Supplementary Information 4 for Cultural influences on word meanings revealed throughlarge-scale semantic alignment: Cross-cultural analyses

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This document contains the supporting information on cross-cultural analyses for *Cultural influences on word meanings revealed throughlarge-scale semantic alignment*. Most of the contents are compiled R markdown documents, showing the R code for analysing the data. The full repository can be found here:

https://github.com/seannyD/ImputeEACulturalDifferences

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4.1 Main cross-cultural analysis (Wikipedia data)

Predicting semantic alignment by cultural similarity

Bill Thompson, Seán Roberts & Gary Lupyan

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Introduction

We compare cultural distances between socieites with linguistic similarities between societies, controlling for shared history in two ways.

The first test uses mixed effects modelling. The pairing of the language family of each language (according to Glottolog, Hammarstrom et al., 2018) is used as a random effect. That means that the model can capture the likelihood that two languages from the Indo-European language family will be more similar to each other than two languages from different language families. The same is done with geographic area according to Autotyp (Nichols et al., 2013). The model also included a fixed effect for for the number of lexical comparisons that went into the mean semantic alignment estimates (generally, more available comparisons indicate more possible comparisons, i.e. more similar languages).

The second test controls for history using distances from a phylogenetic tree. The tree comes from Bouckaert et al. (2012). Patristic distances between languages are used as a measure of historical distance between societies in a Mantel test. Note that the Mantel test assumes a strict distance metric, which is not necessarily the case with this data, but there are few other ways to deal with continuous pairwise distances. Mantel tests were used to test the relationship between semantic alignment and historical proximity and between semantic alignment and geographic proximity. Geographic proximity was measured as the great circle distance between the cultural centres of each language as defined in Glottolog. For the analysis within domains, partial Mantel tests were used to estimate the correlation between the semantic alignment and the cultural/geographic/historical proximity while partialling out the effect of the other two proximity measures.

The effect was also confirmed using multiple regression on distance matrices (Lichstein, 2007). The results above were robustly replicated using the filtered data and also alternative sources for semantic alignment (common crawl, see file $AnalyseCorrelation_cc.pdf$). The correlation was not robust to all tests or for data derived from the subtitles dataset (see file $AnalyseCorrelation_subs.pdf$), possibly because there were only 20 languages available to analyse.

The final section looks at relationships between sub-domains. The first section describes how the cultural similarity measure was calculated.

Calculating cultural similarity

The aim is to produce a set of distances between societies based on their cultural traits. The Ethnographic Atlas (Murdock et al., 1999) as made available in D-PLACE (Kirby et al., 2016) could do this, but there is a lot of missing data (about 25% in the whole dataset), which means that distances can't be computed easily. One approach is to impute the missing data (guess their values based on existing data). It's unlikely that any imputation method will be completely accurate, but for our purposes we don't need to be accurate, just unbiased. That is, the imputed values should not bias the estimates of the distances between cultures.

In this case, we use multiple imputation: calculating many possible alternative imputations and taking the mean distances over all imputations.

Imputing missing values in the Ethnographic Atlas

We use the imputation package mice for R (van Buuren & Groothuis-Oudshoorn, 2011). We compared various settings of the imputation method, and found that using classification and regression trees (CART) with the standard parameters produced the best results. CART works by building a decision tree: an optimal set of yes-no questions to ask about predictor variables in order to guess the value of a target variable. The tree divides the data into partitions which look similar. The algorithm works out which partition a missing data point would belong to, then samples the target variable distribution from that partition. To account for historical relationships, we included language family according to Glottolog and geographic area according to Autotyp as additional factors on which the imputation process could draw.

We ran CART multiple imputation on the Ethnographic Atlas. We excluded population size, one more variable that was coded for less than 33% of societies, and any societies that had fewer than 33% variables coded. This left 92 variables for 962 languages with 16% missing data.

We tested the imputation by taking the full Ethnographic Atlas data, creating some new missing values in random places and then re-imputing those missing values. We can then asses how accurate the imputation was for those values. Since the main analysis would only be using a small sub-set of the data, it is important to assess performance on these in particular, rather than the entire set of languages. Missing data was only inserted for languages in the main analysis of semantic alignment below. CART imputation guessed the correct value of missing data 74% of the time (average over 100 imputations). This is reasonably good, considering that most variables have between 4 and 8 possible values (median = 6). For example, this is 8.6 standard deviations better than choosing randomly (accuracy = 19%) and 5.6 standard deviations better than sampling from the known distribution of the target variable (accuracy = 37% on the same missing data). This is not good enough to use in analyses that look at individual traits, but serves our purposes to estimate overall distances between languages.

We produced 100 imputation sets with the final settings. These were then used to create distance matrices using Gower distance between discrete traits (mean correlation between sets r=0.94, estimates of distance vary by around 2% on average). The final distance matrix was the mean of each of the 100 distance matrices. Distances were also calculate for sub-domains of the data.

The full scripts and data can be found at https://github.com/seannyD/ImputeEACulturalDifferences. Reviewers can follow this link: https://figshare.com/s/06378bc59a771d28b1d0

Load libraries

```
library(ape)
library(ecodist)
library(lme4)
library(sjPlot)
library(ggplot2)
library(igraph)
library(lattice)
library(dplyr)
```

Parameters (using data from Northuralex and Wikipedia, k=100, unfiltered):

```
datasetName = "wikipedia-main"
lingDistancesFile = "../data/FAIR/nel-wiki-k100-alignments-by-language-pair.csv"
lingDistancesFileNK = "../data/FAIR/nel-wiki-k100-alignments-by-language-pair-without-kinship.csv"
lingDistancesByDomainFile = "../results/EA_distances/nel-wiki-k100_with_ling.csv"
# (generated by ../processing/combineCultAndLingDistances.R)
```

All domains

Load data

Read the cultural distances:

```
cult = read.csv("../results/EA_distances/CulturalDistances_Long.csv", stringsAsFactors = F)
names(cult) = c("11","12","cult.dist")

Add language family:

l = read.csv("../data/FAIR_langauges_glotto_xdid.csv", stringsAsFactors = F)

g = read.csv("../data/glottolog-languoid.csv/languoid.csv", stringsAsFactors = F)

l$family = g[match(1$glotto,g$id),]$family_pk

l$family = g[match(1$family,g$pk),]$name

Read the semantic distances

ling = read.csv(lingDistancesFile, stringsAsFactors = F)
```

There are very few possible comparisons for Slovenian and Northern Sami, so we'll remove these:

```
ling = ling[!(ling$11=="se" | ling$12 == "se"),]
ling = ling[!(ling$11=="s1" | ling$12 == "s1"),]
```

Combine the linguistic and cultural distances. Note that we flip the cultural measure from a distance measure to a similarity measure.

```
cult$11.iso2 = 1[match(cult$11,1$Language2),]$iso2
cult$12.iso2 = 1[match(cult$12,1$Language2),]$iso2
fairisos = unique(c(ling$11,ling$12))
cultisos = unique(c(cult$11.iso2, cult$12.iso2))
cult = cult[(cult$11.iso2 %in% fairisos) & (cult$12.iso2 %in% fairisos),]
ling = ling[(ling$11 %in% cultisos) & (ling$12 %in% cultisos),]
matches = sapply(1:nrow(ling), function(i){
 which(cult$11.iso2==ling$11[i] & cult$12.iso2==ling$12[i])
})
ling$cult.dist = cult[matches,]$cult.dist
# Flip
ling$cult.dist = 1 - ling$cult.dist
# Scale
ling$cult.dist.center = scale(ling$cult.dist)
cdc.s = attr(ling$cult.dist.center, "scaled:scale")
cdc.c = attr(ling$cult.dist.center, "scaled:center")
ling$cult.dist.center = as.numeric(ling$cult.dist.center)
ling$comparison_count.center =
  scale(ling$comparison_count)
ling$family1 = l[match(ling$11, l$iso2),]$family
ling$family2 = 1[match(ling$12, 1$iso2),]$family
1[1$Language=="Arabic",]$autotyp.area= "Greater Mesopotamia"
1[1$Language=="Persian",]$autotyp.area= "Greater Mesopotamia"
ling$area1 = l[match(ling$11, l$iso2),]$autotyp.area
```

```
ling$area2 = 1[match(ling$12, 1$iso2),]$autotyp.area
fgroup = cbind(ling$family1,ling$family2)
fgroup = apply(fgroup,1,sort)
ling$family.group = apply(fgroup,2,paste,collapse=":")
agroup = cbind(ling$area1,ling$area2)
agroup = apply(agroup,1,sort)
ling$area.group = apply(agroup,2,paste,collapse=":")
ling$rho.center = scale(ling$local_alignment)
Each observation is now assocaited with a language family pair:
head(ling[,c("11","12","local_alignment",'family.group')])
##
      11 12 local_alignment
                                           family.group
                                   Abkhaz-Adyge:Japonic
## 7
      ja ab 0.01930414
## 8
     ab zh
                 0.02225169 Abkhaz-Adyge:Sino-Tibetan
## 10 cv xal
                 0.02765860
                                        Mongolic:Turkic
                 0.02832668
## 11 xal ja
                                       Japonic:Mongolic
## 12 xal zh
                 0.02895876
                                  Mongolic:Sino-Tibetan
                 0.03192066 Abkhaz-Adyge:Indo-European
## 14 bn ab
And the same is true for area:
tail(ling[,c("11","12","local_alignment",'area.group')])
       11 12 local_alignment
                                        area.group
## 2522 fr es
              0.3936442
                                     Europe: Europe
## 2524 cs uk
                  0.4023323
                                Europe:Inner Asia
## 2528 cs ru
                  0.4082099
                                 Europe: Inner Asia
                  0.4129814 Inner Asia:Inner Asia
## 2529 be ru
## 2532 uk be
                   0.4276664 Inner Asia:Inner Asia
## 2535 uk ru
                   0.5079911 Inner Asia:Inner Asia
Number of observations:
# Number of datapoints:
nrow(ling)
## [1] 731
# Number of unique languages:
length(unique(unlist(ling[,c("11","12")])))
## [1] 39
# Number of unique language families:
uniqueFamilies = unique(unlist(ling[,c("family1","family2")]))
length(uniqueFamilies)
## [1] 10
# Number of unique areas:
uniqueAreas = unique(unlist(ling[,c("area1", "area2")]))
length(uniqueAreas)
```

