frequency.
- "NorseDiagnosticScore": Whether or not the forms differ according to a specific segments that are characteristicly diagnostic of Norse etymology.

2 Load Libraries

```
library(openxlsx)
library(sjPlot)
library(lme4)
library(mgcv)
library(party)
library(ggplot2)
library(phangorn)
library(gridExtra)
library(GGally)
library(brms)
library(ggeffects)
library(gamm4)
```

3 Load data

Load data created by the python program and convert variables to their proper type:

```
d = read.xlsx("../data/IntegrationDistances.xlsx",1)
# Ignore numerals in set name
d$Set = gsub("[0-9]","",d$Set)
d$Set = as.factor(d$Set)
d$EngLexeme = factor(d$EngLexeme)
d$ELen = nchar(d$EngForm)
d$NLen = nchar(d$NorseForm)
norseFrequencyColumns = c("NFreqFCPC", "NFreqGawainPoet",
                          "NFreqGenAndEx", "NFreqHavelok",
                          "NFreqMannyng", "NFreqOrmulum",
                          'NFreqWarsAlexander',
                          "NFreqStErkenwald", "NFreqCursorMundi",
                          "NFreqLinc", "NFreqNort", "NFreqRolle")
englishFrequencyColumns = c("EFreqFCPC","EFreqGawainPoet",
                            "EFreqGenAndEx", "EFreqHavelok",
                            "EFreqMannyng", "EFreqOrmulum",
                            "EFreqWarsAlexander",
                            "EFreqStErkenwald", "EFreqCursorMundi",
                            "EFreqLinc", "EFreqNort", "EFreqRolle")
```

Convert frequencies to numeric type:

```
for(col in c(norseFrequencyColumns,englishFrequencyColumns)){
  d[,col] = as.numeric(d[,col])
}
```

Calculate the total frequency across all sources and the proportion of Norse forms compared to all forms:

```
d$totalNFreq = rowSums(d[,norseFrequencyColumns],na.rm = T)
d$totalEFreq = rowSums(d[,englishFrequencyColumns],na.rm = T)
d$NorseProp = d$totalNFreq/ rowSums(d[,c("totalNFreq","totalEFreq")],na.rm = T)

d = d[!is.na(d$NorseProp),]
```

Scale the distances to lie between 0 and 1:

```
d$NormDistance = as.numeric(d$NormDistance)
d$NormFeatureDistance = as.numeric(d$NormFeatureDistance)
```

```
d$NormHistoricalDistance = as.numeric(d$NormHistoricalDistance)

# Scale distances
normX = function(X){
    (X-min(X))/(max(X)-min(X))
}
d$NormDistance = normX(d$NormDistance)
d$NormFeatureDistance = normX(d$NormFeatureDistance)
d$NormHistoricalDistance = normX(d$NormHistoricalDistance)

d$NormDistance.rank =
    rank(d$NormDistance, ties.method = "max")
d$NormFeatureDistance.rank =
    rank(d$NormFeatureDistance, ties.method = "max")
d$NormHistoricalDistance.rank =
    rank(d$NormHistoricalDistance, ties.method = "max")
```

3.1 Examples

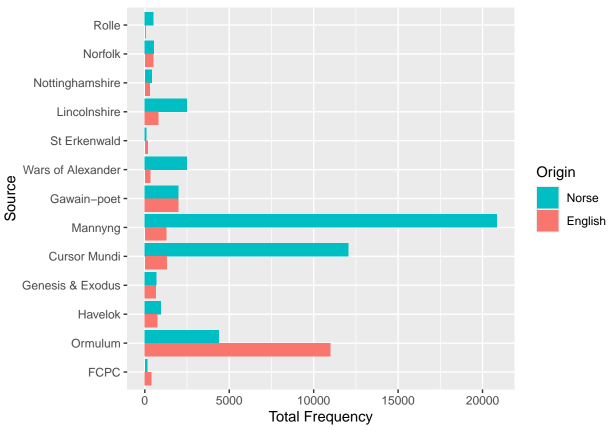
4 Descriptive statistics

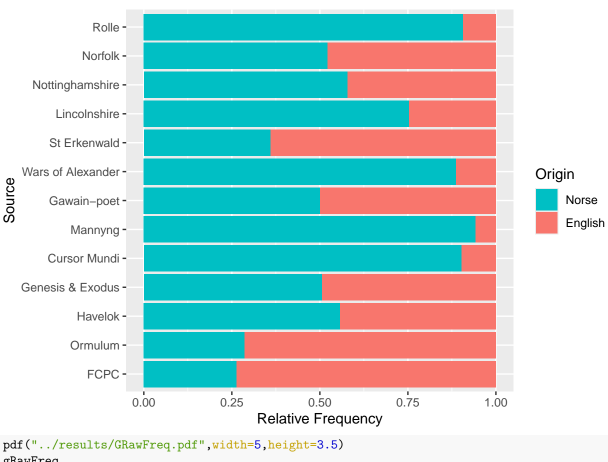
Number of English forms: 135 Number of Norse forms: 126 Number of comparisons: 1636

Number of sets: 67

Proportion of Norse and English terms (relative total frequency of each type):

```
dx = rbind(
 data.frame(
    Lang = "Norse",
    Freq = colSums(d[,norseFrequencyColumns],na.rm = T),
    Source = c("FCPC", "Gawain-poet", "Genesis & Exodus", "Havelok",
                "Mannyng", "Ormulum", "Wars of Alexander",
                "St Erkenwald", "Cursor Mundi",
               "Lincolnshire", "Nottinghamshire",
               "Norfolk", "Rolle")),
 data.frame(
    Lang = "English",
    Freq = colSums(d[,englishFrequencyColumns],na.rm = T),
    Source = c("FCPC", "Gawain-poet", "Genesis & Exodus", "Havelok",
                "Mannyng", "Ormulum", "Wars of Alexander",
                "St Erkenwald", "Cursor Mundi",
               "Lincolnshire", "Nottinghamshire",
               "Norfolk", "Rolle")))
dx$Source = factor(dx$Source,
 levels=c("FCPC","Ormulum","Havelok",
                 "Genesis & Exodus",
                 "Cursor Mundi", "Mannyng",
                 "Gawain-poet", "Wars of Alexander",
                 "St Erkenwald",
           "Lincolnshire", "Nottinghamshire",
               "Norfolk", "Rolle"))
gRawFreq = ggplot(dx,aes(x=Source,y=Freq,fill=Lang)) +
 geom_bar(stat="identity",position = 'dodge') +
 ylab("Total Frequency") +
 scale_fill_discrete(breaks=c('Norse', "English"),
                      name = "Origin") +
  coord_flip()
gRawFreq
```





```
gRawFreq
dev.off()
## pdf
##
pdf("../results/GPropFreq.pdf",width=5,height=3.5)
gPropFreq
dev.off()
## pdf
##
pdf("../results/GRawAndPropFreq.pdf",width=6,height=3)
  grid.arrange(gRawFreq+
               theme(legend.position = "none"),
             gPropFreq +
               theme(axis.text.y = element_blank(),
                      axis.title.y = element_blank()),
             ncol=2)
dev.off()
## pdf
Correlations between distance measures:
```

cor.test(d\$NormDistance,d\$NormFeatureDistance)

```
##
## Pearson's product-moment correlation
##
## data: d$NormDistance and d$NormFeatureDistance
## t = 95.432, df = 1634, p-value < 2.2e-16</pre>
```

```
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.9130857 0.9278599
## sample estimates:
         cor
## 0.9208025
cor.test(d$NormDistance,d$NormHistoricalDistance)
## Pearson's product-moment correlation
##
## data: d$NormDistance and d$NormHistoricalDistance
## t = 76.001, df = 1634, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.8717247 0.8931370
## sample estimates:
##
        cor
## 0.882889
cor.test(d$NormFeatureDistance,d$NormHistoricalDistance)
##
## Pearson's product-moment correlation
##
## data: d$NormFeatureDistance and d$NormHistoricalDistance
## t = 76.008, df = 1634, p-value < 2.2e-16
## alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.8717445 0.8931537
## sample estimates:
##
        cor
## 0.8829071
Function for extracting stats:
getStatReport = function(m0,m1,m2,m3,finalModel,outFile){
 modelComparison = as.data.frame(anova(m0,m1,m2,m3))
 modelComparison$pChi = round(modelComparison$`Pr(>Chisq)`,3)
 modelComparison$pChi[modelComparison$pChi==0] = "< 0.001"</pre>
 modelComparison$lldiff = NA
 modelComparison$lldiff[2:4] = diff(modelComparison$logLik)
 modelComparison = modelComparison[2:4,]
 modelComparison$Term = c("Linear", "Quadratic", "Cubic")
 coef = as.data.frame(summary(finalModel)$coefficients)
 rx = which(grepl("Norm",rownames(coef)))
  coef[rx, "Estimate"] = round(coef[rx, "Estimate"],3)
  coef[rx,"z value"] = round(coef[rx,"z value"],3)
  coef[,"Pr(>|z|)"] = round(coef[,"Pr(>|z|)"],3)
  coef[,"Pr(>|z|)"][coef[,"Pr(>|z|)"] ==0] = "< 0.001"
 modelComparison$Estimate = coef[rx,"Estimate"]
 modelComparison$z = coef[rx,"z value"]
 modelComparison$p = coef[rx,"Pr(>|z|)"]
 modelComparison = modelComparison[,
                                     c("Term", "BIC", "lldiff",
                                      "Chisq", "Df", "pChi",
                                       "Estimate", "z", "p")]
```

```
mcx = paste(paste0(c("Linear: ","Quadratic: ","Cubic: "),
        "beta = ",coef[rx,"Estimate"],
        ", z = ", coef[rx, "z value"],
        ", Wald p = ", coef[rx, "Pr(>|z|)"],
        ", LLDiff = ",round(modelComparison$lldiff,1),
        ", df = ", modelComparison$Df,
        ", p = ",modelComparison$pChi),collapse="; ")
  mcx = gsub("= <","<",mcx)</pre>
  cat(mcx,file=outFile)
 # mx = rbind(c("", "Model Comparison", "", "", "", "Model Estimate", "", ""),
         names (modelComparison),
         modelComparison)
 #
  modelComparison$BIC = round(modelComparison$BIC,1)
  modelComparison$lldiff = round(modelComparison$lldiff,1)
  modelComparison$Chisq = round(modelComparison$Chisq,1)
  modelComparison$Estimate = round(modelComparison$Estimate,2)
  write.csv(modelComparison,file=gsub("\\.txt",".csv",outFile))
  return(mcx)
}
```

5 Total frequency analysis

The analyses below predict the total frequency across all sources, using the orthographic form as the basis for observations.

5.1 Simple distance (total frequency)

We use nested model comparison to evaluate whether higher-order variables should be added (i.e. how non-linear the relationships is).

```
d$NormDistanceChosen = d$NormDistance
# Baseline model
m0 = glmer(cbind(totalNFreq,totalEFreq) ~
             1 + (1|Set) + (1|EngLexeme),
           data = d, family = "binomial")
# Add the distance measure
m1 = update(m0,~.+NormDistanceChosen)
# Add quadratic term
m2 = update(m1,~.+I(NormDistanceChosen^2))
# Add cubic term
m3 = update(m2,~.+I(NormDistanceChosen^3))
# Compare fit of models
anova(m0,m1,m2,m3)
## Data: d
## Models:
## m0: cbind(totalNFreq, totalEFreq) ~ 1 + (1 | Set) + (1 | EngLexeme)
## m1: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) + NormDistanceChosen
## m2: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) + NormDistanceChosen + I(NormDist
## m3: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) + NormDistanceChosen + I(NormDist
             AIC
                  BIC logLik deviance
                                          Chisq Df Pr(>Chisq)
     npar
         3 32132 32149 -16063
## mO
                                 32126
## m1
         4 32134 32156 -16063
                                 32126
                                         0.0208
                                                1
                                                       0.8854
## m2
        5 32078 32105 -16034
                                 32068 58.4612
                                                 1 2.073e-14 ***
## m3
         6 31842 31875 -15915
                                 31830 237.7808 1 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Summary of final model:
summary(m3)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial ( logit )
## Formula: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) +
       NormDistanceChosen + I(NormDistanceChosen^2) + I(NormDistanceChosen^3)
##
##
      Data: d
##
                       logLik deviance df.resid
##
                 BIC
##
   31842.1 31874.5 -15915.1 31830.1
                                           1630
##
## Scaled residuals:
       Min
                  1Q
                      Median
                                    3Q
                                            Max
## -13.5685 -2.5875 -0.6287
                                0.7102
                                        24.4508
##
## Random effects:
                          Variance Std.Dev.
## Groups
             Name
## EngLexeme (Intercept) 1.998
                                   1.413
```

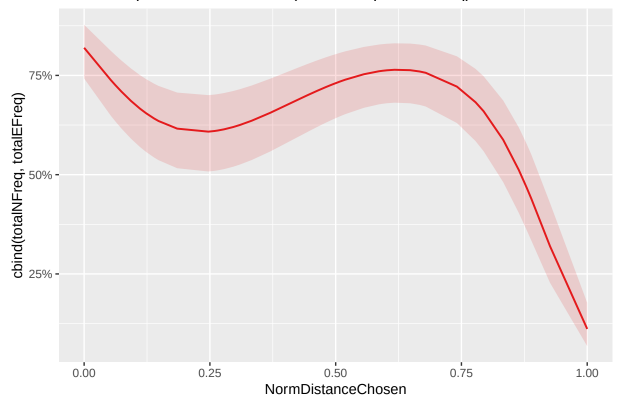
1.208

(Intercept) 1.458

Set

```
## Number of obs: 1636, groups: EngLexeme, 135; Set, 67
##
## Fixed effects:
##
                           Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                             1.5135
                                        0.2325
                                                 6.509 7.57e-11 ***
## NormDistanceChosen
                           -10.5625
                                        0.9471 -11.153
                                                        < 2e-16 ***
## I(NormDistanceChosen^2)
                                        2.2254 13.996
                          31.1462
                                                        < 2e-16 ***
## I(NormDistanceChosen^3) -24.1724
                                        1.5321 -15.778
                                                        < 2e-16 ***
##
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
##
## Correlation of Fixed Effects:
##
               (Intr) NrmDsC I(NDC^2
## NrmDstncChs -0.435
## I(NrmDsC^2)
               0.388 -0.966
## I(NrmDsC^3) -0.346 0.906 -0.982
pSimpleTotal = plot_model(m3,'eff',
  terms="NormDistanceChosen[all]")
pSimpleTotal
```