Cross-over between language famlies and areas:

	Europe	${\tt Greater}$	Mesopotamia	${\tt Indic}$	Inner	Asia N	I Coast	Asia
Abkhaz-Adyge	0		1	0		0		0
Afro-Asiatic	0		1	0		0		0
Dravidian	0		0	3		0		0
Indo-European	11		2	1		5		0
Japonic	0		0	0		0		1
Koreanic	0		0	0		0		1
Mongolic	0		0	0		1		0
Sino-Tibetan	0		0	0		0		0
Turkic	0		1	0		5		0
Uralic	1		0	0		4		0
	Southea	ast Asia						
	Afro-Asiatic Dravidian Indo-European Japonic Koreanic Mongolic Sino-Tibetan Turkic	Abkhaz-Adyge 0 Afro-Asiatic 0 Dravidian 0 Indo-European 11 Japonic 0 Koreanic 0 Mongolic 0 Sino-Tibetan 0 Turkic 0 Uralic 1	Abkhaz-Adyge 0 Afro-Asiatic 0 Dravidian 0 Indo-European 11 Japonic 0 Koreanic 0 Mongolic 0 Sino-Tibetan 0 Turkic 0	Abkhaz-Adyge 0 1 Afro-Asiatic 0 1 Dravidian 0 0 Indo-European 11 2 Japonic 0 0 Koreanic 0 0 Mongolic 0 0 Sino-Tibetan 0 0 Turkic 0 1 Uralic 1 0	Abkhaz-Adyge 0 1 0 Afro-Asiatic 0 1 0 Dravidian 0 0 3 Indo-European 11 2 1 Japonic 0 0 0 Koreanic 0 0 0 Mongolic 0 0 0 Sino-Tibetan 0 0 0 Turkic 0 1 0 Uralic 1 0 0	Abkhaz-Adyge 0 1 0 Afro-Asiatic 0 1 0 Dravidian 0 0 3 Indo-European 11 2 1 Japonic 0 0 0 Koreanic 0 0 0 Mongolic 0 0 0 Sino-Tibetan 0 0 0 Uralic 1 0 0	Abkhaz-Adyge 0 1 0 0 Afro-Asiatic 0 1 0 0 Dravidian 0 0 3 0 Indo-European 11 2 1 5 Japonic 0 0 0 0 Koreanic 0 0 0 0 Mongolic 0 0 0 1 Sino-Tibetan 0 0 0 0 Turkic 0 1 0 5 Uralic 1 0 0 4	Afro-Asiatic 0 1 0 0 Dravidian 0 0 3 0 Indo-European 11 2 1 5 Japonic 0 0 0 0 Koreanic 0 0 0 0 Mongolic 0 0 0 1 Sino-Tibetan 0 0 0 0 Turkic 0 1 0 5 Uralic 1 0 0 4

Abkhaz-Adyge 0 Afro-Asiatic ## 0 ## Dravidian 0 ## Indo-European 0 ## Japonic 0 Koreanic 0 ## 0 Mongolic ## ## Sino-Tibetan 1 Turkic ## 0 ## Uralic

LMER models

Mixed effects model, predicting Linguistic similaritys from cultural similarity, with random intercept for family and area and random slope for cultural similarity for family and area.

We start with a null model with random intercepts for family and area, and random slopes for cultural similarity by both. We add a fixed effect of the number of comparisons made for each datapoint (number of concepts that were available to compare). Then we add a fixed effect of cultural similarity

```
m0 = lmer(
  rho.center ~ 1 +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling
## boundary (singular) fit: see ?isSingular
m0.5 = lmer(
 rho.center ~ 1 +
    comparison_count.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling
)
## boundary (singular) fit: see ?isSingular
m1 = lmer(
 rho.center ~ 1 +
    comparison_count.center +
    cult.dist.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling
## boundary (singular) fit: see ?isSingular
an1 = anova(m0, m0.5, m1)
## refitting model(s) with ML (instead of REML)
an1
## Data: ling
## Models:
## m0: rho.center ~ 1 + (1 + cult.dist.center | family.group) + (1 +
           cult.dist.center | area.group)
## m0.5: rho.center ~ 1 + comparison_count.center + (1 + cult.dist.center |
## m0.5:
            family.group) + (1 + cult.dist.center | area.group)
## m1: rho.center ~ 1 + comparison_count.center + cult.dist.center +
           (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
## m1:
## m1:
           area.group)
##
       Df
             AIC
                    BIC logLik deviance
                                            Chisq Chi Df Pr(>Chisq)
        8 1654.6 1691.3 -819.30
## mO
                                 1638.6
## m0.5 9 1293.0 1334.3 -637.50
                                  1275.0 363.597
                                                       1 < 2.2e-16 ***
                                                       1 4.704e-05 ***
## m1
       10 1278.4 1324.4 -629.22 1258.4 16.564
## ---
```

```
Cultural similarity is not significantly correlated with Linguistic similarity. Here are the model estimates:
summary(m1)
## Linear mixed model fit by REML ['lmerMod']
## Formula: rho.center ~ 1 + comparison_count.center + cult.dist.center +
       (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
##
       area.group)
##
      Data: ling
##
## REML criterion at convergence: 1271.9
##
## Scaled residuals:
       Min 1Q Median
                                3Q
                                        Max
## -2.6249 -0.6167 0.1103 0.6571 4.7402
##
## Random effects:
## Groups
                                   Variance Std.Dev. Corr
##
   family.group (Intercept)
                                  0.1612630 0.40158
##
                 cult.dist.center 0.0001817 0.01348
##
                 (Intercept)
                                  0.0510850 0.22602
   area.group
##
                 cult.dist.center 0.0036658 0.06055 -1.00
##
                                   0.2885416 0.53716
  Residual
## Number of obs: 731, groups: family.group, 48; area.group, 20
##
## Fixed effects:
##
                           Estimate Std. Error t value
## (Intercept)
                           -0.39270 0.09073 -4.328
## comparison_count.center 0.61196
                                       0.02688 22.770
## cult.dist.center
                            0.19678
                                       0.03275
                                                 6.008
##
## Correlation of Fixed Effects:
##
               (Intr) cmpr_.
## cmprsn_cnt. 0.090
## clt.dst.cnt -0.194 -0.201
## convergence code: 0
## boundary (singular) fit: see ?isSingular
Plot the estimates, rescaling the variables back to the original units:
trans = function(X){
 X * attr(ling$rho.center,"scaled:scale") +
  attr(ling$rho.center,"scaled:center")
}
gx = plot_model(m1,'pred',terms='cult.dist.center')
gx$data$predicted = trans(gx$data$predicted)
gx$data$conf.low = trans(gx$data$conf.low)
gx$data$conf.high = trans(gx$data$conf.high)
gx$data$x = gx$data$x *
  cdc.s +cdc.c
gx = gx + \#coord\_cartesian(ylim=c(0, 0.5),
                           xlim=c(0.15,0.85)) +
```

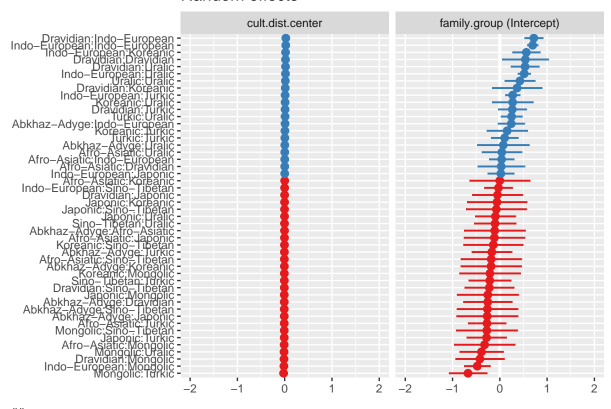
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

xlab("Cultural similarity") +

```
ylab("Semantic alignment") +
  ggtitle("") +
  geom_point(data=ling,aes(x=cult.dist,y=local_alignment))
gx
   0.5 -
   0.4 -
Semantic alignment
   0.1 -
   0.0 -
          0.2
                                      0.4
                                                                  0.6
                                                                                              0.8
                                           Cultural similarity
pdf(paste0("../results/stats/",datasetName,"/CulturalDistance_Rho_Graph.pdf"),
    height=2.5, width=2.5)
gx
dev.off()
## pdf
##
Plot the random effects:
plot_model(m1,'re', sort.est = "cult.dist.center")
```

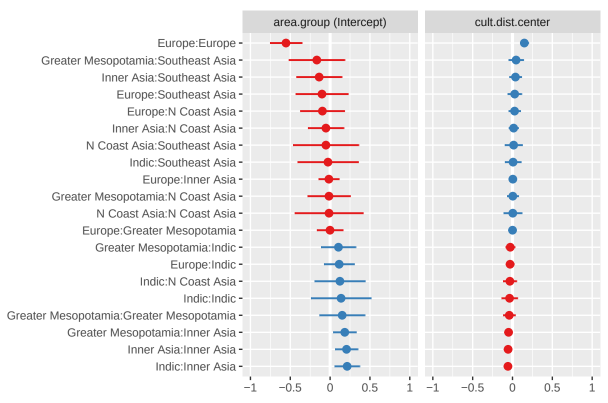
[[1]]

Random effects



[[2]]

Random effects



Without Kinship data

The analyses below show that the strongest relationship is with Kinship. Here we run the analysis as above, but using semantic distances computed without concepts that relate to kinship. Note that the local alignment values correlate with r > 0.99.

Code for constructing the data is hidden, but it is the same as above and available in the Rmd file:

Run the lmer models:

```
mONK = lmer(
  rho.center ~ 1 +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = lingNK
## boundary (singular) fit: see ?isSingular
m0.5NK = lmer(
 rho.center ~ 1 +
   comparison_count.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = lingNK
## boundary (singular) fit: see ?isSingular
m1NK = lmer(
 rho.center ~ 1 +
   comparison_count.center +
   cult.dist.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = lingNK
## boundary (singular) fit: see ?isSingular
anova(mONK,mO.5NK,m1NK)
## refitting model(s) with ML (instead of REML)
## Data: lingNK
## Models:
## mONK: rho.center ~ 1 + (1 + cult.dist.center | family.group) + (1 +
            cult.dist.center | area.group)
## m0.5NK: rho.center ~ 1 + comparison_count.center + (1 + cult.dist.center |
              family.group) + (1 + cult.dist.center | area.group)
## mO.5NK:
## m1NK: rho.center ~ 1 + comparison_count.center + cult.dist.center +
            (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
## m1NK:
## m1NK:
             area.group)
##
         Df
              AIC
                      BIC logLik deviance
                                              Chisq Chi Df Pr(>Chisq)
## mONK
          8 1654.6 1691.3 -819.30
                                    1638.6
## m0.5NK 9 1293.0 1334.3 -637.50
                                    1275.0 363.597
                                                         1 < 2.2e-16 ***
## m1NK
        10 1278.4 1324.4 -629.22
                                   1258.4 16.564
                                                         1 4.704e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

summary(m1NK)

```
## Linear mixed model fit by REML ['lmerMod']
## Formula: rho.center ~ 1 + comparison_count.center + cult.dist.center +
##
      (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
##
      area.group)
##
     Data: lingNK
##
## REML criterion at convergence: 1271.9
##
## Scaled residuals:
##
      Min
           1Q Median
                               ЗQ
## -2.6249 -0.6167 0.1103 0.6571 4.7402
## Random effects:
## Groups
                                 Variance Std.Dev. Corr
               Name
                                0.1612630 0.40158
## family.group (Intercept)
                cult.dist.center 0.0001817 0.01348 1.00
##
##
                (Intercept)
                             0.0510850 0.22602
  area.group
##
                cult.dist.center 0.0036658 0.06055
## Residual
                                 0.2885416 0.53716
## Number of obs: 731, groups: family.group, 48; area.group, 20
## Fixed effects:
##
                          Estimate Std. Error t value
## (Intercept)
                          -0.39270 0.09073 -4.328
## comparison_count.center 0.61196 0.02688 22.770
## cult.dist.center
                          0.19678 0.03275
##
## Correlation of Fixed Effects:
##
              (Intr) cmpr_.
## cmprsn_cnt. 0.090
## clt.dst.cnt -0.194 -0.201
## convergence code: 0
## boundary (singular) fit: see ?isSingular
```

MRM

Use multiple regression on distance matrices (Lichstein, 2007) to do the same test as above. The code below loads the data into a matrix format:

```
# Use graph method to make distance matrix
grph <- graph.data.frame(ling[,c("11",'12','local_alignment')], directed=FALSE)</pre>
# add value as a weight attribute
ling.m = get.adjacency(grph, attr="local_alignment", sparse=FALSE)
rownames(ling.m) = 1[match(rownames(ling.m), 1$iso2), ]$Language2
colnames(ling.m) = 1[match(colnames(ling.m), 1$iso2), ]$Language2
# Same for comparison_count.center
grph <- graph.data.frame(ling[,c("11",'12','comparison_count')], directed=FALSE)</pre>
# add value as a weight attribute
cc.m = get.adjacency(grph, attr="comparison_count", sparse=FALSE)
rownames(cc.m) = 1[match(rownames(cc.m),1$iso2),]$Language2
colnames(cc.m) = 1[match(colnames(cc.m),1$iso2),]$Language2
cult.m = read.csv("../results/EA distances/CulturalDistances.csv", stringsAsFactors = F)
rownames(cult.m) = cult.m[,1]
cult.m = cult.m[,2:ncol(cult.m)]
cult.m = as.matrix(cult.m)
# Flip cultural value to distance
cult.m = 1-cult.m
mx = match(rownames(ling.m),rownames(cult.m))
cult.m = cult.m[mx,mx]
colnames(cult.m) = rownames(cult.m)
# Same/different matrix for language family
family.matrix = l[match(rownames(ling.m), l$Language), ]$family
family.matrix = outer(family.matrix,family.matrix,"!=") *1
# Load ASJP distances for second test
asjp = readRDS("../data/ASJP/asjp17-dists FAIR.RData")
ling.m.glotto = 1[match(rownames(cult.m), 1$Language2), ]$glotto
ling.m.glotto = ling.m.glotto[ling.m.glotto %in% rownames(asjp)]
asjp.m = asjp[ling.m.glotto,ling.m.glotto]
asjp.lang.names = 1[match(rownames(asjp.m), 1$glotto), ]$Language2
# Matrices for second analysis with asjp
ling.m2 = ling.m[asjp.lang.names,asjp.lang.names]
cult.m2 = cult.m[asjp.lang.names,asjp.lang.names]
cc.m2 = cc.m[asjp.lang.names,asjp.lang.names]
# Load the geographic distances:
geoDist = read.csv("../data/GeographicDistances.csv",stringsAsFactors = F)
geoDist.m = as.matrix(geoDist)
geoDist.m = geoDist.m[!is.na(geoDist.m[,1]),!is.na(geoDist.m[1,])]
# Convert to log distance in thousand km
geoDist.m = log10(geoDist.m/1000)
geoDist.m[is.infinite(geoDist.m)] = 0
colnames(geoDist.m) = gsub("\\."," ",colnames(geoDist.m))
rownames(geoDist.m) = colnames(geoDist.m)
geoDist.m1 = geoDist.m[rownames(ling.m),rownames(ling.m)]
geoDist.m2 = geoDist.m[rownames(ling.m2),rownames(ling.m2)]
```

```
# center and scale values
ling.m = matrix(scale(as.vector(cc.m)),nrow=nrow(ling.m))
cc.m = matrix(scale(as.vector(cc.m)),nrow=nrow(cc.m))
cult.m = matrix(scale(as.vector(cult.m)),nrow=nrow(cult.m))
geoDist.m1 = matrix(scale(as.vector(geoDist.m1)),nrow=nrow(geoDist.m1))

asjp.m = matrix(scale(as.vector(asjp.m)),nrow=nrow(asjp.m))
ling.m2 = matrix(scale(as.vector(ling.m2)),nrow=nrow(ling.m2))
cc.m2 = matrix(scale(as.vector(cc.m2)),nrow=nrow(cc.m2))
cult.m2 = matrix(scale(as.vector(cult.m2)),nrow=nrow(cult.m2))
geoDist.m2 = matrix(scale(as.vector(geoDist.m2)),nrow=nrow(geoDist.m2))
```

Run the MRM model, predicting semantic alignment by cultural distance, controlling for family distance, geographic ditance, and the comparison count (number of observations). Here, the family distance between two languages is just whether they are part of the same family. Note that this does not take into account particular values for particular families, nor the random slopes within families.

```
## $coef
##
                           as.dist(ling.m)
## Int
                                0.22628455 0.0452
## as.dist(cult.m)
                                0.27056532 0.0117
                               -0.22892903 0.1053
## as.dist(family.matrix)
## as.dist(geoDist.m1)
                                0.00141692 0.9863
## as.dist(cc.m)
                                0.60941424 0.0001
##
## $r.squared
##
          R2
                  pval
## 0.6104647 0.0001000
##
## $F.test
##
          F
              F.pval
## 288.3577
              0.0001
```

Semantic alignment is significantly correlated with cultural distance.

In the result above, geographic distance is not correlated with semantic distance. Geographic distance turns out to be moderately correlated with cultural distance:

```
## $coef
                                           pval
##
                    as.dist(geoDist.m1)
## Int
                            -0.02111026 0.7707
                            -0.50170516 0.0001
## as.dist(cult.m)
##
## $r.squared
##
          R2
                  pval
## 0.1529605 0.0001000
##
## $F.test
##
          F
              F.pval
## 133.4504
              0.0001
```

Even when testing for non-linear geographic effects, the main result still holds:

```
ecodist::MRM(as.dist(ling.m) ~
    as.dist(cult.m) +
    as.dist(family.matrix) +
    as.dist(geoDist.m1) +
    as.dist(geoDist.m1) +
```

\$F.test ## F F.pval ## 191.9400 0.0001

Below, we run the same test, but using average string distances in basic vocabulary from the ASJP (Wichmann, Holman & Brown, 2018) as controls for history. We used the distances as calculated in Jäger (2018), which used them to construct historical phylogenies.