Predicted probabilities of cbind(totalNFreq, totalEFreq)



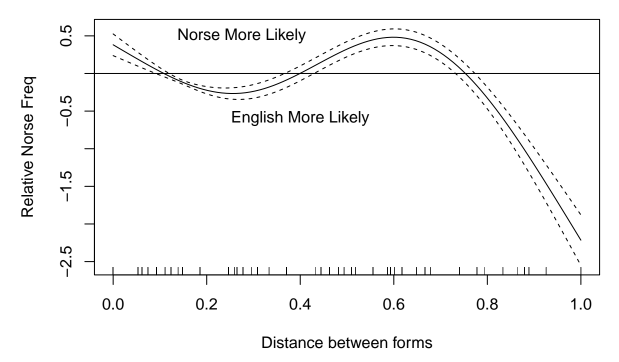
Marginal represents the variance explained by the fixed effects. Conditional represents the variance explained by the entire model.

```
## Warning: 'r.squaredGLMM' now calculates a revised statistic. See the help page.
## Warning: the null model is correct only if all variables used by the original
## model remain unchanged.
## R2m R2c
## theoretical 0.04500701 0.9783670
## delta 0.04474303 0.9726285
```

Statistics: (Linear: beta = -10.563, z = -11.153, Wald p < 0.001, LLDiff = 0, df = 1, p = 0.885; Quadratic: beta = 31.146, z = 13.996, Wald p < 0.001, LLDiff = 29.2, df = 1, p < 0.001; Cubic: beta = -24.172, z = -15.778, Wald p < 0.001, LLDiff = 118.9, df = 1, p < 0.001)

Same analysis using GAM with binomial distribution:

```
mGAM = gam(cbind(totalNFreq,totalEFreq) ~
            s(NormDistanceChosen, k=4) +
            s(Set.bs="re") +
            s(EngLexeme, bs="re"),
           data = d, family = "binomial")
summary(mGAM)
##
## Family: binomial
## Link function: logit
##
## Formula:
## cbind(totalNFreq, totalEFreq) ~ s(NormDistanceChosen, k = 4) +
      s(Set, bs = "re") + s(EngLexeme, bs = "re")
##
##
## Parametric coefficients:
              Estimate Std. Error z value Pr(>|z|)
                           0.2287 2.937 0.00331 **
## (Intercept)
                0.6717
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Approximate significance of smooth terms:
                           edf Ref.df
                                         Chi.sq p-value
## s(NormDistanceChosen) 2.997
                                   3
                                          253.3 <2e-16 ***
## s(Set)
                        44.342
                                   66 1327777.6 0.369
## s(EngLexeme)
                        81.735
                                  134 354268.6
                                                  0.521
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) = 0.44 Deviance explained = 46.1\%
## UBRE = 16.052 Scale est. = 1
                                        n = 1636
plot.gam(mGAM,
        ylab="Relative Norse Freq",
        xlab="Distance between forms",
        select = 1)
abline(h=0)
text(0.275,0.5,"Norse More Likely")
text(0.4,-0.6,"English More Likely")
```



The raw frequency of Norse forms is negatively correlated with distance:

d\$totalAllFreq = d\$totalNFreq + d\$totalEFreq

Chain 1: Iteration:

Chain 1: Iteration:

Chain 1: Iteration:

Chain 1: Iteration: 1000 / 2000 [50%]

Chain 1: Iteration: 1001 / 2000 [50%]

Chain 1: Iteration: 1200 / 2000 [60%]

Chain 1: Iteration: 1400 / 2000 [70%]

The distribution of Norse proportions is bimodal, with most terms having either low or high proportions. This isn't ideal for the binomial models above, so we also test the same model using a beta binomial distribution, which can fit bimodal binomial distributions. The result is very similar to the others:

```
mBinomBeta = brm(totalNFreq | trials(totalAllFreq) ~
               1 + NormDistanceChosen +
               I(NormDistanceChosen^2) +
               I(NormDistanceChosen^3) +
             (1|Set) + (1|EngLexeme),
           data = d, family = "beta_binomial")
## Compiling Stan program...
## Start sampling
##
## SAMPLING FOR MODEL '7404dd55f235b75d82269cea3d068515' NOW (CHAIN 1).
## Chain 1:
## Chain 1: Gradient evaluation took 0.000564 seconds
## Chain 1: 1000 transitions using 10 leapfrog steps per transition would take 5.64 seconds.
## Chain 1: Adjust your expectations accordingly!
## Chain 1:
## Chain 1:
## Chain 1: Iteration:
                                            (Warmup)
                          1 / 2000 [ 0%]
## Chain 1: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
```

(Warmup)

(Warmup)

(Warmup)

(Warmup)

(Sampling)

(Sampling)

(Sampling)

400 / 2000 [20%]

600 / 2000 [30%]

800 / 2000 [40%]

```
## Chain 1: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 1: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 1: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 1:
## Chain 1: Elapsed Time: 70.7199 seconds (Warm-up)
## Chain 1:
                           64.0899 seconds (Sampling)
## Chain 1:
                           134.81 seconds (Total)
## Chain 1:
##
## SAMPLING FOR MODEL '7404dd55f235b75d82269cea3d068515' NOW (CHAIN 2).
## Chain 2:
## Chain 2: Gradient evaluation took 0.000462 seconds
## Chain 2: 1000 transitions using 10 leapfrog steps per transition would take 4.62 seconds.
## Chain 2: Adjust your expectations accordingly!
## Chain 2:
## Chain 2:
## Chain 2: Iteration:
                          1 / 2000 [ 0%]
                                            (Warmup)
## Chain 2: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 2: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
## Chain 2: Iteration: 600 / 2000 [ 30%]
                                            (Warmup)
## Chain 2: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 2: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 2: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 2: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 2: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 2: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 2: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 2: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 2:
## Chain 2:
            Elapsed Time: 73.5453 seconds (Warm-up)
## Chain 2:
                           68.7505 seconds (Sampling)
## Chain 2:
                           142.296 seconds (Total)
## Chain 2:
##
## SAMPLING FOR MODEL '7404dd55f235b75d82269cea3d068515' NOW (CHAIN 3).
## Chain 3:
## Chain 3: Gradient evaluation took 0.000463 seconds
## Chain 3: 1000 transitions using 10 leapfrog steps per transition would take 4.63 seconds.
## Chain 3: Adjust your expectations accordingly!
## Chain 3:
## Chain 3:
## Chain 3: Iteration:
                          1 / 2000 [ 0%]
                                            (Warmup)
## Chain 3: Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 3: Iteration: 400 / 2000 [ 20%]
                                            (Warmup)
## Chain 3: Iteration: 600 / 2000 [ 30%]
                                            (Warmup)
## Chain 3: Iteration: 800 / 2000 [ 40%]
                                            (Warmup)
## Chain 3: Iteration: 1000 / 2000 [ 50%]
                                            (Warmup)
## Chain 3: Iteration: 1001 / 2000 [ 50%]
                                            (Sampling)
## Chain 3: Iteration: 1200 / 2000 [ 60%]
                                            (Sampling)
## Chain 3: Iteration: 1400 / 2000 [ 70%]
                                            (Sampling)
## Chain 3: Iteration: 1600 / 2000 [ 80%]
                                            (Sampling)
## Chain 3: Iteration: 1800 / 2000 [ 90%]
                                            (Sampling)
## Chain 3: Iteration: 2000 / 2000 [100%]
                                            (Sampling)
## Chain 3:
## Chain 3: Elapsed Time: 66.5127 seconds (Warm-up)
## Chain 3:
                           62.0208 seconds (Sampling)
## Chain 3:
                           128.533 seconds (Total)
## Chain 3:
```

```
## SAMPLING FOR MODEL '7404dd55f235b75d82269cea3d068515' NOW (CHAIN 4).
## Chain 4:
## Chain 4: Gradient evaluation took 0.000457 seconds
## Chain 4: 1000 transitions using 10 leapfrog steps per transition would take 4.57 seconds.
## Chain 4: Adjust your expectations accordingly!
## Chain 4:
## Chain 4:
## Chain 4: Iteration:
                          1 / 2000 [ 0%]
                                           (Warmup)
## Chain 4: Iteration: 200 / 2000 [ 10%]
                                           (Warmup)
## Chain 4: Iteration: 400 / 2000 [ 20%]
                                           (Warmup)
## Chain 4: Iteration: 600 / 2000 [ 30%]
                                           (Warmup)
## Chain 4: Iteration: 800 / 2000 [ 40%]
                                           (Warmup)
## Chain 4: Iteration: 1000 / 2000 [ 50%]
                                           (Warmup)
## Chain 4: Iteration: 1001 / 2000 [ 50%]
                                           (Sampling)
## Chain 4: Iteration: 1200 / 2000 [ 60%]
                                           (Sampling)
## Chain 4: Iteration: 1400 / 2000 [ 70%]
                                           (Sampling)
## Chain 4: Iteration: 1600 / 2000 [ 80%]
                                           (Sampling)
## Chain 4: Iteration: 1800 / 2000 [ 90%]
                                           (Sampling)
## Chain 4: Iteration: 2000 / 2000 [100%]
                                           (Sampling)
## Chain 4:
## Chain 4: Elapsed Time: 73.2741 seconds (Warm-up)
## Chain 4:
                           57.0322 seconds (Sampling)
## Chain 4:
                           130.306 seconds (Total)
## Chain 4:
## Warning: Bulk Effective Samples Size (ESS) is too low, indicating posterior means and medians may
## Running the chains for more iterations may help. See
## https://mc-stan.org/misc/warnings.html#bulk-ess
## Warning: Tail Effective Samples Size (ESS) is too low, indicating posterior variances and tail qu
## Running the chains for more iterations may help. See
## https://mc-stan.org/misc/warnings.html#tail-ess
summary(mBinomBeta)
## Family: beta_binomial
##
    Links: mu = logit; phi = identity
## Formula: totalNFreq | trials(totalAllFreq) ~ 1 + NormDistanceChosen + I(NormDistanceChosen^2) + I
     Data: d (Number of observations: 1638)
     Draws: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
           total post-warmup draws = 4000
##
## Group-Level Effects:
## ~EngLexeme (Number of levels: 135)
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
                               0.09
## sd(Intercept)
                     0.69
                                        0.53
                                                 0.87 1.01
                                                                 410
                                                                          968
## ~Set (Number of levels: 67)
##
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sd(Intercept)
                     0.33
                               0.15
                                        0.02
                                                 0.60 1.02
                                                                 177
                                                                          294
##
## Population-Level Effects:
                         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS
##
## Intercept
                             0.69
                                      0.29
                                                0.12
                                                         1.26 1.01
## NormDistanceChosen
                            -4.72
                                       2.45
                                               -9.58
                                                        -0.00 1.01
                                                                         855
## INormDistanceChosenE2
                            13.32
                                                2.49
                                                        24.34 1.00
                                                                         900
                                       5.61
                                             -17.88
                                                        -3.21 1.00
                                                                         978
## INormDistanceChosenE3
                           -10.53
                                       3.76
##
                         Tail_ESS
## Intercept
                             1597
```

```
## NormDistanceChosen
                              1410
## INormDistanceChosenE2
                              1399
## INormDistanceChosenE3
                              1595
## Family Specific Parameters:
##
       Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
                     0.08
                               1.85
                                        2.16 1.00
                                                       3460
## phi
##
## Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
pred = ggpredict(mBinomBeta,terms="NormDistanceChosen[all]")
## Note: uncertainty of error terms are not taken into account. Consider
     setting `interval` to "prediction". This will call `posterior_predict()`
     instead of `posterior_epred()`.
## Warning in checkMatrixPackageVersion(): Package version inconsistency detected.
## TMB was built with Matrix version 1.6.1
## Current Matrix version is 1.6.5
## Please re-install 'TMB' from source using install.packages('TMB', type = 'source') or ask CRAN fo
ggplot(pred,aes(x=x,y=predicted,ymin=conf.low,ymax=conf.high)) +
  geom_ribbon(alpha=0.3)+
  geom_line() +
  xlab("Distance") +
  ylab("Predicted Likelihood")
  8 -
  6 -
Predicted Likelihood
  2 -
```

0.50

Distance

0.75

1.00

0.25

0.00

5.2 Feature-based distance (total frequency)

We use nested model comparison to evaluate whether higher-order variables should be added (i.e. how non-linear the relationships is).

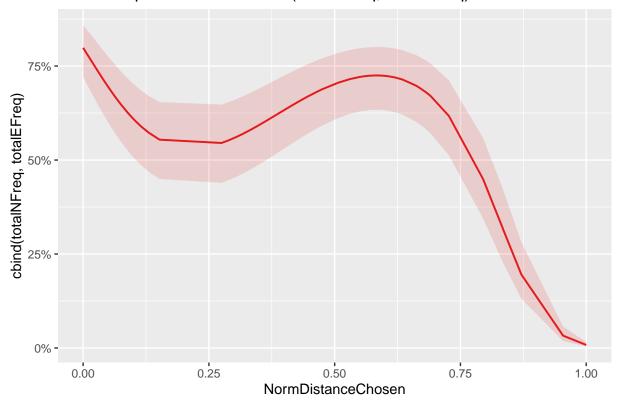
```
d$NormDistanceChosen = d$NormFeatureDistance
# Baseline model
m0 = glmer(cbind(totalNFreq,totalEFreq) ~
             1 + (1|Set) + (1|EngLexeme),
           data = d, family = "binomial")
# Add the distance measure
m1 = update(m0,~.+NormDistanceChosen)
# Add quadratic term
m2 = update(m1,~.+I(NormDistanceChosen^2))
# Add cubic term
m3 = update(m2,~.+I(NormDistanceChosen^3))
# Compare fit of models
anova(m0,m1,m2,m3)
## Data: d
## Models:
## m0: cbind(totalNFreq, totalEFreq) ~ 1 + (1 | Set) + (1 | EngLexeme)
## m1: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) + NormDistanceChosen
## m2: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) + NormDistanceChosen + I(NormDist
## m3: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) + NormDistanceChosen + I(NormDist
                  BIC logLik deviance
##
     npar
            AIC
                                        Chisq Df Pr(>Chisq)
## mO
        3 32262 32278 -16128
                                 32256
        4 32247 32269 -16120
## m1
                                32239
                                       16.464 1 4.958e-05 ***
        5 32159 32186 -16074
                                32149 90.456 1 < 2.2e-16 ***
## m2
## m3
        6 31856 31888 -15922
                                31844 305.264 1 < 2.2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Summary of final model:
summary(m3)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) +
##
      NormDistanceChosen + I(NormDistanceChosen^2) + I(NormDistanceChosen^3)
##
      Data: d
##
                BIC logLik deviance df.resid
       AIC
## 31855.8 31888.2 -15921.9 31843.8
##
## Scaled residuals:
##
       Min
                 1Q
                      Median
                                    3Q
## -13.5687 -2.5875 -0.6362
                               0.7073
                                       24.4503
##
## Random effects:
  Groups
             Name
                          Variance Std.Dev.
## EngLexeme (Intercept) 2.002
                                  1.415
             (Intercept) 1.551
                                   1.245
## Number of obs: 1638, groups: EngLexeme, 135; Set, 67
## Fixed effects:
                           Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                            1.3771
                                       0.2192 6.282 3.33e-10 ***
```

0.9665 -13.580 < 2e-16 ***

-13.1251

NormDistanceChosen

Predicted probabilities of cbind(totalNFreq, totalEFreq)

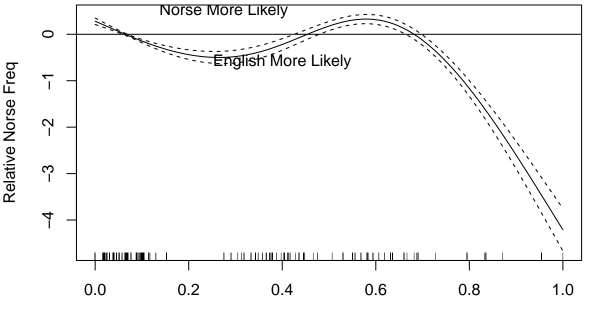


Statistics: (Linear: beta = -13.125, z = -13.58, Wald p < 0.001, LLDiff = 8.2, df = 1, p < 0.001; Quadratic: beta = 41.402, z = 15.854, Wald p < 0.001, LLDiff = 45.2, df = 1, p < 0.001; Cubic: beta = -34.456, z = -17.797, Wald p < 0.001, LLDiff = 152.6, df = 1, p < 0.001)

Same analysis using GAM with binomial distribution:

```
##
## Family: binomial
## Link function: logit
##
## Formula:
## cbind(totalNFreq, totalEFreq) ~ s(NormDistanceChosen, k = 4) +
```

```
##
      s(Set, bs = "re") + s(EngLexeme, bs = "re")
##
## Parametric coefficients:
              Estimate Std. Error z value Pr(>|z|)
                0.6916
                           0.2334
                                    2.964 0.00304 **
## (Intercept)
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Approximate significance of smooth terms:
##
                           edf Ref.df
                                         Chi.sq p-value
## s(NormDistanceChosen) 2.999
                                          348.8 <2e-16 ***
                                    3
## s(Set)
                        45.288
                                   66 1224049.3
                                                 0.399
## s(EngLexeme)
                        80.832
                                  134 344925.2
                                                  0.644
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) =
                 0.44 Deviance explained = 46.1%
## UBRE = 16.045 Scale est. = 1
                                        n = 1638
plot.gam(mGAM,
        ylab="Relative Norse Freq",
        xlab="Distance between forms",
        select = 1)
abline(h=0)
text(0.275,0.5,"Norse More Likely")
text(0.4,-0.6,"English More Likely")
```



Distance between forms

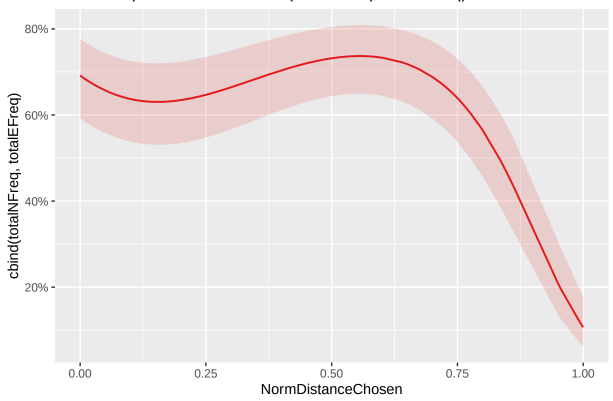
5.3 Historical distance (total frequency)

We use nested model comparison to evaluate whether higher-order variables should be added (i.e. how non-linear the relationships is).

```
d$NormDistanceChosen = d$NormHistoricalDistance
# Baseline model
m0 = glmer(cbind(totalNFreq,totalEFreq) ~
             1 + (1|Set) + (1|EngLexeme),
           data = d, family = "binomial")
# Add the distance measure
m1 = update(m0,~.+NormDistanceChosen)
# Add quadratic term
m2 = update(m1,~.+I(NormDistanceChosen^2))
# Add cubic term
m3 = update(m2,~.+I(NormDistanceChosen^3))
# Compare fit of models
anova(m0,m1,m2,m3)
## Data: d
## Models:
## m0: cbind(totalNFreq, totalEFreq) ~ 1 + (1 | Set) + (1 | EngLexeme)
## m1: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) + NormDistanceChosen
## m2: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) + NormDistanceChosen + I(NormDist
## m3: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) + NormDistanceChosen + I(NormDist
     npar
                  BIC logLik deviance
##
            AIC
                                          Chisq Df Pr(>Chisq)
## mO
        3 32262 32278 -16128
                                 32256
## m1
        4 32263 32284 -16127
                                 32255
                                         1.3764 1
                                                       0.2407
        5 32146 32173 -16068
                                 32136 118.7880
## m2
                                                 1
                                                       <2e-16 ***
## m3
        6 32080 32112 -16034
                                 32068 67.9115 1
                                                       <2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Summary of final model:
summary(m3)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
  Family: binomial (logit)
## Formula: cbind(totalNFreq, totalEFreq) ~ (1 | Set) + (1 | EngLexeme) +
##
      NormDistanceChosen + I(NormDistanceChosen^2) + I(NormDistanceChosen^3)
##
      Data: d
##
                BIC logLik deviance df.resid
       AIC
   32079.9 32112.3 -16033.9 32067.9
##
## Scaled residuals:
##
       Min
                  1Q
                      Median
                                    3Q
## -13.6467 -2.5893 -0.6312
                                0.7375
                                       24.4681
##
## Random effects:
  Groups
             Name
                          Variance Std.Dev.
## EngLexeme (Intercept) 1.976
                                   1.406
              (Intercept) 1.486
                                   1.219
## Number of obs: 1638, groups: EngLexeme, 135; Set, 67
## Fixed effects:
                           Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                             0.8076
                                       0.2234 3.616 0.000299 ***
## NormDistanceChosen
                            -3.9175
                                        0.9838 -3.982 6.83e-05 ***
```

```
## I(NormDistanceChosen^2) 16.2533
                                       2.5269
                                              6.432 1.26e-10 ***
## I(NormDistanceChosen^3) -15.2648
                                       1.7883 -8.536 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
               (Intr) NrmDsC I(NDC^2
##
## NrmDstncChs -0.350
## I(NrmDsC^2) 0.320 -0.978
## I(NrmDsC^3) -0.296 0.934 -0.985
pHistoricalTotal = plot_model(m3,'eff',
 terms="NormDistanceChosen[all]")
pHistoricalTotal
```

Predicted probabilities of cbind(totalNFreq, totalEFreq)



Statistics: (Linear: beta = -3.918, z = -3.982, Wald p < 0.001, LLDiff = 0.7, df = 1, p = 0.241; Quadratic: beta = 16.253, z = 6.432, Wald p < 0.001, LLDiff = 59.4, df = 1, p < 0.001; Cubic: beta = -15.265, z = -8.536, Wald p < 0.001, LLDiff = 34, df = 1, p < 0.001)

Same analysis using GAM with binomial distribution:

```
##
## Family: binomial
## Link function: logit
##
## Formula:
## cbind(totalNFreq, totalEFreq) ~ s(NormDistanceChosen, k = 4) +
```

```
##
      s(Set, bs = "re") + s(EngLexeme, bs = "re")
##
## Parametric coefficients:
              Estimate Std. Error z value Pr(>|z|)
                0.5980
                          0.2378
                                   2.514
## (Intercept)
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Approximate significance of smooth terms:
##
                           edf Ref.df
                                      Chi.sq p-value
## s(NormDistanceChosen) 2.998
                                  3
                                        163.6 <2e-16 ***
## s(Set)
                        45.968
                                  66 698441.1 0.5322
## s(EngLexeme)
                        80.247
                                 134 698557.1 0.0652 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) = 0.435 Deviance explained = 45.8%
## UBRE = 16.164 Scale est. = 1
                                       n = 1638
plot.gam(mGAM,
        ylab="Relative Norse Freq",
        xlab="Distance between forms",
        select = 1)
abline(h=0)
text(0.275,0.5,"Norse More Likely")
text(0.4,-0.6,"English More Likely")
     2
                     Norse More Likely
     0
Relative Norse Freq
     S
                             English More Likely
     -1.5
            0.0
                        0.2
                                     0.4
                                                  0.6
                                                               0.8
                                                                            1.0
                                 Distance between forms
```

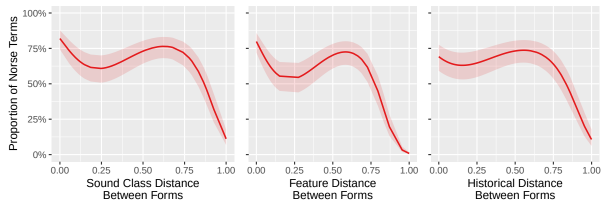
Summary of results for total frequency analyses:

```
bigPlot = grid.arrange(
             pSimpleTotal +
               ggtitle("Total Frequency Analysis") +
               coord_cartesian(ylim = c(0,1))+
               xlab("Sound Class Distance\nBetween Forms")+
               ylab("Proportion of Norse Terms"),
             pFeatureTotal+
               ggtitle("") +
               theme(axis.title.y = element_blank(),
                     axis.text.y=element_blank())+
               coord_cartesian(ylim = c(0,1))+
               xlab("Feature Distance\nBetween Forms"),
             pHistoricalTotal+
               ggtitle("") +
               theme(axis.title.y = element_blank(),
                     axis.text.y=element_blank())+
               coord_cartesian(ylim = c(0,1))+
               xlab("Historical Distance\nBetween Forms"),
             nrow=1,widths=c(1.3,1,1))
```

```
pdf("../results/BigEffectsPlot_totalFreq.pdf",width = 8,height=3)
    plot(bigPlot)
dev.off()
```

pdf ## 2

Total Frequency Analysis



5.4 Exploratory analyses

5.4.1 Word Class

It is possible that lexical choices differ by word class. For example, verbs may be more resistant to integration. We test this by adding word class as a predictor. Since some word forms appear as several classes, we treat word class as a series of independent binary variables indicating the possibility or not of the word appearing as this class.

```
classes = unique(trimws(unlist(strsplit(unique(d$Class),"/"))))
classes = classes[!is.na(classes)]
classNames = paste0("Class.",gsub(" ",".",classes))
for(i in 1:length(classes)){
 d[,classNames[i]] = grepl(paste0("^",classes[i]),d$Class)
```

Build a null model with non-linear distance, as above:

Approximation) [glmerMod] ## Family: binomial (logit)

```
d$NormDistanceChosen = d$NormDistance
mC0 = glmer(cbind(totalNFreq,totalEFreq) ~
             1 + NormDistanceChosen +
               I(NormDistanceChosen^2) +
               I(NormDistanceChosen^3) +
             (1|Set) + (1|EngLexeme),
           data = d, family = "binomial")
```

```
Adding the verb class variable does not improve the fit of the model:
mC1a = update(mC0,~.+Class.verb)
anova(mCO,mC1a)
## Data: d
## Models:
## mCO: cbind(totalNFreq, totalEFreq) ~ 1 + NormDistanceChosen + I(NormDistanceChosen^2) + I(NormDis
## mC1a: cbind(totalNFreq, totalEFreq) ~ NormDistanceChosen + I(NormDistanceChosen^2) + I(NormDistan
                     BIC logLik deviance Chisq Df Pr(>Chisq)
               AIC
           6 31975 32007 -15981
                                    31963
## mCO
           7 31977 32014 -15981
## mC1a
                                    31963 0.0928 1
                                                        0.7607
Adding all other classes does not improve the fit of the model:
mC1b = update(mC0,~.+Class.adverb + Class.preposition + Class.noun +
               Class.numeral + Class.adjective + Class.verb +
               Class.indefinite.pronoun + Class.interjection)
## fixed-effect model matrix is rank deficient so dropping 2 columns / coefficients
## Warning in checkConv(attr(opt, "derivs"), opt$par, ctrl = control$checkConv, :
## Model failed to converge with max|grad| = 0.00689701 (tol = 0.002, component 1)
anova(mCO,mC1b)
## Data: d
## Models:
## mCO: cbind(totalNFreq, totalEFreq) ~ 1 + NormDistanceChosen + I(NormDistanceChosen^2) + I(NormDis
## mC1b: cbind(totalNFreq, totalEFreq) ~ NormDistanceChosen + I(NormDistanceChosen^2) + I(NormDistan
##
                     BIC logLik deviance Chisq Df Pr(>Chisq)
           6 31975 32007 -15981
## mCO
                                    31963
          12 31980 32044 -15978
                                    31956 7.0668 6
## mC1b
                                                        0.3147
summary(mC1b)
## Generalized linear mixed model fit by maximum likelihood (Laplace
```

```
## Formula:
## cbind(totalNFreq, totalEFreq) ~ NormDistanceChosen + I(NormDistanceChosen^2) +
      I(NormDistanceChosen^3) + (1 | Set) + (1 | EngLexeme) + Class.adverb +
      Class.preposition + Class.noun + Class.numeral + Class.adjective +
      Class.verb + Class.indefinite.pronoun + Class.interjection
##
     Data: d
##
##
                BIC
                      logLik deviance df.resid
##
   31979.6 32044.4 -15977.8 31955.6
##
## Scaled residuals:
                     Median
       Min 1Q
## -13.6715 -2.5875 -0.6247
                               0.7114 24.4498
##
## Random effects:
## Groups
            Name
                         Variance Std.Dev.
## EngLexeme (Intercept) 1.992
                                  1.411
             (Intercept) 1.178
                                  1.085
## Number of obs: 1638, groups: EngLexeme, 135; Set, 67
## Fixed effects:
                               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                                1.2177
                                        0.7815 1.558 0.119
## NormDistanceChosen
                               -10.4432
                                           0.9922 -10.526
                                                            <2e-16 ***
## I(NormDistanceChosen^2)
                                30.8080
                                           2.3382 13.176
                                                            <2e-16 ***
## I(NormDistanceChosen^3)
                               -23.9496
                                           1.6074 -14.899
                                                            <2e-16 ***
## Class.adverbTRUE
                                0.9804
                                           0.9147
                                                   1.072
                                                             0.284
## Class.nounTRUE
                                 0.1828
                                           0.8329
                                                   0.220
                                                            0.826
## Class.numeralTRUE
                                                   1.538
                                                           0.124
                                 1.8335
                                           1.1919
## Class.adjectiveTRUE
                                -0.4274
                                           0.9183 - 0.465
                                                             0.642
## Class.verbTRUE
                                 0.1505
                                           0.8427
                                                    0.179
                                                             0.858
## Class.indefinite.pronounTRUE
                                1.0948
                                           1.6814
                                                   0.651
                                                           0.515
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
              (Intr) NrmDsC I(NDC^2 I(NDC^3 Clss.dvTRUE Clss.nnTRUE Clss.nmTRUE
##
## NrmDstncChs -0.129
## I(NrmDsC^2) 0.114 -0.969
## I(NrmDsC^3) -0.101 0.915 -0.983
## Clss.dvTRUE -0.833 -0.013 0.010 -0.008
## Clss.nnTRUE -0.917 -0.005 0.008 -0.009
                                            0.781
## Clss.nmTRUE -0.636 0.009 -0.008
                                    0.005 0.541
                                                        0.594
## Clss.djTRUE -0.831 -0.006 0.005 -0.005
                                            0.708
                                                        0.785
                                                                    0.539
## Clss.vrTRUE -0.861 -0.004 0.007 -0.007
                                             0.732
                                                        0.805
                                                                    0.557
## Clss.n.TRUE -0.454 -0.014 0.016 -0.019 0.388
                                                        0.427
                                                                    0.296
              Clss.djTRUE Clss.vTRUE
## NrmDstncChs
## I(NrmDsC^2)
## I(NrmDsC^3)
## Clss.dvTRUE
## Clss.nnTRUE
## Clss.nmTRUE
## Clss.djTRUE
## Clss.vrTRUE 0.729
                           0.400
## Clss.n.TRUE 0.387
## fit warnings:
## fixed-effect model matrix is rank deficient so dropping 2 columns / coefficients
```

```
## optimizer (Nelder_Mead) convergence code: 0 (OK)
## Model failed to converge with max|grad| = 0.00689701 (tol = 0.002, component 1)
```

5.4.2 Diagnostic consonants

Some Norse words have more salient diagnostic segments - they sound 'more Norse'. For example, the presence /sk/ vs post-alveolar fricative. Below, we create a variable that distinguishes pairs where the Norse includes /sk/ (and the English includes / /) from other cases, and enter this as a variable into the model to test whether it improves the prediction of the proportion of Norse usage.

```
d$NorseDiagnosticSK =
 grepl("sk",d$NorseForm) &
 grepl(" ",d$EngForm) &
 !grepl("sk",d$EngForm)
# (this isn't valid for all possible words,
# but does identify the relevant data in
# our sample)
d$NorseDiagnosticSK = factor(d$NorseDiagnosticSK)
d$NormDistanceChosen = d$NormDistance
mD0 = glmer(cbind(totalNFreq,totalEFreq) ~
             1 + NormDistanceChosen +
               I(NormDistanceChosen^2) +
               I(NormDistanceChosen^3) +
             (1|Set) + (1|EngLexeme),
           data = d, family = "binomial")
mD1 = update(mD0, ~.+NorseDiagnosticSK)
anova(mD0,mD1)
## Data: d
## Models:
## mD0: cbind(totalNFreq, totalEFreq) ~ 1 + NormDistanceChosen + I(NormDistanceChosen^2) + I(NormDis
## mD1: cbind(totalNFreq, totalEFreq) ~ NormDistanceChosen + I(NormDistanceChosen^2) + I(NormDistanceChosen^2)
              AIC BIC logLik deviance Chisq Df Pr(>Chisq)
##
      npar
## mDO
         6 31975 32007 -15981
                                  31963
## mD1
          7 31966 32004 -15976
                                  31952 10.236 1 0.001378 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
summary(mD1)
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
## Formula:
## cbind(totalNFreq, totalEFreq) ~ NormDistanceChosen + I(NormDistanceChosen^2) +
       I(NormDistanceChosen^3) + (1 | Set) + (1 | EngLexeme) + NorseDiagnosticSK
##
##
      Data: d
##
##
        ATC:
                 BIC
                       logLik deviance df.resid
   31966.4 32004.2 -15976.2 31952.4
##
                                           1631
##
## Scaled residuals:
##
       Min
                 1Q
                      Median
                                    30
                                            Max
## -13.6634 -2.5875 -0.6253
                                0.7084 24.4509
##
## Random effects:
## Groups Name
                         Variance Std.Dev.
## EngLexeme (Intercept) 2.041
```

```
## Set
              (Intercept) 1.563
                                    1.250
## Number of obs: 1638, groups: EngLexeme, 135; Set, 67
## Fixed effects:
                            Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                              1.5681
                                         0.2365
                                                 6.630 3.37e-11 ***
                                         0.9237 -11.401 < 2e-16 ***
## NormDistanceChosen
                            -10.5306
## I(NormDistanceChosen^2) 31.1074
                                         2.1647
                                                14.370
                                                         < 2e-16 ***
## I(NormDistanceChosen^3) -24.1643
                                         1.4907 -16.210 < 2e-16 ***
## NorseDiagnosticSKTRUE
                             -2.0690
                                         0.6480 -3.193 0.00141 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
               (Intr) NrmDsC I(NDC^2 I(NDC^3
##
## NrmDstncChs -0.421
## I(NrmDsC^2) 0.374 -0.965
## I(NrmDsC^3) -0.333 0.901 -0.981
## NrsDgSKTRUE -0.090 0.030 -0.040
plot_model(mD1,'eff', terms="NorseDiagnosticSK") +
  ylab("Likelihood of choosing Norse term") +
  xlab("Presence of /sk/") +
  ggtitle("")
  60%
ikelihood of choosing Norse term
  40%
  20%
           FALSE
                                                                                TRUE
                                        Presence of /sk/
get_model_data(mD1,'eff', terms="NorseDiagnosticSK")
## # Predicted probabilities of cbind(totalNFreq, totalEFreq)
##
                                        95% CI | group_col
## NorseDiagnosticSK | Predicted |
##
                   1 |
                             0.62 | 0.52, 0.71 |
                             0.17 | 0.05, 0.43 |
##
                   2 |
```

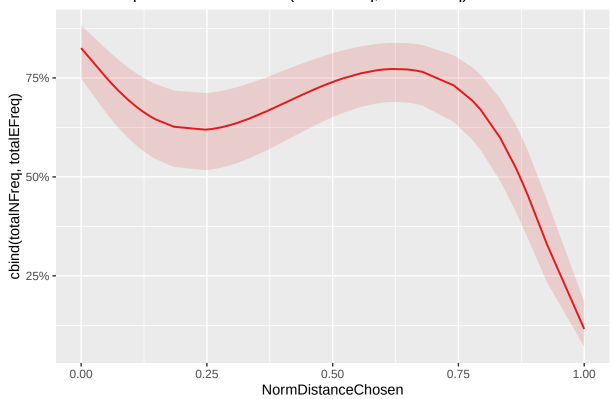
Adding the Norse Diagnostic Score to the model significantly improves the fit of the model. The model

suggests that Norse forms which are more obviously Norse have a lower chance of being selected overall. This is in line with the general findings: words that are obviously more different to the English forms are less likely to be integrated.

The effect of form distance is unaffected by adding the presence of 'sk' as a variable:

```
plot_model(mD1,'eff',terms="NormDistanceChosen[all]")
```

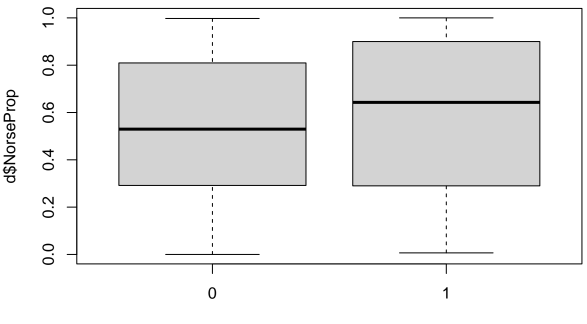
Predicted probabilities of cbind(totalNFreq, totalEFreq)



There are other consonant contrasts which are also diagnostic of Norse origin. For example, the presence of /g/vs/j/v or /k/vs post-alveolar affricate. However, these are only truly diagnostic in production when followed by a front vowel. Still, ignoring this complication for now, we calculate a Norse Diagnostic Score: cases where these segments are diagnostic of the differences between Norse and English forms (the score is potentially more than one, but in our data there are only zero and one):

```
d$NorseDiagnosticScore =
  (grepl("g",d$NorseForm) & !grepl("g",d$EngForm))+
  (grepl("",d$NorseForm) & !grepl("",d$EngForm)) +
   (grepl("sk",d$NorseForm) & !grepl("sk",d$EngForm))+
  # (k in Norse vs in English)
   (grepl("k",d$NorseForm) & grepl(" ",d$EngForm))

d$NorseDiagnosticScore = factor(d$NorseDiagnosticScore)
```



d\$NorseDiagnosticScore

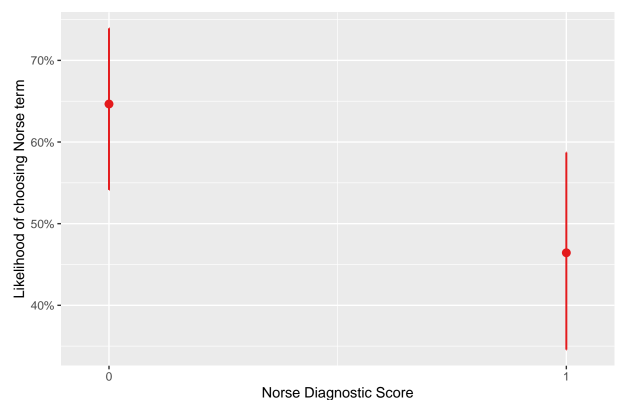
t.test(d\$NorseProp~ d\$NorseDiagnosticScore)

```
##
## Welch Two Sample t-test
##
## data: d$NorseProp by d$NorseDiagnosticScore
## t = -2.2293, df = 1532.9, p-value = 0.02594
## alternative hypothesis: true difference in means between group 0 and group 1 is not equal to 0
## 95 percent confidence interval:
## -0.065524499 -0.004187601
## sample estimates:
## mean in group 0 mean in group 1
## 0.5501452    0.5850013
```

Adding the Norse Diagnostic Score to the model significantly improves the fit of the model. The model suggests that Norse forms which are more obviously Norse have a lower chance of being selected overall.

summary(mD1)

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
## Family: binomial (logit)
## Formula:
## cbind(totalNFreq, totalEFreq) ~ NormDistanceChosen + I(NormDistanceChosen^2) +
##
       I(NormDistanceChosen^3) + (1 | Set) + (1 | EngLexeme) + NorseDiagnosticScore
##
      Data: d
##
##
       AIC
                BIC
                      logLik deviance df.resid
## 31953.8 31991.6 -15969.9 31939.8
##
## Scaled residuals:
       Min 1Q
                     Median
                                   30
## -13.6651 -2.5587 -0.6102 0.7161 24.4515
##
## Random effects:
## Groups
             Name
                         Variance Std.Dev.
## EngLexeme (Intercept) 2.024
                                  1.423
                                  1.332
## Set
             (Intercept) 1.774
## Number of obs: 1638, groups: EngLexeme, 135; Set, 67
##
## Fixed effects:
##
                          Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                            1.6604
                                       0.2476
                                              6.705 2.01e-11 ***
## NormDistanceChosen
                          -10.3816
                                       0.9876 -10.512 < 2e-16 ***
## I(NormDistanceChosen^2) 30.6307
                                       2.3291 13.151 < 2e-16 ***
## I(NormDistanceChosen^3) -23.7685
                                       1.6035 -14.823 < 2e-16 ***
## NorseDiagnosticScore1
                           -0.7469
                                       0.1583 -4.717 2.39e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
               (Intr) NrmDsC I(NDC^2 I(NDC^3
## NrmDstncChs -0.422
## I(NrmDsC<sup>2</sup>) 0.379 -0.969
## I(NrmDsC^3) -0.341 0.913 -0.983
## NrsDgnstcS1 -0.138 -0.010 0.011
                                    -0.014
plot_model(mD1,'eff', terms="NorseDiagnosticScore") +
 ylab("Likelihood of choosing Norse term") +
 xlab("Norse Diagnostic Score") +
 ggtitle("")
```



get_model_data(mD1,'eff', terms="NorseDiagnosticScore")

```
## # Predicted probabilities of cbind(totalNFreq, totalEFreq)
##
## NorseDiagnosticScore | Predicted |
                                          95% CI | group_col
##
                      0 |
                               0.65 | 0.54, 0.74 |
                                                            1
                      1 |
                               0.46 | 0.35, 0.59 |
```

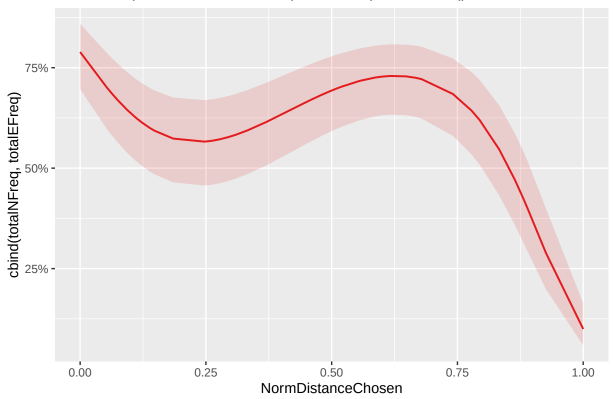
The effect of form distance is unaffected:

##

```
plot_model(mD1,'eff',terms="NormDistanceChosen[all]")
```

1

Predicted probabilities of cbind(totalNFreq, totalEFreq)



6 Source-level frequency analysis

This analysis uses source-level observations: each observation is the frequency of a pair of forms within a particular source. This also lets us control for source-level features like the source itself as a random effect, the age of the source, and whether alliteration might affect decisions.

There may be multiple orthographic forms for each unique full form for comparison, so first, we collapse the data over unique pairs. We also restructure the data to be in 'long' format.