```
ecodist::MRM(as.dist(ling.m2) ~
    as.dist(cult.m2) +
    as.dist(asjp.m) +
    as.dist(geoDist.m2) +
    as.dist(cc.m2),nperm = 10000)
```

```
## $coef
##
                        as.dist(ling.m2)
                                           pval
## Int
                              0.09389590 0.0004
## as.dist(cult.m2)
                              0.26068360 0.0226
## as.dist(asjp.m)
                             -0.25717809 0.0001
## as.dist(geoDist.m2)
                              0.03920662 0.6016
## as.dist(cc.m2)
                              0.61582319 0.0001
##
## $r.squared
##
          R2
                  pval
## 0.6251321 0.0001000
##
## $F.test
##
          F
              F.pval
              0.0001
## 260.5635
```

Mantel tests

Read the historical distances for Indo-European, based on the phylogenetic distances.

Data prep

The geographic distances are loaded above (from "../data/GeographicDistances.csv").

Load historical distances:

```
hist = read.csv("../data/trees/IndoEuropean_historical_distances.csv", stringsAsFactors = F)
hist = hist[!duplicated(hist[,1]),!duplicated(hist[,1])]
rownames(hist) = hist[,1]
hist = hist[,2:ncol(hist)]
hist.m = as.matrix(hist)
colnames(hist.m) = rownames(hist.m)
hist.m = hist.m/max(hist.m)
```

Read the cultural distance as a matrix:

```
cult.m = read.csv("../results/EA_distances/CulturalDistances.csv", stringsAsFactors = F)
rownames(cult.m) = cult.m[,1]
cult.m = cult.m[,2:ncol(cult.m)]
```

Flip the cultural distance into a cultural similarity measure:

```
cult.m = 1-cult.m
```

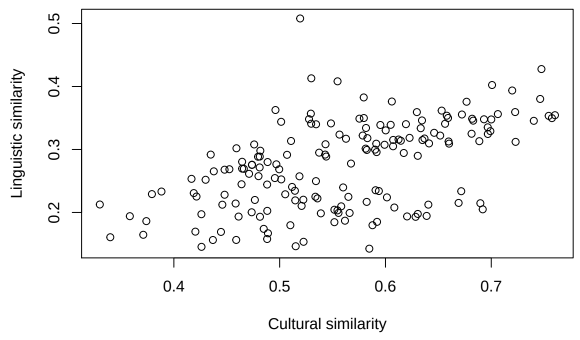
Convert the linguistic similarities to a matrix. This uses igraph to make an undirected graph from the long format with local_alignment as the edge weights, then output a matrix of adjacencies.

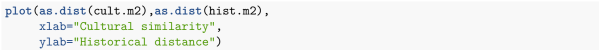
```
grph <- graph.data.frame(ling[,c("l1",'l2','local_alignment')], directed=FALSE)
# add value as a weight attribute
ling.m = get.adjacency(grph, attr="local_alignment", sparse=FALSE)
rownames(ling.m) = l[match(rownames(ling.m),l$iso2),]$Language2
colnames(ling.m) = l[match(colnames(ling.m),l$iso2),]$Language2</pre>
```

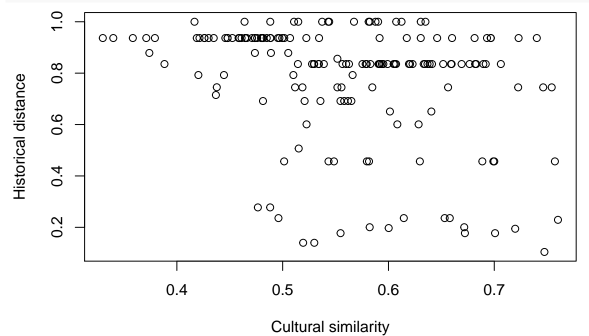
Match the distance matrices

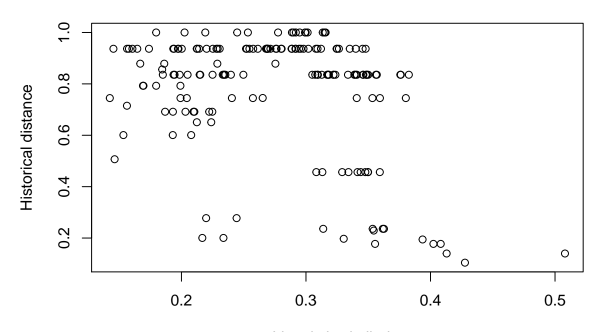
```
in.analysis = intersect(rownames(ling.m),rownames(cult.m))
in.analysis = intersect(in.analysis, rownames(hist.m))
cult.m2 = cult.m[in.analysis,in.analysis]
ling.m2 = ling.m[in.analysis,in.analysis]
hist.m2 = hist.m[in.analysis,in.analysis]
geo.m2 = geoDist.m[in.analysis,in.analysis]
```

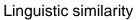
Note that there are only 19 languages with data on linguistic, cultural and historical distance.

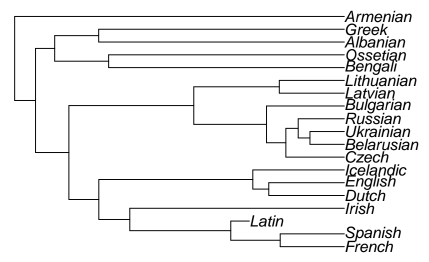












Tests

The results of the test list the following measures:

```
mantelr: Mantel correlation coefficient.
pval1: one-tailed p-value (null hypothesis: r <= 0).</li>
pval2: one-tailed p-value (null hypothesis: r >= 0).
pval3: two-tailed p-value (null hypothesis: r = 0).
llim: lower confidence limit for r.
```

• ulim: upper confidence limit for r.

```
set.seed(1498)
```

Run tests between each pair of measures.

```
distms = list("Cultrual"= cult.m2,
              "Linguistic" = ling.m2,
              "Historical" = hist.m2,
              "Geographic" = geo.m2)
for(i in 1:3){
  for(j in (i+1):4){
    var1 = names(distms)[i]
    var2 = names(distms)[j]
    print(paste("Correlation between",
                var1, "and", var2))
    stat = ecodist::mantel(as.dist(distms[[i]]) ~
                as.dist(distms[[j]]),
                nperm = 100000)
    print(stat)
    stat = round(stat,2)
    pval = round(min(c(stat[2],stat[3])),3)
    if(pval==0){pval = "$<$ 0.001"}</pre>
    stat2 = sprintf("$r$ = %s, 95\\%% CI = [%s,%s], one-tailed $p$ = %s",
      round(stat[1],3),
      round(stat[5],3),
      round(stat[6],3),
      pval)
    stat2 = gsub("0\\.",".",stat2)
    cat(stat2,file=
          paste0("../results/stats/tex/Mantel",var1,"Vs",var2,"Distance.tex"))
 }
}
```

```
## [1] "Correlation between Cultrual and Linguistic"
##
                          pval2
                                    pval3 llim.2.5% ulim.97.5%
     mantelr
                pval1
## 0.5243289 0.0050000 0.9950100 0.0050300 0.3796035 0.6586819
## [1] "Correlation between Cultrual and Historical"
##
                                    pval3 llim.2.5% ulim.97.5%
     mantelr
                pval1
                          pval2
## [1] "Correlation between Cultrual and Geographic"
                pval1
                          pval2
                                    pval3 llim.2.5% ulim.97.5%
##
     mantelr
## -0.4495398 0.9967200 0.0032900 0.0032900 -0.5754918 -0.3109193
## [1] "Correlation between Linguistic and Historical"
                pval1
                          pval2
                                    pval3 llim.2.5% ulim.97.5%
## -0.3372882 0.9859600 0.0140500 0.0167300 -0.5019408 -0.1639425
## [1] "Correlation between Linguistic and Geographic"
```

```
## mantelr pval1 pval2 pval3 llim.2.5% ulim.97.5%
## -0.2594386 0.9182700 0.0817400 0.1195200 -0.3694719 -0.1840035
## [1] "Correlation between Historical and Geographic"
## mantelr pval1 pval2 pval3 llim.2.5% ulim.97.5%
## 0.4210629 0.0004100 0.9996000 0.0004100 0.3313578 0.5176683
```

Run a mantel test comparing the Linguistic alignment to the cultural similarity, controlling for the historical distance between languages:

```
## mantelr pval1 pval2 pval3 llim.2.5% ulim.97.5%
## 0.4659407 0.0100000 0.9900100 0.0107800 0.3408500 0.5938397
```

Main Test: Run a mantel test comparing the Linguistic alignment to the cultural similarity, controlling for the historical distance and geographic distance between languages:

```
mainMantel = ecodist::mantel(as.dist(ling.m2)~
                  as.dist(cult.m2) +
                  as.dist(hist.m2) +
                  as.dist(geo.m2),
                nperm = 100000)
mainMantel
                                         pval3 llim.2.5% ulim.97.5%
     mantelr
                   pval1
                              pval2
   0.4508309 0.0114200
                         0.9885900 0.0119500 0.2962271 0.5993660
mainMantel = round(mainMantel,2)
mainMantel2 = sprintf("$r$ = %s, 95\\%% CI = [%s,%s], one-tailed $p$ = %s",
        round(mainMantel[1],3),
        round(mainMantel[5],3),
        round(mainMantel[6],3),
        round(mainMantel[2],3)
     )
mainMantel2 = gsub("0\\.",".",mainMantel2)
cat(mainMantel2,
```

file="../results/stats/tex/MantelCultrualVsLinguisticDistance_Partial.tex")

MRM

Perform the main test, but using multiple regression on distance matrices (MRM).