```
d2 = data.frame()
for(i in 1:length(englishFrequencyColumns)){
 dx = data.frame(
   Set = d$Set,
   EngForm = d$EngForm,
   NorseForm = d$NorseForm,
   NormDistance = d$NormDistance,
  NormFeatureDistance = d$NormFeatureDistance,
  NormHistoricalDistance = d$NormHistoricalDistance,
  NFreq = d[,norseFrequencyColumns[i]],
  EFreq = d[,englishFrequencyColumns[i]],
   Source = englishFrequencyColumns[i],
   Alliteration = d$Alliteration)
 dx = dx[rowSums(dx[,c("NFreq","EFreq")],na.rm = T)>0,]
 uforms = paste(dx$Set,dx$EngForm,dx$NorseForm)
 dx = data.frame(
   Set = tapply(dx$Set,uforms,head,n=1),
   EngForm = tapply(dx$EngForm,uforms,head,n=1),
  NorseForm = tapply(dx$NorseForm,uforms,head,n=1),
  NormDistance = tapply(dx$NormDistance,uforms,head,n=1),
  NormFeatureDistance = tapply(dx$NormFeatureDistance,uforms,head,n=1),
  NormHistoricalDistance = tapply(dx$NormHistoricalDistance,uforms,head,n=1),
  NFreq = tapply(dx$NFreq,uforms,sum,na.rm=T),
```

```
EFreq = tapply(dx$EFreq,uforms,sum,na.rm=T),
    Source = englishFrequencyColumns[i],
    Alliteration = tapply(dx$Alliteration,uforms,head,n=1))
d2 = rbind(d2,dx)
}

d2$Set = factor(d2$Set)
d2$Source = factor(d2$Source)

# Make a variable for each unique form within a set
d2$EngForm2 = paste(d2$Set,d2$EngForm)
d2$NorseForm2 = paste(d2$Set,d2$NorseForm)
d2$EngForm2 = factor(d2$EngForm2)

# Remove any cases with zero frequency
d2 = d2[!(d2$NFreq==0 & d2$EFreq==0),]

# Proportion of norse forms
d2$NProp = 100*(d2$NFreq/(d2$NFreq+d2$EFreq))
```

Modelling age is difficult, since exact dates are not known. It would be possible to model the date of publication as an ordinal variable, for example assuming FCPC > Ormulum > {Havelok, GenAndEx} > CursorMundi > {Mannyng, GawainPoet} > WarsAlexander > StErkenwald. However, the texts from the Corpus of Middle English span several centuries, and so are hard to place in this order. Instead, we simply use the century as an ordered category.

```
AgeCategories =
    c("EFreqFCPC"=12,
      "EFreqOrmulum"=12,
     "EFreqHavelok"=13,
     "EFreqGenAndEx" = 14,
     "EFreqCursorMundi" = 14,
     "EFreqGawainPoet" = 14,
     "EFreqStErkenwald" = 15,
     "EFreqMannyng" = 15,
     "EFreqWarsAlexander" = 15,
     "EFreqLinc"=15,
       "EFreqNott"=15,
       "EFreqNorf"=15,
       "EFreqRolle"=15)
d2$Age = AgeCategories[d2$Source]
d2$Age = factor(d2$Age,ordered=T)
contrasts(d2$Age) = contr.sum(length(unique(d2$Age)))
mean(d2$NFreq/(d2$NFreq+d2$EFreq))
## [1] 0.5579405
range(d2$NFreq/(d2$NFreq+d2$EFreq))
## [1] 0 1
```

[1] 0.3594087

mean(d2\$NormDistance)

Alliteration only applies to poetry sources: Gawain, St Erkenwold, and Wars of Alexander. So turn all others to 'false':

```
d2[!d2$Source %in%
     c("EFreqGawainPoet",
       "EFreqStErkenwald",
       "EFreqWarsAlexander",
       "EFreqLinc",
         "EFreqNott",
         "EFreqNorf",
         "EFreqRolle"),]$Alliteration = FALSE
d2$Alliteration = factor(d2$Alliteration)
gPairs = ggpairs(d2[,c("NormDistance","NormFeatureDistance",
              "NormHistoricalDistance", "NProp")],
        columnLabels = c("Sound Class Distance",
                          "Feature Distance",
                          "Historical Distance",
                          "Freq (% Norse)"),
        rowLabels = c("Sound Class Distance",
                          "Feature Distance",
                          "Historical Distance",
                          "Freq (% Norse)"))
pdf("../results/GPairs.pdf", width=6, height=5.5)
 gPairs
dev.off()
## pdf
## 2
```

6.1 Simple distance

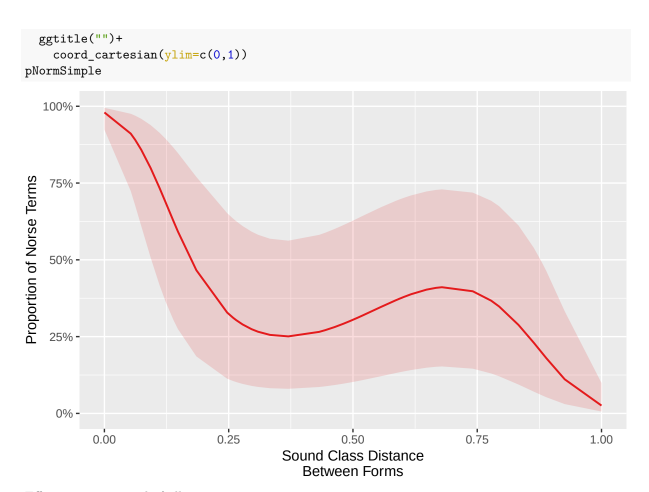
Use a mixed effects model to predict the frequency of Norse and English forms by a random effect for cognate Set and Source. We add a main effect of age of source, then introduce the normalised distance measure along with its non-linear terms.

Compare fit of models:

```
anova(m0,m1,m2,m3)
```

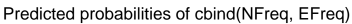
```
## Data: d2
## Models:
## m0: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2)
## m1: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2) + NormDi
## m2: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2) + NormDi
## m3: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2) + NormDi
##
            AIC BIC logLik deviance
                                       Chisq Df Pr(>Chisq)
## mO
      8 21104 21145 -10544
                                21088
## m1
        9 20936 20981 -10459
                                20918 170.622 1 < 2.2e-16 ***
## m2 10 20875 20926 -10428 20855 62.509 1 2.652e-15 ***
```

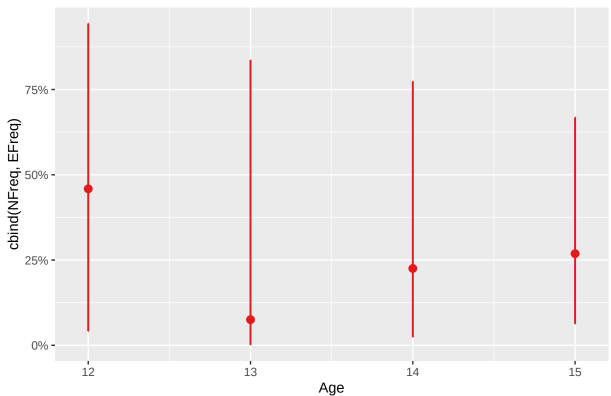
```
## m3
       11 20547 20602 -10262
                                 20525 330.399 1 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Summary of final model:
summary(m3)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) +
       (1 | EngForm2) + NormDistance + I(NormDistance^2) + I(NormDistance^3)
##
##
      Data: d2
## Control: glmerControl(optimizer = "bobyqa")
##
##
                 BIC
                       logLik deviance df.resid
##
   20546.6 20602.4 -10262.3 20524.6
                                           1162
##
## Scaled residuals:
        Min
                  1Q
                       Median
                                    3Q
                                            Max
## -105.030
                                 1.127
                                        175.531
              -1.362
                        0.114
##
## Random effects:
## Groups
            Name
                         Variance Std.Dev.
## EngForm2 (Intercept) 4.302
                                  2.074
             (Intercept) 5.354
                                  2.314
             (Intercept) 4.389
                                  2.095
## Source
## Number of obs: 1173, groups: EngForm2, 129; Set, 67; Source, 13
##
## Fixed effects:
##
                       Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                       3.554398
                                  0.834099
                                             4.261 2.03e-05 ***
                                             0.831
                                                      0.406
## Age1
                       1.062382
                                  1.278355
## Age2
                      -1.281640
                                  1.644782
                                            -0.779
                                                      0.436
## Age3
                      -0.007456
                                  1.124018
                                            -0.007
                                                      0.995
## AlliterationTRUE
                       2.233356
                                  0.071790 31.110
                                                   < 2e-16 ***
## NormDistance
                     -32.968226
                                  1.575406 -20.927
                                                    < 2e-16 ***
## I(NormDistance^2) 68.761447
                                  3.535850 19.447 < 2e-16 ***
## I(NormDistance^3) -43.320731
                                  2.343141 -18.488 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
                                           AlTRUE NrmDst I(ND^2
##
               (Intr) Age1
                             Age2
                                    Age3
## Age1
                0.010
## Age2
                0.398 - 0.521
               -0.185 -0.260 -0.490
## Age3
## AlltrtnTRUE -0.012 0.015 -0.010 0.000
## NormDistanc -0.211 -0.007 0.005 0.001 -0.026
## I(NrmDst^2) 0.189 0.007 -0.005 0.000 0.025 -0.968
## I(NrmDst^3) -0.169 -0.007 0.006 0.000 -0.026 0.908 -0.981
Norse frequency varies with distance. The plot below shows the marginal effects, holding discrete pre-
dictors constant at their proportions (not reference level):
pNormSimple =
 plot_model(m3,'eff', terms="NormDistance[all]") +
 xlab("Sound Class Distance\nBetween Forms") +
 ylab("Proportion of Norse Terms") +
```



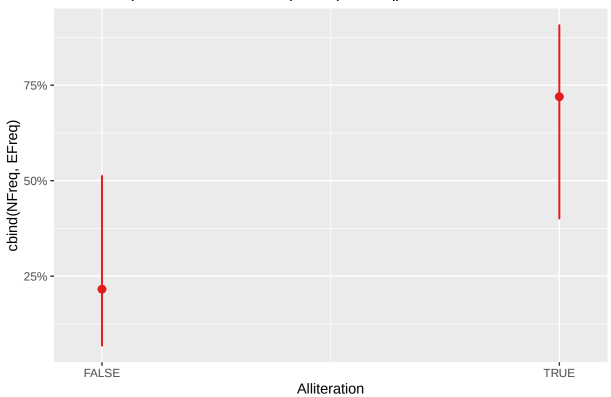
Effects over time and of alliteration:

plot_model(m3,'eff', terms="Age [all]")



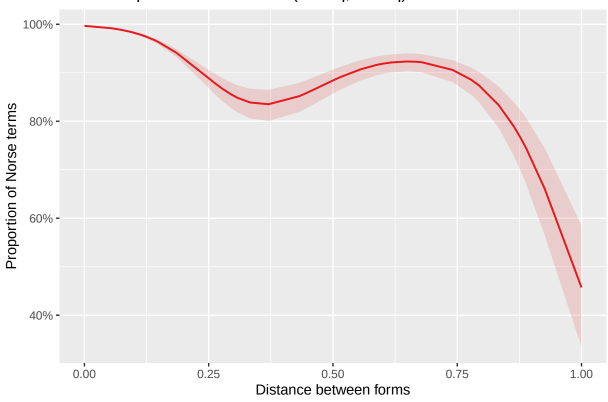


```
plot_model(m3,'eff', terms="Alliteration")
```



The GAM model seems a little different, showing a smaller effect at lower distances.