```
set.seed(21889)
mainMRM = ecodist::MRM(as.dist(ling.m2)~
                  as.dist(cult.m2) +
                  as.dist(hist.m2) +
                  as.dist(geo.m2), nperm=10000)
mainMRM
## $coef
##
                    as.dist(ling.m2)
                                        pval
## Int
                         0.123122021 0.9147
## as.dist(cult.m2)
                         0.350192033 0.0108
## as.dist(hist.m2)
                         -0.059044640 0.1623
```

```
## as.dist(geo.m2) 0.008177519 0.8212
##
## $r.squared
##
       R2
                pval
## 0.3072419 0.0073000
##
## $F.test
## F.pval
## 24.68846 0.00730
mainMRM2 = sprintf("$\\beta= $\%s, $p=$\%s",
                 round(mainMRM$coef[2,1],2),
                 round(mainMRM$coef[2,2],2))
cat(mainMRM2,
 file="../results/stats/tex/MRMCultrualVsLinguisticDistance_Partial.tex")
```

Analysis of filtered data

The analyses in this section use local alignment values based on (a) data that passes the wikipedia filter, and (b) data that passes the semantic filter.

Wikipedia filter

```
ling.filtered = read.csv(
   "../data/FAIR/nel-wiki-k100-alignments-by-language-pair_Filtered.csv",
   stringsAsFactors = F)
```

Note that the semantic alignment for the filtered and unfiltered data are essentially exactly the same, but for fewer languages:

```
ling.filtered$unfiltered.rho =
  apply(ling.filtered[,
       c("iso2_l1","iso2_l2")],1,
  function(X){
    ling[(ling$11==X[1] & ling$12==X[2]) |
            (ling$11==X[2] & ling$12==X[1]),]$local_alignment[1]
  })
cor(ling.filtered$unfiltered.rho,ling.filtered$rho,use = "complete.obs")
```

[1] 0.9999918

Continue to build data for replication:

```
ling.filtered$area1 = 1[match(ling.filtered$name_11,1$Language),]$autotyp.area
ling.filtered$area2 = 1[match(ling.filtered$name_12,1$Language),]$autotyp.area
fgroup = cbind(ling.filtered$family1,ling.filtered$family2)
fgroup = apply(fgroup,1,sort)
ling.filtered$family.group = apply(fgroup,2,paste,collapse=":")
agroup = cbind(ling.filtered$area1,ling.filtered$area2)
agroup = apply(agroup,1,sort)
ling.filtered$area.group = apply(agroup,2,paste,collapse=":")
ling.filtered$rho.center = scale(ling.filtered$rho)
ling.filtered$comparison_count.center = scale(ling.filtered$comparison_count)
matches = sapply(1:nrow(ling.filtered), function(i){
 x = which((cult$11==ling.filtered$name_l1[i] &
          cult$12==ling.filtered$name_12[i]) |
            (cult$12==ling.filtered$name_l1[i] &
          cult$11==ling.filtered$name_12[i]))
 x[1]
})
ling.filtered$cult.dist = cult[matches,]$cult.dist
# flip
ling.filtered$cult.dist = 1 - ling.filtered$cult.dist
ling.filtered = ling.filtered[!is.na(ling.filtered$cult.dist),]
ling.filtered$cult.dist.center = scale(ling.filtered$cult.dist)
```

```
mOF = lmer(
  rho.center ~ 1 +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling.filtered
## boundary (singular) fit: see ?isSingular
m0.5F = lmer(
  rho.center ~ 1 +
    comparison_count.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling.filtered
## boundary (singular) fit: see ?isSingular
m1F = lmer(
  rho.center \sim 1 +
    comparison_count.center +
    cult.dist.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling.filtered
an1F = anova(mOF,mO.5F,m1F)
## refitting model(s) with ML (instead of REML)
an1F
## Data: ling.filtered
## Models:
## mOF: rho.center ~ 1 + (1 + cult.dist.center | family.group) + (1 +
           cult.dist.center | area.group)
## m0.5F: rho.center ~ 1 + comparison_count.center + (1 + cult.dist.center |
            family.group) + (1 + cult.dist.center | area.group)
## m1F: rho.center ~ 1 + comparison_count.center + cult.dist.center +
## m1F:
            (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
## m1F:
            area.group)
                                             Chisq Chi Df Pr(>Chisq)
##
        Df AIC BIC logLik deviance
         8 446.96 473.74 -215.48
                                    430.96
## m0.5F 9 428.63 458.76 -205.32
                                    410.63 20.3271
                                                         1 6.527e-06 ***
        10 425.68 459.15 -202.84
## m1F
                                    405.68 4.9519
                                                         1
                                                              0.02606 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Cultural similarity is significantly correlated with Linguistic similarity, even in the filtered data. Here are the
model estimates:
summary(m1F)
## Linear mixed model fit by REML ['lmerMod']
## Formula: rho.center ~ 1 + comparison_count.center + cult.dist.center +
##
       (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
##
       area.group)
```

```
##
     Data: ling.filtered
##
## REML criterion at convergence: 412.8
##
## Scaled residuals:
     Min 1Q Median
                            3Q
##
                                    Max
## -1.8211 -0.5105 -0.0662 0.4157 6.2698
## Random effects:
## Groups
               Name
                               Variance Std.Dev. Corr
## family.group (Intercept) 0.17449 0.4177
               cult.dist.center 0.07149 0.2674
##
                                                 -0.21
               (Intercept) 0.64616 0.8038
## area.group
                \verb|cult.dist.center| 0.02791 \quad 0.1670
##
                                                1.00
                               0.26481 0.5146
## Residual
## Number of obs: 210, groups: family.group, 31; area.group, 19
##
## Fixed effects:
##
                         Estimate Std. Error t value
## (Intercept)
                         -0.82996 0.21638 -3.836
## comparison_count.center 0.41070
                                 0.08668 4.738
## cult.dist.center
                        0.29681 0.11189 2.653
##
## Correlation of Fixed Effects:
      (Intr) cmpr_.
## cmprsn_cnt. 0.133
## clt.dst.cnt 0.387 -0.054
```

Semantic filter

```
ling.semFiltered = read.csv(
  "../data/FAIR/nel-wiki-k100-alignments-by-language-pair SemanticFiltered.csv",
  stringsAsFactors = F)
ling.semFiltered$area1 = 1[match(ling.semFiltered$name_11,1$Language),]$autotyp.area
ling.semFiltered$area2 = 1[match(ling.semFiltered$name_12,1$Language),]$autotyp.area
fgroup = cbind(ling.semFiltered$family1,ling.semFiltered$family2)
fgroup = apply(fgroup,1,sort)
ling.semFiltered$family.group = apply(fgroup,2,paste,collapse=":")
agroup = cbind(ling.semFiltered$area1,ling.semFiltered$area2)
agroup = apply(agroup,1,sort)
ling.semFiltered$area.group = apply(agroup,2,paste,collapse=":")
ling.semFiltered$rho.center = scale(ling.semFiltered$rho)
ling.semFiltered$comparison_count.center = scale(ling.semFiltered$comparison_count)
matches = sapply(1:nrow(ling.semFiltered), function(i){
  x = which((cult$11==ling.semFiltered$name_l1[i] &
          cult$12==ling.semFiltered$name_12[i]) |
            (cult$12==ling.semFiltered$name_11[i] &
          cult$11==ling.semFiltered$name_12[i]))
  x[1]
})
ling.semFiltered$cult.dist = cult[matches,]$cult.dist
ling.semFiltered$cult.dist = 1 - ling.semFiltered$cult.dist
ling.semFiltered = ling.semFiltered[!is.na(ling.semFiltered$cult.dist),]
ling.semFiltered$cult.dist.center = scale(ling.semFiltered$cult.dist)
mOF = lmer(
  rho.center ~ 1 +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling.semFiltered
)
## boundary (singular) fit: see ?isSingular
m0.5F = lmer(
  rho.center ~ 1 +
    comparison_count.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling.semFiltered
## boundary (singular) fit: see ?isSingular
m1F = lmer(
 rho.center ~ 1 +
```

```
comparison_count.center +
    cult.dist.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling.semFiltered
## boundary (singular) fit: see ?isSingular
an1F = anova(mOF, mO.5F, m1F)
## refitting model(s) with ML (instead of REML)
## Data: ling.semFiltered
## Models:
## mOF: rho.center ~ 1 + (1 + cult.dist.center | family.group) + (1 +
           cult.dist.center | area.group)
## m0.5F: rho.center ~ 1 + comparison_count.center + (1 + cult.dist.center |
            family.group) + (1 + cult.dist.center | area.group)
## m1F: rho.center ~ 1 + comparison count.center + cult.dist.center +
## m1F:
            (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
## m1F:
            area.group)
                                              Chisq Chi Df Pr(>Chisq)
##
        Df
               AIC
                      BIC logLik deviance
## mOF
         8 1660.3 1697.1 -822.17
                                    1644.3
## m0.5F 9 1300.8 1342.1 -641.39
                                     1282.8 361.576
                                                         1 < 2.2e-16 ***
## m1F
         10 1286.2 1332.1 -633.09
                                     1266.2 16.595
                                                         1 4.627e-05 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Cultural similarity is significantly correlated with Linguistic similarity, even in the semantic filtered data.
Here are the model estimates:
summary(m1F)
## Linear mixed model fit by REML ['lmerMod']
## Formula: rho.center ~ 1 + comparison_count.center + cult.dist.center +
##
       (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
##
       area.group)
      Data: ling.semFiltered
##
##
## REML criterion at convergence: 1279.6
##
## Scaled residuals:
##
       Min
                1Q Median
                                30
## -2.6257 -0.6155 0.1002 0.6528 4.7581
##
## Random effects:
##
                                  Variance Std.Dev. Corr
   Groups
                 Name
   family.group (Intercept)
                                  0.1627947 0.40348
##
                 cult.dist.center 0.0001944 0.01394
##
                 (Intercept)
                                  0.0515199 0.22698
   {\tt area.group}
                 cult.dist.center 0.0037519 0.06125
##
                                                     -1.00
                                  0.2916401 0.54004
## Residual
```

Number of obs: 731, groups: family.group, 48; area.group, 20

Both filters

Main test on data where both the wikipedia and semantic filter are on.

```
ling.bothFiltered = read.csv(
  "../data/FAIR/nel-wiki-k100-alignments-by-language-pair_BothFiltered.csv",
  stringsAsFactors = F)
ling.bothFiltered$area1 = 1[match(ling.bothFiltered$name_11,1$Language),]$autotyp.area
ling.bothFiltered$area2 = 1[match(ling.bothFiltered$name_12,1$Language),]$autotyp.area
fgroup = cbind(ling.bothFiltered$family1,ling.bothFiltered$family2)
fgroup = apply(fgroup,1,sort)
ling.bothFiltered$family.group = apply(fgroup,2,paste,collapse=":")
agroup = cbind(ling.bothFiltered$area1,ling.bothFiltered$area2)
agroup = apply(agroup,1,sort)
ling.bothFiltered$area.group = apply(agroup,2,paste,collapse=":")
ling.bothFiltered$rho.center = scale(ling.bothFiltered$rho)
ling.bothFiltered$comparison_count.center = scale(ling.bothFiltered$comparison_count)
matches = sapply(1:nrow(ling.bothFiltered), function(i){
  x = which((cult$11==ling.bothFiltered$name_l1[i] &
          cult$12==ling.bothFiltered$name_12[i]) |
            (cult$12==ling.bothFiltered$name_l1[i] &
          cult$11==ling.bothFiltered$name_12[i]))
 x[1]
})
ling.bothFiltered$cult.dist = cult[matches,]$cult.dist
# flip
ling.bothFiltered$cult.dist = 1 - ling.bothFiltered$cult.dist
ling.bothFiltered = ling.bothFiltered[!is.na(ling.bothFiltered$cult.dist),]
ling.bothFiltered$cult.dist.center = scale(ling.bothFiltered$cult.dist)
mOF = lmer(
  rho.center ~ 1 +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling.bothFiltered
## boundary (singular) fit: see ?isSingular
m0.5F = lmer(
  rho.center \sim 1 +
    comparison_count.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling.bothFiltered
```

boundary (singular) fit: see ?isSingular

```
m1F = lmer(
  rho.center ~ 1 +
    comparison_count.center +
    cult.dist.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling.bothFiltered
an1F = anova(m0F, m0.5F, m1F)
## refitting model(s) with ML (instead of REML)
an1F
## Data: ling.bothFiltered
## Models:
## mOF: rho.center ~ 1 + (1 + cult.dist.center | family.group) + (1 +
            cult.dist.center | area.group)
## m0.5F: rho.center ~ 1 + comparison_count.center + (1 + cult.dist.center |
             family.group) + (1 + cult.dist.center | area.group)
## mO.5F:
## m1F: rho.center ~ 1 + comparison_count.center + cult.dist.center +
## m1F:
            (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
## m1F:
            area.group)
##
         Df
              AIC
                      BIC logLik deviance
                                              Chisq Chi Df Pr(>Chisq)
## mOF
         8 447.24 474.02 -215.62
                                     431.24
## m0.5F 9 429.45 459.58 -205.73
                                     411.45 19.7903
                                                         1 8.642e-06 ***
         10 426.47 459.95 -203.24
                                     406.47 4.9762
## m1F
                                                         1
                                                               0.0257 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Cultural similarity is significantly correlated with Linguistic similarity, even in the fully filtered data. Here
are the model estimates:
summary(m1F)
## Linear mixed model fit by REML ['lmerMod']
## Formula: rho.center ~ 1 + comparison_count.center + cult.dist.center +
       (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
##
##
       area.group)
##
      Data: ling.bothFiltered
##
## REML criterion at convergence: 413.6
##
## Scaled residuals:
##
       Min
                1Q Median
                                3Q
## -1.8275 -0.5144 -0.0659 0.4117 6.2874
##
## Random effects:
##
    Groups
                 Name
                                  Variance Std.Dev. Corr
##
    family.group (Intercept)
                                  0.17288 0.4158
##
                 cult.dist.center 0.07138 0.2672
                                                     -0.21
##
                 (Intercept)
                                  0.64816 0.8051
    area.group
##
                 cult.dist.center 0.02802 0.1674
                                                     1.00
                                   0.26612 0.5159
## Residual
```

Number of obs: 210, groups: family.group, 31; area.group, 19

##

Comparison between domains

The code that produce the results of this section can be found in analysis/compareDomains.R.

Part 1: Compare each linguistic domain to the overall cultural similarity

We fit a mixed effects model to compare the linguistic similarity in a given domain to the overall cultural distance. The linguistic similarity for the given domain is the dependent variable. There are random intercepts for language family and area pairs, and random slopes for overall cultural similarity by language family and by area. The comparison_count variable is added as a fixed effect. This null model is compared to a model with an additional fixed effect for the overall cultural similarity.

There are 21 linguistic domains with enough data. All correlations are positive and 11 are significant at the 0.05 level (adjusted for multiple comparisons).

The full results are in the file:

 $../results/stats/wikipedia-main/Cor_LingAlignmentByDomains_vs_OverallCulturalSimilarity.csv\\ Summary:$

```
p1res
                                                                Adjusted p sig
##
                              Domain
                                           Beta
## 2
                      Food and drink 0.29039152 3.842274e-08 8.068775e-07
## 6
        Miscellaneous function words 0.31349670 9.672370e-08 2.031198e-06
## 9
                            The body 0.23183657 8.711593e-07 1.829434e-05
## 13
                             Animals 0.26483784 4.281952e-06 8.992099e-05
## 21
                                Time 0.26708073 3.341177e-05 7.016471e-04
## 3
          Agriculture and vegetation 0.21319270 4.954909e-05 1.040531e-03
## 16
                        Modern world 0.15392213 2.860946e-04 6.007988e-03
## 14
                  The physical world 0.15530592 6.771587e-04 1.422033e-02
## 11
                   Spatial relations 0.11188738 1.323355e-03 2.779045e-02
## 20
                             Kinship 0.25408132 1.332699e-03 2.798669e-02
## 7
               Clothing and grooming 0.16478921\ 2.245060e-03\ 4.714625e-02
## 10
                    Sense perception 0.11168260 2.806430e-03 5.893504e-02
## 15 Social and political relations 0.10210872 6.603817e-03 1.386802e-01
## 1
                           The house 0.10743767 1.485121e-02 3.118755e-01
## 18
                            Quantity 0.13804241 1.691864e-02 3.552914e-01
## 8
                 Speech and language 0.11367507 3.209804e-02 6.740588e-01
        Basic actions and technology 0.06996793 7.317704e-02 1.000000e+00
## 19
## 17
                           Cognition 0.06413968 9.294337e-02 1.000000e+00
## 12
                 Emotions and values 0.06324463 9.437249e-02 1.000000e+00
                          Possession 0.07833831 1.102507e-01 1.000000e+00
## 5
## 4
                              Motion 0.05544251 2.537090e-01 1.000000e+00
```

Part 2: Compare each linguistic domain to the cultural similarity of each original D-PLACE domain

The method is the same as for part 1, except the cultural distance for a particular cultural domain is used instead of the overall cultural distance.

The full results are in the file:

```
../results/stats/wikipedia-main/Cor\_LingAlignmentByDomains\_vs\_DPlaceCulturalDomains.csv
```

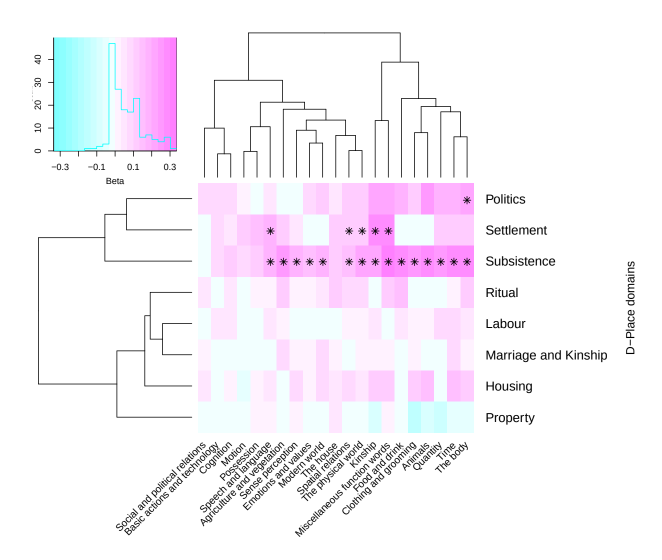
The graph below shows the mixed effects model coefficient estimate for the relationship between each linguistic domain and each cultural domain. Pink colours indicate positive correlations and blue colours indicate negative correlations. Stronger colours indicate stronger correlations. An asterisk indicates that the correlation is stronger than would be expected by chance, when adjusting the p-value for multiple comparisons.

The insert in the top left shows the distribution of Beta values.

The domains are clustered using higherarchical clustering. This is for visualisation and reflects similarity in the numeric relations, not history or conceptual hierarchies.

List of significant correlations (after adjusting p-value for multiple comparisons):

##		Ling Domain	Cult Domain	Beta	Adjusted p
##	68	The body	Politics	0.2139638	1.292033e-03
##	155	Kinship	Settlement	0.2785591	8.766279e-03
##	43	Miscellaneous function words	Settlement	0.2899807	7.022120e-07
##	83	Spatial relations	Settlement	0.1122428	1.431007e-03
##	59	Speech and language	Settlement	0.1915031	3.729177e-05
##	107	The physical world	Settlement	0.1321783	4.102261e-02
##	17	Agriculture and vegetation	Subsistence	0.2513632	2.887202e-05
##	97	Animals	Subsistence	0.2942266	7.865336e-06
##	49	Clothing and grooming	Subsistence	0.2484714	1.338589e-04
##	89	Emotions and values	${\tt Subsistence}$	0.1524638	7.946118e-04
##	9	Food and drink	${\tt Subsistence}$	0.3005301	1.084517e-06
##	153	Kinship	${\tt Subsistence}$	0.2346825	4.160376e-03
##	41	Miscellaneous function words	${\tt Subsistence}$	0.3353616	2.648483e-05
##	121	Modern world	${\tt Subsistence}$	0.1851981	7.434982e-06
##	137	Quantity	${\tt Subsistence}$	0.2454534	7.092645e-03
##	73	Sense perception	${\tt Subsistence}$	0.1850017	6.280926e-04
##	113	${\tt Social} \ {\tt and} \ {\tt political} \ {\tt relations}$	${\tt Subsistence}$	0.1544007	3.637901e-05
##	81	Spatial relations	${\tt Subsistence}$	0.1504141	6.897505e-04
##	57	Speech and language	${\tt Subsistence}$	0.1991926	2.699664e-04
##	65	The body	${\tt Subsistence}$	0.2764446	5.291731e-08
##	105	The physical world	${\tt Subsistence}$	0.2221189	1.157346e-04
##	161	Time	Subsistence	0.2921557	1.640226e-05



Concepticon domains

Figure 1:

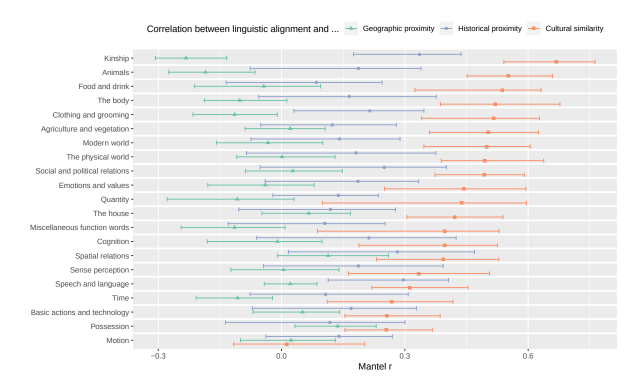


Figure 2:

Part 3: Compare each linguistic domain to the phylogenetic and geographic distance

This test compares each linguistic similarity scores to each of three target distances: the cultural distance, the historical distance and the geographic distance. We use a partial Mantel test (from the package ecodist) to estimate the strength of the relationship between the linguistic domain and the target distance, while controlling for the other two distances. The test uses 100,000 permutations.

The full results are in the file:

${\tt Cor_LingAlignmentByDomains_vs_HistoricalAndGeographicalDistance.csv}$

The graph below shows the results. Point estimates are the estimated Mantel R. The error bars show the 95% confidence intervals from the permutation test.

There appears to be a trade-off: The stronger the relationship with geographic distance, the weaker the relationship with cultural distance (r = -0.529, t = -2.72, df=19, p = 0.014). This does not hold for historical and cultural distance (r = 0.27, t = 1.22, df=19, p = 0.24).

Note that, after controlling for multiple comparisons, only 2 domains are significant:

```
## domain comparison mantelr lower upper pval3 p.adjusted
## 37 Animals lingVCult 0.5518312 0.4515039 0.6591382 0.00129 0.02709
## 58 Kinship lingVCult 0.6687835 0.5407906 0.7629987 0.00012 0.00252
```

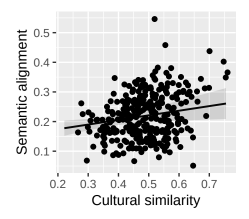


Figure 3: Semantic alignment and cultural similarity for data using the Common Crawl alignments

Summary of alternative data sources

The github repository also includes the main tests in this document, but applied to data where the alignments are done according to the Common Crawl or Subtitles databases.

Common Crawl

Mixed effects model: The correlation between semantic alignment and cultural similarity was significant (β = 0.182, $\chi^2(1)$ = 4.94, p=0.026). See figure 4.

MRM results:

	Estimate	p-value
Intercept	0.011	0.84
Cultural distance	0.086	0.172
Language family	0.001	0.987
Geographic distance	0.043	0.318
Comparison count	0.871	0.0001

Table 1: MRM analysis predicting semantic alignment (Common Crawl), with family control. $R^2=0.761$

	Estimate	p-value
Intercept	0.034	0.0311
Cultural distance	0.072	0.315
ASJP	-0.168	0.0009
Geographic distance	0.075	0.108
Comparison count	0.841	0.0001

Table 2: MRM analysis predicting semantic alignment (Common Crawl), with ASJP control. $R^2=0.744$

Mantel tests:

	Var1	Var2	r	llim	ulim	p
2	Cultrual	Linguistic	-0.084	-0.201	0.000885	0.277
3	Cultrual	Historical	-0.315	-0.447	-0.204	0.0211
4	Cultrual	Geographic	-0.461	-0.586	-0.312	0.00295
5	Linguistic	Historical	-0.25	-0.357	-0.114	0.015
6	Linguistic	Geographic	0.136	-0.0134	0.235	0.121
7	Historical	Geographic	0.405	0.31	0.519	0.00101
71	Linguistic	Cultural **	-0.0831	-0.152	-0.00152	0.27

Table 3: Mantel tests (Common Crawl). ** = partial Mantel test, controlling for historical and geographical distance.

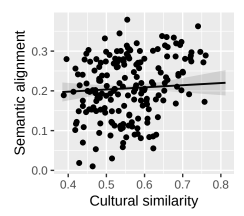


Figure 4: Semantic alignment and cultural similarity for data using the Subtitles alignments

Subtitles

Mixed effects model: The correlation between semantic alignment and cultural similarity was not significant (β = 0.0606, $\chi^2(1)$ = 0.74, p=0.39). See figure 5

MRM results:

	Estimate	p-value
Intercept	0.073	0.517
Cultural distance	0.125	0.413
Language family	-0.041	0.838
Geographic distance	-0.013	0.887
Comparison count	0.806	0.0002

Table 4: MRM analysis predicting semantic alignment (Subtitles), with family control. R^2 =0.744

Mantel tests:

	Estimate	p-value
Intercept	0.083	0.297
Cultural distance	-0.023	0.884
ASJP	-0.282	0.0083
Geographic distance	0.019	0.854
Comparison count	0.831	0.0003

Table 5: MRM analysis predicting semantic alignment (Subtitles), with ASJP control. R^2 =0.803

	Var1	Var2	r	$_{ m llim}$	$_{ m ulim}$	p
2	Cultrual	Linguistic	0.351	0.241	0.538	0.0884
3	Cultrual	Historical	-0.17	-0.286	-0.0153	0.155
4	Cultrual	Geographic	-0.34	-0.57	-0.172	0.0276
5	Linguistic	Historical	-0.352	-0.503	-0.0822	0.0301
6	Linguistic	Geographic	-0.273	-0.462	-0.0818	0.0806
7	Historical	Geographic	0.346	0.181	0.522	0.01
71	Linguistic	Cultural **	0.281	0.129	0.466	0.135

 $\label{eq:manuel} \text{Table 6: Mantel tests (Subtitles). *** = partial Mantel test, controlling for historical and geographical distance. }$

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4.2 Analysis of numerals

Semantic alignments in number words

Bill Thompson, Seán Roberts & Gary Lupyan

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Introduction

What predicts the semantic alignment of number words?

Although translations may be more direct for number words, there are still differences between languages regarding their semantic associations. Numerals in 16 languages were analysed using binary decision trees to find coherent clusters then a Generalised Additive Model to explain differences in alignment.

Variables:

- l1: Iso2 code for language 1
- l2: Iso2 code for language 2
- wordform_l1: Orthogaphic word form for language 1
- wordform_l2: Orthogaphic word form for language 2
- local_alignment: Local semantic alignment
- $\bullet \ global_alignment$: Global semantic alignment
- freq_l1: Frequency of orthographic form in l1
- freq_l2: Frequency of orthographic form in l1
- global_density_l1: Density
- $global_density_l2$: Density
- local_density_l1: Density

- local_density_l2: Density
- editdistance: Edit distance between orthographic forms
- k: Parameter for number of neighbours (constant at k=100).
- n: Number of possible comparisons
- number: Number concept
- number_numeric: Numeric value of number
- name 11: Name of language 11
- family_l1: Family of language l1
- name l2: Name of language l2
- family_l2: Family of language 12
- same.family: Compared languages are part of same family?
- hist.dist: Historical distance according to phylogenetic tree.
- l1_typology: Number line typology according to Calude & Verkerk (2016)
- *l2_typology*: Number line typology according to Calude & Verkerk (2016)
- sameNumeralTypology: Comparison of number line typology.
- seven: True if the numeral is 7. (see below)
- homophone: Word has a homophone in the North Euralex database.
- freqDiff: Difference in orthographic frequency (absolute, log scale)

A note on EDF values and random effects in Generalised Additive Models

(This note taken from Monaghan & Roberts (2019) Cognitive influences in language evolution: Psycholinguistic predictors of loan word borrowing. *Cognition*, 186, 147-158.)

The Estimated Degrees of Freedom (EDF) is an indication of how non-linear a smooth term is (higher = less linear). It is intended as a diagnostic measure of the shape of the curve, rather than a value used in estimating significance. The EDF is not the same as a simple polynomial curve's degree. Instead of a single polynomial curve, each smooth term is a collection of underlying basis functions (simpler curves). When each basis function is weighted by a coefficient, they add up to fit the data. The model attempts to find a collection of basis functions and weighting coefficients that add up to fit the data. Smaller collections of basis functions (simpler models) are preferred and larger (more complex) collections are penalised.

When the model converges on a solution, each smooth term is a collection of simpler curves. Each curve might have a polynomial degree, but it is also useful to have an estimate of the linearity/non-linearity of the whole smooth term. This is what the EDF provides: an EDF of 1 indicates a linear relationship, and higher values indicate more non-linear relationship. The definition of EDF in the implementation we use is described in Wood (2008):

"Associated with each smooth function is one or more measures of function 'wiggliness' $\beta_T^j \tilde{S}_j \beta_j$ where \tilde{S}_j is a matrix of known coefficients. Typically the wiggliness measure evaluates something like the univariate spline penalty."

That is, an EDF value is a combination of non-linearity measures of the basis functions, weighted by the weighting coefficient of each basis function. So, in general, a curve with an EDF of around 2 will look like a quadratic curve, and an EDF of around 3 will look like a cubic curve. However, this does not have to be the case: a smooth term could have a strong linear term, and a very weak non-linear term. The EDF captures this possibility as a continuous value. The simplest way to actually assess the smooth term is to plot it.

Random effects in the GAM implementation we use are treated just like a smooth term with the identity matrix as the penalty coefficient matrix. When entering a language pair as a random (intercept) effect, coefficients are created for each pair, modelled as independent and identically distributed normal random variables. The values are defined as discrete points along a smooth function. So, just like in a mixed effects model, the predicted alignment can be adjusted by a random intercept (the coefficients), e.g. the model can represent the alignment between English and French as higher overall, and the alignment between English and Bulgarian as being slightly lower etc. Stronger differences between levels of the random effect would need be represented by more complex functions, which would be penalised (similar to how a linear mixed effect

model penalises random effect coefficient estimates which deviate from a normal distribution). The EDF value for the random effects relates to the 'wiggliness' of these coefficients when plotted in a regular space. This makes the EDF difficult to interpret. A random effect where there were no differences between levels would have an EDF of 1 (a flat line), but it would also be 1 when there were consistent distances between each level. So a high EDF would indicate something like an imbalance in the distribution of coefficients, e.g. a range of values that does not fit a normal distribution. In fact, there are several language pairs with lower alignment (pairs from different language families), and few with very high alignment (possibly a ceiling effect).

Load libraries, graphing theme