```
##
## Family: binomial
## Link function: logit
##
## Formula:
## cbind(NFreq, EFreq) ~ s(NormDistance, k = 4) + s(Age, bs = "re") +
       s(Set, bs = "re") + s(Source, bs = "re") + s(EngForm2, bs = "re")
##
##
## Parametric coefficients:
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.001498
                           2.230288 -0.001
##
## Approximate significance of smooth terms:
                        edf Ref.df
                                      Chi.sq p-value
## s(NormDistance) 2.99967
                                3 4.539e+02 < 2e-16 ***
                                3 2.027e+03 0.99821
## s(Age)
                   0.03305
## s(Set)
                                66 3.235e+07 0.10624
                   49.04776
## s(Source)
                   11.96498
                               12 1.112e+07 < 2e-16 ***
## s(EngForm2)
                   72.08085
                             128 3.721e+07 0.00747 **
```



Statistics for the simple measure:

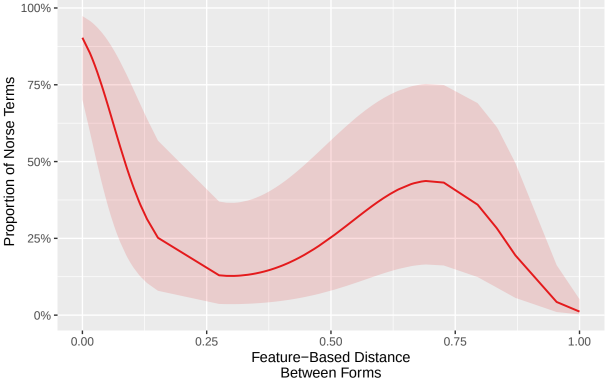
(Linear: beta = -32.968, z = -20.927, Wald p < 0.001, LLDiff = 85.3, df = 1, p < 0.001; Quadratic: beta = 68.761, z = 19.447, Wald p < 0.001, LLDiff = 31.3, df = 1, p < 0.001; Cubic: beta = -43.321, z = -18.488, Wald p < 0.001, LLDiff = 165.2, df = 1, p < 0.001)

6.2 Feature-based distance

Same analysis as the simple distance above:

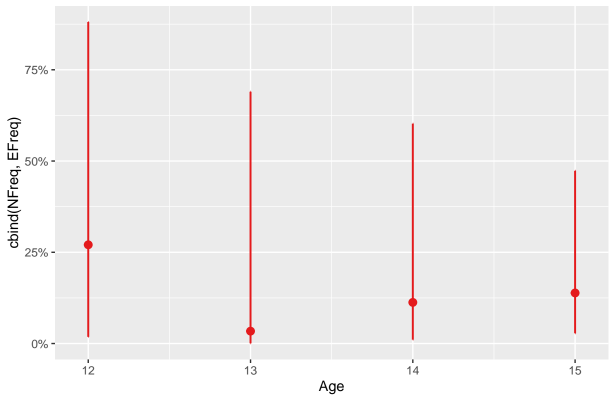
```
m0 = glmer(cbind(NFreq,EFreq) ~ Age + Alliteration
             + (1|Set) + (1|Source) +
            (1|EngForm2),
          data = d2, family = "binomial")
m1 = update(m0,~.+NormFeatureDistance)
m2 = update(m1,~.+I(NormFeatureDistance^2))
m3 = update(m2,~.+I(NormFeatureDistance^3))
## Warning in checkConv(attr(opt, "derivs"), opt$par, ctrl = control$checkConv, :
## Model failed to converge with max|grad| = 0.0469466 (tol = 0.002, component 1)
anova(m0,m1,m2,m3)
## Data: d2
## Models:
## m0: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2)
## m1: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2) + NormFe
## m2: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2) + NormFe
## m3: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2) + NormFe
          AIC BIC logLik deviance
                                       Chisq Df Pr(>Chisq)
     npar
## mO
       8 21104 21145 -10544
## m1
        9 20901 20946 -10442
                               20883 205.280 1 < 2.2e-16 ***
## m3 11 20582 20638 -10280 20560 276.202 1 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
summary(m3)
## Generalized linear mixed model fit by maximum likelihood (Laplace
   Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) +
      (1 | EngForm2) + NormFeatureDistance + I(NormFeatureDistance^2) +
##
      I(NormFeatureDistance^3)
     Data: d2
##
##
       AIC
                BIC
                     logLik deviance df.resid
## 20582.3 20638.1 -10280.2 20560.3
                                         1162
##
## Scaled residuals:
      Min 1Q Median
                                  3Q
                                          Max
## -102.933 -1.381 0.116
                               1.096 178.703
##
## Random effects:
                       Variance Std.Dev.
## Groups Name
## EngForm2 (Intercept) 4.740
                                2.177
## Set
            (Intercept) 5.840
                                2.417
## Source
            (Intercept) 4.365
                                2.089
## Number of obs: 1173, groups: EngForm2, 129; Set, 67; Source, 13
##
## Fixed effects:
                            Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                      0.829291 2.290
                                                          0.022 *
                            1.899237
                                       1.278915 0.834
                            1.066301
                                                          0.404
## Age1
## Age2
                           -1.291551
                                      1.646010 -0.785
                                                          0.433
                           -0.005518
## Age3
                                      1.123471 -0.005
                                                          0.996
```

```
## AlliterationTRUE
                                         0.071937 31.301
                              2.251681
                                                            <2e-16 ***
                                         1.781901 -18.200
## NormFeatureDistance
                            -32.430222
                                                            <2e-16 ***
                                         4.551277 17.015
## I(NormFeatureDistance^2) 77.441357
                                                            <2e-16 ***
## I(NormFeatureDistance^3) -51.690896
                                         3.157168 -16.373
                                                            <2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
                                           AlTRUE NrmFtD I(NFD^2
##
               (Intr) Age1
                             Age2
                                    Age3
## Age1
                0.006
                0.404 -0.522
## Age2
## Age3
               -0.187 -0.259 -0.490
## AlltrtnTRUE -0.014 0.015 -0.010 -0.001
## NrmFtrDstnc -0.144 0.005 -0.017 0.011 -0.024
## I(NrmFtD^2) 0.135 -0.006 0.017 -0.011 0.026 -0.981
## I(NrmFtD^3) -0.125  0.006 -0.017  0.010 -0.028  0.934 -0.984
## optimizer (Nelder_Mead) convergence code: 0 (OK)
## Model failed to converge with max|grad| = 0.0469466 (tol = 0.002, component 1)
pNormFeature =
  plot_model(m3,'eff', terms="NormFeatureDistance[all]")+
    xlab("Feature-Based Distance\nBetween Forms") +
    ylab("Proportion of Norse Terms") +
    ggtitle("")+
    coord_cartesian(ylim=c(0,1))
pNormFeature
  100% -
```



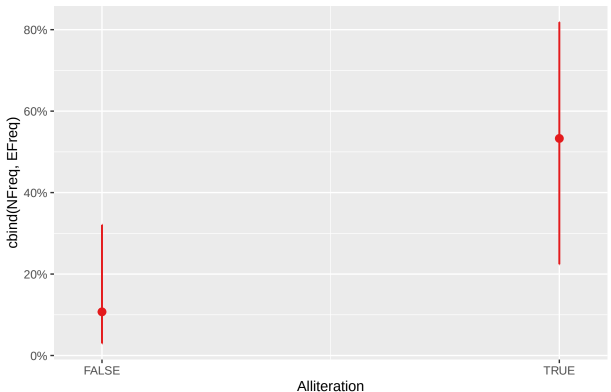
Effects over time and of alliteration:

```
plot_model(m3,'eff', terms="Age [all]")
```



plot_model(m3,'eff', terms="Alliteration")

Predicted probabilities of cbind(NFreq, EFreq)



GAM model:

```
s(Age, bs = "re") +
             s(Set,bs="re") +
             s(Source, bs="re")+
             s(EngForm2, bs="re"),
           data = d2, family = "binomial")
summary (mGAM)
##
## Family: binomial
## Link function: logit
##
## Formula:
## cbind(NFreq, EFreq) ~ s(NormFeatureDistance, k = 4) + s(Age,
       bs = "re") + s(Set, bs = "re") + s(Source, bs = "re") + s(EngForm2,
       bs = "re")
##
##
## Parametric coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) 0.09027
                           2.81544
                                     0.032
##
## Approximate significance of smooth terms:
                                edf Ref.df
                                              Chi.sq p-value
## s(NormFeatureDistance) 2.99978
                                        3 3.674e+02 < 2e-16 ***
## s(Age)
                           0.04248
                                         3 4.889e+03
                                                       0.9993
                          52.70122
## s(Set)
                                        66 7.617e+07
                                                       0.0244 *
## s(Source)
                           11.95629
                                        12 1.658e+07 6.75e-05 ***
## s(EngForm2)
                          69.37364
                                       128 2.384e+07
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) = 0.711 Deviance explained = 69.1\%
## UBRE = 16.565 Scale est. = 1
                                         n = 1173
plot.gam(mGAM,
         ylab="Relative Norse Freq",
         xlab="Distance between forms",
         select = 1)
Relative Norse Freq
     0
      7
     7
            0.0
                         0.2
                                       0.4
                                                     0.6
                                                                   0.8
                                                                                 1.0
```

Distance between forms

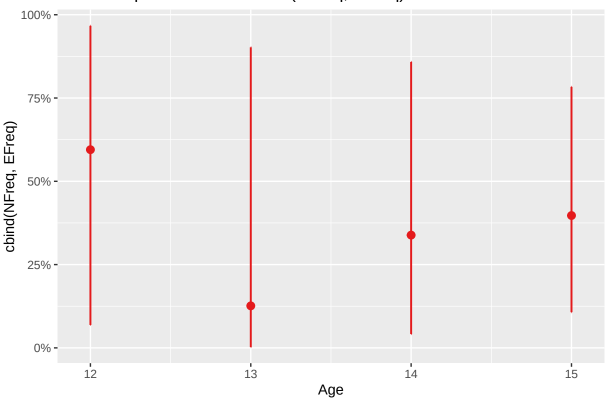
Linear: beta = -32.43, z = -18.2, Wald p < 0.001, LLDiff = 102.6, df = 1, p < 0.001; Quadratic: beta = 77.441, z = 17.015, Wald p < 0.001, LLDiff = 23.2, df = 1, p < 0.001; Cubic: beta = -51.691, z = -16.373, Wald p < 0.001, LLDiff = 138.1, df = 1, p < 0.001

6.3 Historical distance

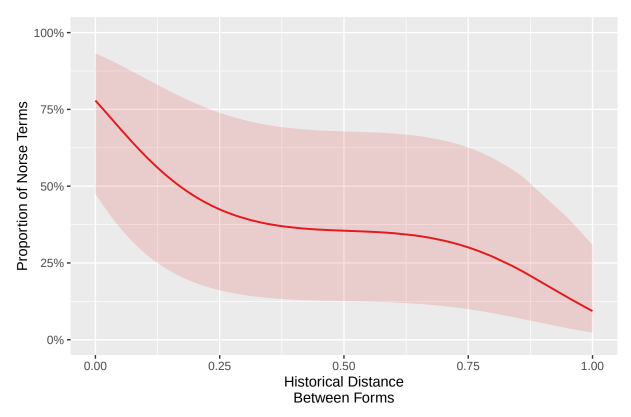
Same analysis as the simple distance above:

```
m0 = glmer(cbind(NFreq,EFreq) ~ Age + Alliteration +
             (1|Set) + (1|Source) +
             (1|EngForm2),
           data = d2, family = "binomial",
           control=glmerControl(optimizer="bobyqa"))
m1 = update(m0,~.+NormHistoricalDistance)
m2 = update(m1,~.+I(NormHistoricalDistance^2),
            control=glmerControl(optimizer="bobyqa"))
m3 = update(m2,~.+I(NormHistoricalDistance^3),
            control=glmerControl(optimizer="bobyqa"))
anova(m0,m1,m2,m3)
## Data: d2
## Models:
## m0: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2)
## m1: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2) + NormHi
## m2: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2) + NormHi
## m3: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) + (1 | EngForm2) + NormHi
##
           AIC BIC logLik deviance
     npar
                                         Chisq Df Pr(>Chisq)
## mO
        8 21104 21145 -10544
                                21088
        9 20904 20949 -10443
                                 20886 202.5263 1 < 2.2e-16 ***
## m1
## m2
       10 20901 20952 -10441
                                20881
                                       4.5342 1
                                                     0.03322 *
## m3
        11 20879 20935 -10429
                                20857 23.9212 1 1.004e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
For this measure, the cubic term is marginal.
This is the cubic model:
summary(m3)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) +
##
       (1 | EngForm2) + NormHistoricalDistance + I(NormHistoricalDistance^2) +
       I(NormHistoricalDistance^3)
##
      Data: d2
##
## Control: glmerControl(optimizer = "bobyqa")
##
##
        AIC
                 BIC
                      logLik deviance df.resid
   20879.2 20934.9 -10428.6 20857.2
##
                                           1162
##
## Scaled residuals:
       Min 1Q
                     Median
                                    3Q
                                           Max
## -110.625 -1.325
                     0.118
                                1.129 171.595
##
## Random effects:
## Groups
           Name
                        Variance Std.Dev.
## EngForm2 (Intercept) 4.065
                                 2.016
## Set
             (Intercept) 5.239
                                  2.289
             (Intercept) 4.425
## Source
                                 2.104
## Number of obs: 1173, groups: EngForm2, 129; Set, 67; Source, 13
##
## Fixed effects:
##
                                Estimate Std. Error z value Pr(>|z|)
```

```
## (Intercept)
                                           0.82206
                                                   1.123
                                                            0.261
                                0.92307
                                                             0.415
## Age1
                                1.04395
                                           1.28208
                                                    0.814
## Age2
                               -1.27604
                                           1.65242 -0.772
                                                             0.440
                                                              0.993
## Age3
                               -0.01052
                                           1.13003 -0.009
## AlliterationTRUE
                                           0.07204 31.190 < 2e-16 ***
                                2.24683
## NormHistoricalDistance
                              -10.42926
                                           1.41162 -7.388 1.49e-13 ***
## I(NormHistoricalDistance^2) 19.96760
                                           3.64632
                                                    5.476 4.35e-08 ***
## I(NormHistoricalDistance^3) -13.06820
                                           2.52493 -5.176 2.27e-07 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
              (Intr) Age1
                            Age2
                                   Age3
                                          AlTRUE NrmHsD I(NHD^2
               0.008
## Age1
               0.407 -0.519
## Age2
## Age3
              -0.189 -0.260 -0.492
## AlltrtnTRUE -0.014 0.015 -0.010 0.000
## NrmHstrclDs -0.128  0.003 -0.001 -0.002 -0.025
## I(NrmHsD^2) 0.114 -0.003 0.001 0.002 0.031 -0.976
## I(NrmHsD^3) -0.104  0.002  0.000 -0.002 -0.040  0.929 -0.984
plot_model(m3,'eff', terms="Age [all]")
```



```
pNormHist =
  plot_model(m3,'eff', terms="NormHistoricalDistance[all]")+
    xlab("Historical Distance\nBetween Forms") +
    ylab("Proportion of Norse Terms") +
    ggtitle("")+
    coord_cartesian(ylim=c(0,1))
pNormHist
```

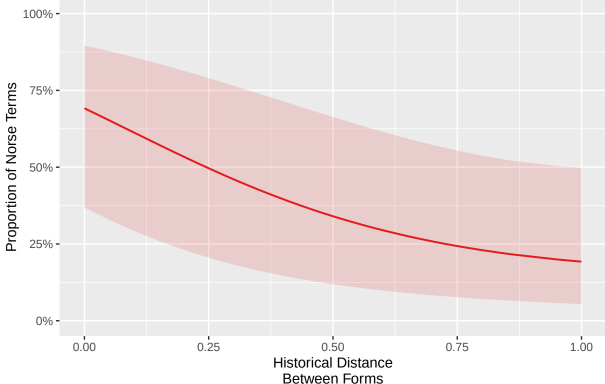


And for comparison, the quadratic model:

```
summary(m2)
```

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
## Family: binomial (logit)
## Formula: cbind(NFreq, EFreq) ~ Age + Alliteration + (1 | Set) + (1 | Source) +
       (1 | EngForm2) + NormHistoricalDistance + I(NormHistoricalDistance^2)
##
      Data: d2
## Control: glmerControl(optimizer = "bobyqa")
##
                BIC
                      logLik deviance df.resid
##
        AIC
   20901.1 20951.8 -10440.6 20881.1
##
##
## Scaled residuals:
       Min
                 1Q
                                    3Q
                      Median
                                            Max
## -110.404
            -1.347
                       0.116
                                 1.150 171.447
##
## Random effects:
  Groups
           Name
                        Variance Std.Dev.
## EngForm2 (Intercept) 4.043
                                  2.011
                                  2.279
## Set
             (Intercept) 5.192
             (Intercept) 4.413
                                  2.101
## Number of obs: 1173, groups: EngForm2, 129; Set, 67; Source, 13
##
## Fixed effects:
##
                               Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                0.47469
                                          0.81655
                                                     0.581
                                                             0.5610
## Age1
                                1.03689
                                           1.28216
                                                     0.809
                                                             0.4187
## Age2
                               -1.26463
                                           1.65211
                                                   -0.765
                                                             0.4440
## Age3
                               -0.01151
                                           1.12830 -0.010
                                                            0.9919
                                           0.07193 31.047 < 2e-16 ***
## AlliterationTRUE
                                2.23317
                                          0.52262 -6.961 3.38e-12 ***
## NormHistoricalDistance
                               -3.63796
```

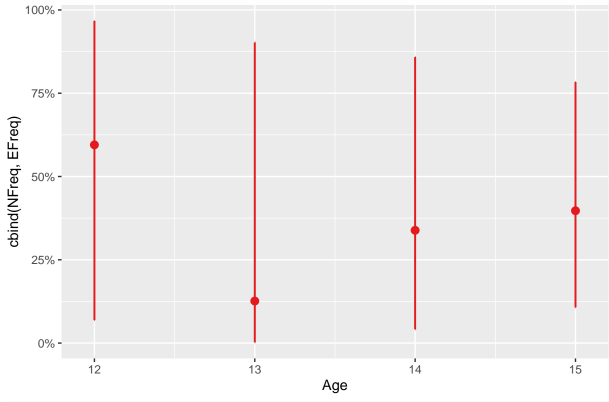
```
## I(NormHistoricalDistance^2) 1.39642
                                          0.65371
                                                   2.136 0.0327 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
##
               (Intr) Age1
                            Age2
                                  Age3
                                          AlTRUE NrmHsD
## Age1
               0.007
## Age2
               0.410 - 0.520
## Age3
              -0.190 -0.259 -0.491
## AlltrtnTRUE -0.019 0.015 -0.010 0.000
## NrmHstrclDs -0.086 0.000 -0.001 0.000 0.037
## I(NrmHsD^2) 0.065 -0.001 0.001 -0.001 -0.053 -0.931
 plot_model(m2,'eff', terms="NormHistoricalDistance[all]")+
   xlab("Historical Distance\nBetween Forms") +
   ylab("Proportion of Norse Terms") +
   ggtitle("")+
   coord_cartesian(ylim=c(0,1))
  100% -
```



There is little qualitative difference, so for sake of easy comparison with the other results, we use the cubic model.

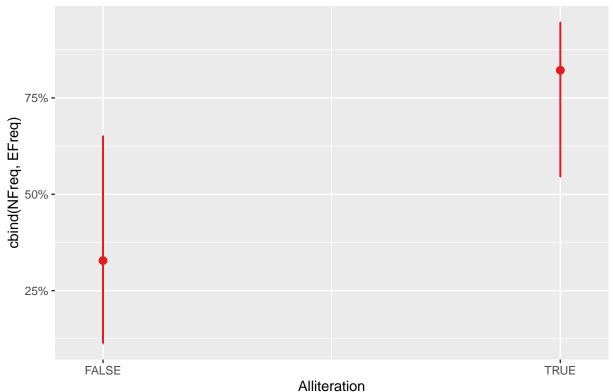
Effects over time and of alliteration:

```
plot_model(m3,'eff', terms="Age [all]")
```



plot_model(m3,'eff', terms="Alliteration")

Predicted probabilities of cbind(NFreq, EFreq)

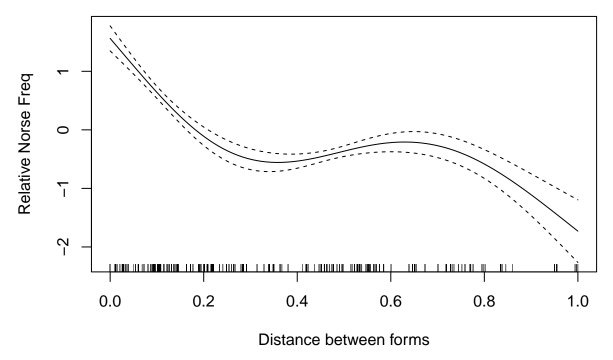


GAM model:

```
mGAM = gam(cbind(NFreq,EFreq) ~ Alliteration +
s(NormHistoricalDistance, k=4) +
```

```
s(Age, bs = "re") +
                                  s(Set,bs="re") +
                                  s(Source, bs="re")+
                                  s(EngForm2, bs="re"),
                             data = d2, family = "binomial")
summary (mGAM)
##
## Family: binomial
## Link function: logit
##
## Formula:
## cbind(NFreq, EFreq) ~ Alliteration + s(NormHistoricalDistance,
                 k = 4) + s(Age, bs = "re") + <math>s(Set, bs = "re") + s(Source, bs = "re") + <math>s(Source, bs = "re") + s(Source, bs 
                  bs = "re") + s(EngForm2, bs = "re")
## Parametric coefficients:
                                                     Estimate Std. Error z value Pr(>|z|)
##
                                                  -0.35137 2.38755 -0.147 0.883
## (Intercept)
                                                                                     0.07209 31.144
## AlliterationTRUE 2.24532
                                                                                                                                  <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Approximate significance of smooth terms:
                                                                                             edf Ref.df
                                                                                                                                 Chi.sq p-value
## s(NormHistoricalDistance) 2.995951 3 2.472e+02 < 2e-16 ***
## s(Age)
                                                                               0.001067
                                                                                                                   3 1.164e+00 0.999276
                                                                                                         66 6.216e+07 0.000642 ***
## s(Set)
                                                                             47.845260
## s(Source)
                                                                             11.996842
                                                                                                               12 1.098e+07 < 2e-16 ***
## s(EngForm2)
                                                                             ## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) = 0.726 Deviance explained = 70.3\%
## UBRE = 15.903 Scale est. = 1
plot.gam(mGAM,
                       ylab="Relative Norse Freq",
                       xlab="Distance between forms",
```

select = 1)



Linear: beta = -10.429, z = -7.388, Wald p < 0.001, LLDiff = 101.3, df = 1, p < 0.001; Quadratic: beta = 19.968, z = 5.476, Wald p < 0.001, LLDiff = 2.3, df = 1, p = 0.033; Cubic: beta = -13.068, z = -5.176, Wald p < 0.001, LLDiff = 12, df = 1, p < 0.001

```
bigPlot = grid.arrange(
       pNormSimple +
          ggtitle("Source-Level Analysis"),
       pNormFeature+
         ggtitle("") +
         theme(axis.title.y=element_blank(),
               axis.text.y=element_blank()),
       pNormHist+
         ggtitle("") +
         theme(axis.title.y=element_blank(),
               axis.text.y=element_blank()),
       nrow=1,widths=c(1.3,1,1))
pdf("../results/BigEffectsPlot.pdf", width = 8, height=3)
 plot(bigPlot)
dev.off()
## pdf
##
       Source-Level Analysis
   100%
   75%
```

Proportion of Norse Terms 50% 25% 0% 0.00 0.25 0.50 0.75 1.00 0.00 0.25 0.50 0.75 1.00 0.00 0.25 0.50 1.00 Sound Class Distance Feature-Based Distance Historical Distance

Between Forms

Between Forms

Between Forms

```
tSimple = read.csv("../results/SimpleRes_totalFreq.csv",
                  stringsAsFactors = F)
tFeature = read.csv("../results/FeatureRes_totalFreq.csv",
                   stringsAsFactors = F)
tHistorical = read.csv("../results/HistoricalRes_totalFreq.csv",
                      stringsAsFactors = F)
sSimple = read.csv("../results/SimpleRes.csv",
                  stringsAsFactors = F)
sFeature = read.csv("../results/FeatureRes.csv",
                   stringsAsFactors = F)
sHistorical = read.csv("../results/HistoricalRes.csv",
                      stringsAsFactors = F)
res = rbind(tSimple,tFeature,tHistorical)
resNames = c(lldiff="Log Likelihood Difference",
            Chisq="Chi Squared",pChi="p")
resNames2 = names(res)
resNames2[resNames2 %in% names(resNames)] =
 resNames[resNames2[resNames2 %in% names(resNames)]]
res = res[,names(res)!="X"]
res = cbind(Measure=rep(c("Sound Class", "Feature", "Historical"), each=3), res)
"Model Estimate","",""),
```

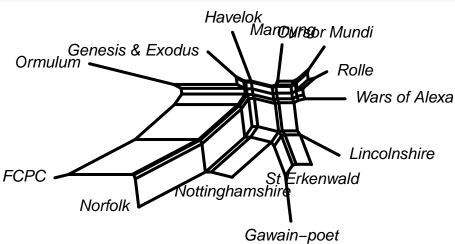
7 Comparison between texts

Compare sources according to the difference in proportion of Norse terms for each set.

```
g = read.xlsx("../data/SharedIntegrationOfCognatesData.xlsx",1)
g = g[g$Etymology %in% c("Norse", "English"),]
# Ignore numerals in set name
g$Set = gsub("[0-9]","",g$Set)
nCols = names(g)[which(names(g)=="No..in.Ormulum"):which(names(g)=="Rolle")]
for(col in nCols){
  g[,col] = as.numeric(g[,col])
## Warning: NAs introduced by coercion
## Warning: NAs introduced by coercion
## Warning: NAs introduced by coercion
allSets = unique(g$Set)
f = data.frame(Set = allSets)
fLog = data.frame(Set = allSets)
for(col in nCols){
 gN = g[g$Etymology == "Norse",]
 gE = g[g$Etymology == "English",]
 fNorse = tapply(gN[,col],gN$Set,sum)[allSets]
 fNorse[is.na(fNorse)] = 0
 fEng = tapply(gE[,col],gE$Set,sum)[allSets]
 fEng[is.na(fEng)] = 0
 f[,col] = fNorse / (fEng+fNorse)
 fLog[,col] = log10(1+fEng) - log10(1+fNorse)
nColsLabels = c("Ormulum", "FCPC", "Havelok", "Genesis & Exodus",
                "Mannyng", "Gawain-poet", "Wars of Alexander",
                "St Erkenwald", "Cursor Mundi", "Lincolnshire",
                "Nottinghamshire", "Norfolk", "Rolle")
mat = matrix(NA, nrow=length(nCols),
            ncol = length(nCols))
rownames(mat) = nColsLabels
colnames(mat) = nColsLabels
matLog = matrix(NA, nrow=length(nCols),
                ncol = length(nCols))
rownames(matLog) = nColsLabels
colnames(matLog) = nColsLabels
for(i in 1:length(nCols)){
  iProp = f[,nCols[i]]
  iPropLog = fLog[,nCols[i]]
 for(j in 1:length(nCols)){
    jProp = f[,nCols[j]]
    jPropLog = fLog[,nCols[j]]
   diffs = abs(iProp-jProp)
   diffs = diffs[!is.nan(diffs)]
   diffs = diffs[!is.na(diffs)]
   mat[nColsLabels[i],nColsLabels[j]] = mean(diffs)
```

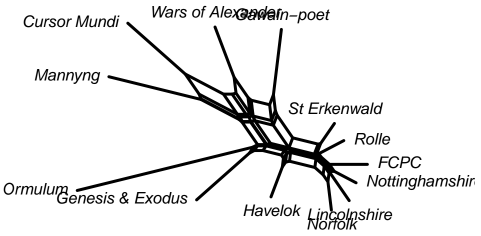
```
diffsLog = abs(iPropLog - jPropLog)
  matLog[nColsLabels[i],nColsLabels[j]] = mean(diffsLog)
}

phy = neighborNet(mat)
plot(phy)
```



```
pdf("../results/NeighbourNet.pdf")
plot(phy)
dev.off()

## pdf
## 2
phyLog = neighborNet(matLog)
plot(phyLog)
```



```
pdf("../results/NeighbourNet_Log.pdf")
plot(phyLog)
dev.off()
```

pdf ## 2

8 Bayesian modelling

8.1 Simple distance

In this section, we replicate the analyses above using a Bayesian framework. As with the example of a Bayesian analysis above, we use a beta binomial framework.

```
d$NormDistanceChosen = d$NormDistance
d$totalAllFreq = d$totalNFreq + d$totalEFreq
mBSimple = brm(totalNFreq | trials(totalAllFreq) ~
               s(NormDistanceChosen, k=4) +
             (1|Set) + (1|EngLexeme),
           data = d, family = "beta_binomial",
         silent = 2,refresh=0)
## Warning: There were 18 divergent transitions after warmup. See
## https://mc-stan.org/misc/warnings.html#divergent-transitions-after-warmup
## to find out why this is a problem and how to eliminate them.
## Warning: Examine the pairs() plot to diagnose sampling problems
## Warning: Bulk Effective Samples Size (ESS) is too low, indicating posterior means and medians may
## Running the chains for more iterations may help. See
## https://mc-stan.org/misc/warnings.html#bulk-ess
## Warning: Tail Effective Samples Size (ESS) is too low, indicating posterior variances and tail qu
## Running the chains for more iterations may help. See
## https://mc-stan.org/misc/warnings.html#tail-ess
summary(mBSimple)
## Warning: There were 18 divergent transitions after warmup. Increasing
## adapt_delta above 0.8 may help. See
## http://mc-stan.org/misc/warnings.html#divergent-transitions-after-warmup
## Family: beta_binomial
    Links: mu = logit; phi = identity
##
## Formula: totalNFreq | trials(totalAllFreq) ~ s(NormDistanceChosen, k = 4) + (1 | Set) + (1 | EngL
     Data: d (Number of observations: 1638)
     Draws: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
            total post-warmup draws = 4000
##
## Smooth Terms:
##
                              Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS
                                  3.05
                                            1.98
                                                     0.47
                                                               8.06 1.01
## sds(sNormDistanceChosen_1)
##
                              Tail ESS
## sds(sNormDistanceChosen_1)
##
## Group-Level Effects:
## ~EngLexeme (Number of levels: 135)
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sd(Intercept)
                     0.68
                               0.08
                                        0.53
                                                 0.84 1.00
                                                                 618
##
## ~Set (Number of levels: 67)
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
                                        0.04
## sd(Intercept)
                     0.34
                               0.14
                                                 0.62 1.01
                                                                 251
                                                                          389
##
## Population-Level Effects:
##
                         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS
                                       0.10
## Intercept
                             0.26
                                                0.07
                                                         0.46 1.00
                                                                        1610
                                       2.58
## sNormDistanceChosen_1
                            -4.96
                                              -10.10
                                                         -0.52 1.01
                                                                        1196
##
                         Tail_ESS
```

```
## Intercept
                              2701
## sNormDistanceChosen_1
                              1707
##
## Family Specific Parameters:
       Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## phi
           1.99
                      0.08
                               1.85
                                         2.14 1.00
                                                       5180
                                                                 2723
##
## Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
pred = ggpredict(mBSimple,
                 terms="NormDistanceChosen")
## Note: uncertainty of error terms are not taken into account. Consider
     setting `interval` to "prediction". This will call `posterior_predict()`
##
     instead of `posterior_epred()`.
##
## Warning in checkMatrixPackageVersion(): Package version inconsistency detected.
## TMB was built with Matrix version 1.6.1
## Current Matrix version is 1.6.5
## Please re-install 'TMB' from source using install.packages('TMB', type = 'source') or ask CRAN fo
# Predict uses 10 trials
pred$predicted = pred$predicted/10
pred$conf.low = pred$conf.low/10
pred$conf.high = pred$conf.high/10
ggplot(pred,aes(x=x,y=predicted,ymin=conf.low,ymax=conf.high)) +
  geom_ribbon(alpha=0.3)+
  geom_line() +
  xlab("Distance") +
  ylab("Predicted Likelihood")
  0.7 -
  0.6 -
Predicted Likelihood
  0.3 -
  0.2 -
                           0.25
                                                                 0.75
        0.00
                                              0.50
                                                                                     1.00
```

In the analyses above, we set the number of knots at 4. We can test this setting by running two different

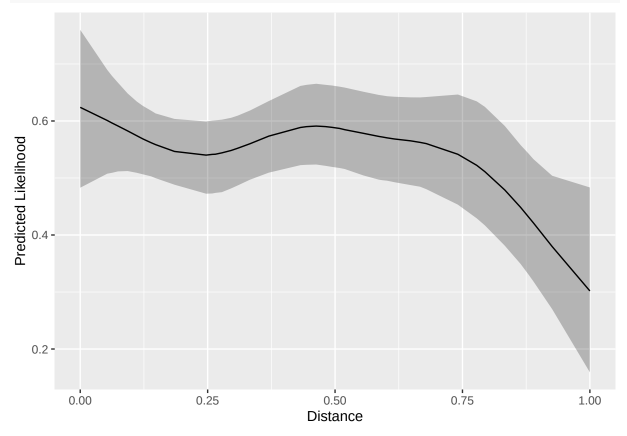
Distance

models with different number of knots:

```
mBSimple.k4 = brm(totalNFreq | trials(totalAllFreq) ~
                s(NormDistanceChosen, k=4) +
              (1|Set) + (1|EngLexeme),
           data = d, family = "beta_binomial",
         silent = 2,refresh=0)
## recompiling to avoid crashing R session
## Warning: There were 9 divergent transitions after warmup. See
## https://mc-stan.org/misc/warnings.html#divergent-transitions-after-warmup
## to find out why this is a problem and how to eliminate them.
## Warning: Examine the pairs() plot to diagnose sampling problems
pred = ggpredict(mBSimple.k4,
                  terms="NormDistanceChosen")
## Note: uncertainty of error terms are not taken into account. Consider
     setting `interval` to "prediction". This will call `posterior_predict()`
##
     instead of `posterior_epred()`.
# Predict uses 10 trials
pred$predicted = pred$predicted/10
pred$conf.low = pred$conf.low/10
pred$conf.high = pred$conf.high/10
ggplot(pred,aes(x=x,y=predicted,ymin=conf.low,ymax=conf.high)) +
  geom ribbon(alpha=0.3)+
  geom_line() +
  xlab("Distance") +
  ylab("Predicted Likelihood")
   0.7 -
   0.6 -
Predicted Likelihood
  0.3 -
   0.2 -
                           0.25
        0.00
                                                                 0.75
                                              0.50
                                                                                     1.00
                                            Distance
mBSimple.k8 = brm(totalNFreq | trials(totalAllFreq) ~
                s(NormDistanceChosen, k=8) +
```

```
(1|Set) + (1|EngLexeme),
           data = d, family = "beta_binomial",
         silent = 2,refresh=0)
## recompiling to avoid crashing R session
## Warning: There were 19 divergent transitions after warmup. See
## https://mc-stan.org/misc/warnings.html#divergent-transitions-after-warmup
## to find out why this is a problem and how to eliminate them.
## Warning: Examine the pairs() plot to diagnose sampling problems
pred = ggpredict(mBSimple.k8,
                 terms="NormDistanceChosen[all]")
## Note: uncertainty of error terms are not taken into account. Consider
     setting `interval` to "prediction". This will call `posterior_predict()`
##
     instead of `posterior_epred()`.
# Predict uses 10 trials
pred$predicted = pred$predicted/10
pred$conf.low = pred$conf.low/10
pred$conf.high = pred$conf.high/10
ggplot(pred,aes(x=x,y=predicted,ymin=conf.low,ymax=conf.high)) +
 geom_ribbon(alpha=0.3)+
 geom_line() +
 xlab("Distance") +
```

ylab("Predicted Likelihood")



There is very little difference in the overall shape of the relationship between k=4 and k=8. At least, there are no conclusions we would draw from the k=8 fit that we would not draw from the k=4 fit. We decided to stick with the simpler model (k=4).

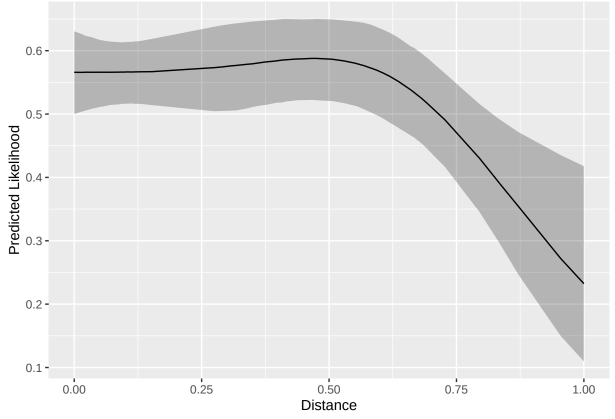
Our conclusion was, as stated in the discussion: "we argue that the overall picture is that the effect exists for extreme ends of form distance". That is, there may be some non-linear effect in the middle, but the

basic picture is that low distance = likely to be Norse, high distance = likely to be Native. All of the analyses fit this general description.

8.2 Feature distance

```
d$NormDistanceChosen = d$NormFeatureDistance
d$totalAllFreq = d$totalNFreq + d$totalEFreq
mBSimple = brm(totalNFreq | trials(totalAllFreq) ~
               s(NormDistanceChosen, k=4) +
             (1|Set) + (1|EngLexeme),
           data = d, family = "beta_binomial",
         silent = 2,refresh=0)
## recompiling to avoid crashing R session
## Warning: There were 3 divergent transitions after warmup. See
## https://mc-stan.org/misc/warnings.html#divergent-transitions-after-warmup
## to find out why this is a problem and how to eliminate them.
## Warning: Examine the pairs() plot to diagnose sampling problems
## Warning: Bulk Effective Samples Size (ESS) is too low, indicating posterior means and medians may
## Running the chains for more iterations may help. See
## https://mc-stan.org/misc/warnings.html#bulk-ess
summary(mBSimple)
## Warning: There were 3 divergent transitions after warmup. Increasing
## adapt_delta above 0.8 may help. See
## http://mc-stan.org/misc/warnings.html#divergent-transitions-after-warmup
## Family: beta_binomial
    Links: mu = logit; phi = identity
## Formula: totalNFreq | trials(totalAllFreq) ~ s(NormDistanceChosen, k = 4) + (1 | Set) + (1 | EngL
     Data: d (Number of observations: 1638)
##
     Draws: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
            total post-warmup draws = 4000
##
##
## Smooth Terms:
                              Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS
##
## sds(sNormDistanceChosen_1)
                                  2.69
                                            1.52
                                                      0.71
                                                               6.58 1.00
##
                              Tail_ESS
## sds(sNormDistanceChosen_1)
                                  2171
## Group-Level Effects:
## ~EngLexeme (Number of levels: 135)
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
                     0.68
                               0.08
## sd(Intercept)
                                        0.53
                                                                 814
                                                                         1562
##
## ~Set (Number of levels: 67)
                 Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
##
                     0.38
                               0.13
                                        0.09
                                                  0.64 1.02
                                                                 294
## sd(Intercept)
                                                                          497
##
## Population-Level Effects:
##
                         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS
## Intercept
                             0.25
                                       0.10
                                                0.05
                                                         0.44 1.00
## sNormDistanceChosen_1
                                                -8.29
                                                         -0.72 1.00
                            -4.11
                                       1.98
                                                                        1339
                         Tail ESS
## Intercept
                             2432
## sNormDistanceChosen_1
                             1843
##
```

```
## Family Specific Parameters:
       Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
##
## phi
                    0.08
                              1.85
                                       2.16 1.00
                                                     4468
##
## Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
pred = ggpredict(mBSimple,
                 terms="NormDistanceChosen")
## Note: uncertainty of error terms are not taken into account. Consider
     setting `interval` to "prediction". This will call `posterior_predict()`
     instead of `posterior_epred()`.
# Predict uses 10 trials
pred$predicted = pred$predicted/10
pred$conf.low = pred$conf.low/10
pred$conf.high = pred$conf.high/10
ggplot(pred,aes(x=x,y=predicted,ymin=conf.low,ymax=conf.high)) +
  geom_ribbon(alpha=0.3)+
  geom_line() +
  xlab("Distance") +
  ylab("Predicted Likelihood")
```



8.3 Historical distance

```
data = d, family = "beta_binomial",
         silent = 2,refresh=0)
## recompiling to avoid crashing R session
## Warning: There were 7 divergent transitions after warmup. See
## https://mc-stan.org/misc/warnings.html#divergent-transitions-after-warmup
## to find out why this is a problem and how to eliminate them.
## Warning: Examine the pairs() plot to diagnose sampling problems
## Warning: Bulk Effective Samples Size (ESS) is too low, indicating posterior means and medians may
## Running the chains for more iterations may help. See
## https://mc-stan.org/misc/warnings.html#bulk-ess
summary(mBSimple)
## Warning: There were 7 divergent transitions after warmup. Increasing
## adapt delta above 0.8 may help. See
## http://mc-stan.org/misc/warnings.html#divergent-transitions-after-warmup
## Family: beta_binomial
   Links: mu = logit; phi = identity
## Formula: totalNFreq | trials(totalAllFreq) ~ s(NormDistanceChosen, k = 4) + (1 | Set) + (1 | EngL
     Data: d (Number of observations: 1638)
     Draws: 4 chains, each with iter = 2000; warmup = 1000; thin = 1;
##
##
            total post-warmup draws = 4000
## Smooth Terms:
                              Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS
##
## sds(sNormDistanceChosen_1)
                                  2.71
                                            1.50
                                                     0.80
                                                              6.60 1.00
                              Tail ESS
## sds(sNormDistanceChosen_1)
                                  2215
##
## Group-Level Effects:
## ~EngLexeme (Number of levels: 135)
##
                Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk ESS Tail ESS
## sd(Intercept)
                     0.67
                               0.08
                                        0.52
                                                 0.84 1.01
                                                                749
## ~Set (Number of levels: 67)
                Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## sd(Intercept)
                    0.36
                               0.14
                                        0.07
                                                0.63 1.02
                                                                296
                                                                         421
## Population-Level Effects:
                         Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS
##
## Intercept
                             0.25
                                  0.10
                                                0.06
                                                     0.44 1.00
                                                                       1837
                            -3.33
                                       2.22
                                               -8.01
                                                         0.50 1.00
## sNormDistanceChosen_1
                                                                       1649
                         Tail_ESS
##
## Intercept
                             2266
## sNormDistanceChosen_1
                             2266
## Family Specific Parameters:
##
      Estimate Est.Error 1-95% CI u-95% CI Rhat Bulk_ESS Tail_ESS
## phi
           1.99
                    0.07
                             1.85
                                       2.14 1.00
                                                     5246
## Draws were sampled using sampling(NUTS). For each parameter, Bulk_ESS
## and Tail_ESS are effective sample size measures, and Rhat is the potential
## scale reduction factor on split chains (at convergence, Rhat = 1).
pred = ggpredict(mBSimple,
                 terms="NormDistanceChosen")
```

```
## Note: uncertainty of error terms are not taken into account. Consider
## setting `interval` to "prediction". This will call `posterior_predict()`
## instead of `posterior_epred()`.

# Predict uses 10 trials
pred$predicted = pred$predicted/10
pred$conf.low = pred$conf.low/10
pred$conf.high = pred$conf.high/10
ggplot(pred,aes(x=x,y=predicted,ymin=conf.low,ymax=conf.high)) +
    geom_ribbon(alpha=0.3)+
    geom_line() +
    xlab("Distance") +
    ylab("Predicted Likelihood")
```