```
library(mgcv) # for gam
library(lmtest) # for model comparison
source("GAM_derivaties.R") # for derivitives plot
library(lme4)
library(tidyverse)
#library(langcog)
library(boot)
library(ggplot2)
library(lazyeval)
library(data.table)
library(MuMIn)
library(REEMtree)
library(rpart)
library(rpart.plot)
library(gridExtra)
library(grid)
library(gridBase)
myThemeBasic =
  theme_bw()+
  theme(panel.grid.minor=element_blank(),
        panel.grid.major=element_blank(),
        panel.background=element_blank())+
  theme(axis.text.x=element_text(size=13),
        axis.text.y=element_text(size=13),
        axis.title.x=element_text(size=12),
        axis.title.y=element_text(size=12))+
  theme(legend.background = element_rect(fill="transparent"))+
  theme(legend.text = element text(size=13))+
  theme(legend.title = element_text(size=13))+
  theme(axis.title.y=element_text(vjust=0.9,size=20))+
  theme(axis.title.x=element_text(vjust=0.9,size=20))+
  theme(axis.title.y=element_text(vjust=0))+
  theme(axis.title.x=element_text(vjust=0))
```

Load data

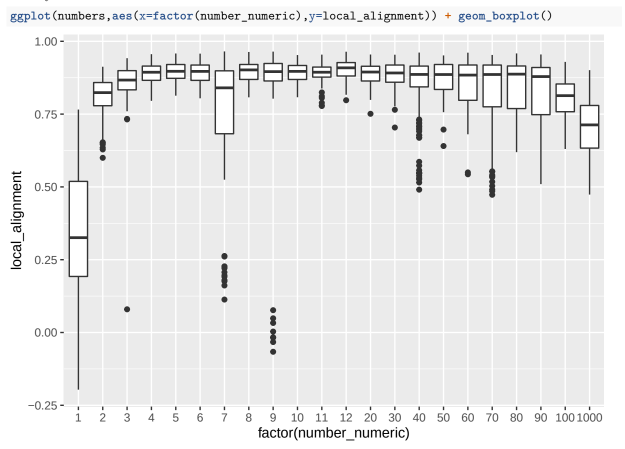
```
numbers <- numbers %>% rename(name_11=Language,family_11=family)
numbers <- left_join(numbers,select(language_info,Language,family,iso2),by=c("l2"="iso2"))</pre>
numbers <- numbers %>% rename(name_12=Language,family_12=family)
irreg_in_danish = c(50,60,70,80,90)
numbers <- numbers %>% mutate(
  same.family = family_11==family_12,
   is_danish = 11=="da" | 12=="da",
   is_single = (number_numeric < 10),</pre>
   is_decade = (number_numeric < 100 & number_numeric \\\% 10 ==0),
   is hundred = (number numeric == 100),
   is_thousand = (number_numeric == 1000),
    irreg_in_danish = (number_numeric %in% irreg_in_danish),
    lang_pair = paste(pmax(11,12),pmin(11,12),sep="-"))
histDistance = read.csv("../../data/trees/IndoEuropean_historical_distances_long.csv",
                        stringsAsFactors = F, encoding = "UTF-8", fileEncoding = "UTF-8")
numbers$hist.dist = histDistance[match(paste(numbers$name_11,numbers$name_12),
                             paste(histDistance$Var1,histDistance$Var2)),]$value
# Refactorise to get rid of non-existant categories
numbers$family_l1 = factor(numbers$family_l1)
numbers$family_12 = factor(numbers$family_12)
# Add typology data
cnv = read.csv("../../data/numbers/Calude_Verkerk_NumberData.csv",stringsAsFactors = F)
numbers <- left_join(numbers,cnv,by=c('l1','l2','number_numeric'))</pre>
numbers$isUralic = numbers$family_11 == "Uralic" | numbers$family_12 == "Uralic"
numbers$danish_irregular = (numbers$number_numeric %in% irreg_in_danish) &
                                     (numbers$name_l1=="Danish" | numbers$name_l2=="Danish")
# historic distance, setting Uralic languages to maximum
numbers$hist.dist2 = numbers$hist.dist
numbers$hist.dist2[is.na(numbers$hist.dist2)] = max(numbers$hist.dist,na.rm = T)
numbers$seven = numbers$number_numeric==7
#some rows got duplicated; remove
numbers <- distinct(numbers)</pre>
# Data on homophones
h = read.csv("../../data/numbers/NumberHomophones.csv",
             stringsAsFactors = F,encoding = "UTF-8",fileEncoding = "UTF-8")
h$code = paste(h$1,h$word)
numbers$homophone = (paste(numbers$11,numbers$wordform_11) %in% h$code) |
                    (paste(numbers$12,numbers$wordform_12) %in% h$code)
numbers$lang_pair.f = factor(numbers$lang_pair)
# Frequency difference (already in log scale)
```

```
numbers$freqDiff = abs(numbers$freq_11-numbers$freq_12)
# 78 frequency observations (3%) are missing, so impute:
freqM = bam(I(1+freqDiff)~
              #s(number_numeric) +
              s(lang_pair.f,bs='re') +
              s(editdistance) +
              s(hist.dist2),
            family = Gamma(link="identity"),
            data = numbers[!is.na(numbers$freqDiff),])
freqMPred = predict(freqM,newdata=numbers)-1
#plot(freqMPred, numbers$freqDiff)
#abline(0,1)
numbers[is.na(numbers$freqDiff),]$freqDiff =
 freqMPred[is.na(numbers$freqDiff)]
Group data by language:
langAverages = data.frame()
for(1 in unique(c(numbers$11,numbers$12))){
 dx = numbers[numbers$11==1 | numbers$12==1,]
  langAverages = rbind(langAverages,
                  data.frame(
                   local_alignment = dx$local_alignment,
                    number_numeric = dx$number_numeric,
                    1 = 1,
                    11 = dxname_11,
                    12 = dx name_12)
}
langAverages$1 = factor(langAverages$1,
    levels = names(sort(tapply(langAverages$local_alignment,langAverages$l,mean))))
```

Overview

Numeric value

Plot by numeric value:

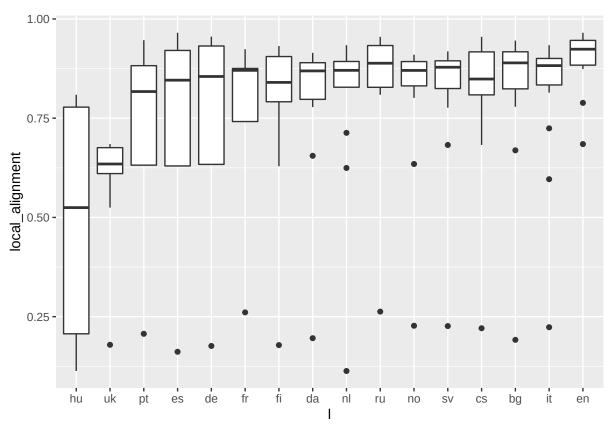


- 1 has a low numeric alignment.
- Alignment rises 2-3.
- 7 has a lower alignment.
- Drop in alignment for 100, 1000.
- Outliers for 7 and 9.
- Outliers for 40 and 70?

1 and 2 are often sources for grammaticalised indefinite/duel markers (Givón, 1981).

The slight decline from 10 to 1000 could be due to the declining frequency of occurances of these numbers (Dehaene & Mehler, 1992), which might affect convergence on meanings, but more directly would affect the co-occurance statistics.

What's driving the difference with 7 and 9? 7 might be linked to there being 7 days in the week, so semantic differences in time might be reflected. Let's look at individual languages:



The plot above shows that the main difference is being driven by comparisons with Hungarian. All the outliers around 0.25 are comparisons with Hungarian. This might be because Hungarian is a Uralic language, but maybe also because the Hungarian word for '7' also directly means "week".

Ukrainian is also low. We note that forms for 7 and 8 are very similar in Ukrainian ($7 = \sin and 8 = visim$).

What are the outliers for 9? These all include comparisons to French:

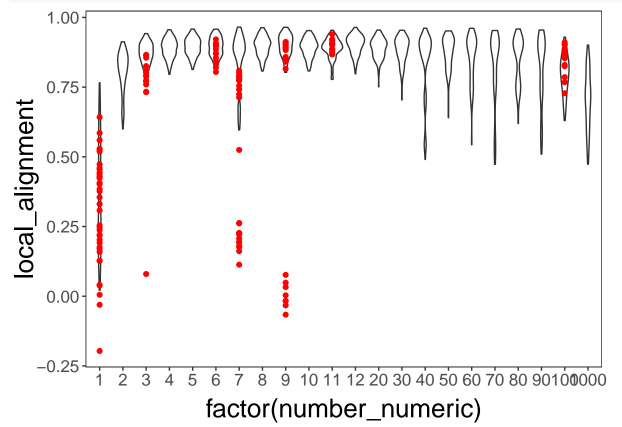
numbers[numbers\$number_numeric==9 & numbers\$local_alignment<0.25,c("11",'12',"local_alignment")]

```
##
        11 12 local_alignment
## 1
                  0.003192686
        fr uk
## 483
                  -0.016457496
        fr da
  786
                  0.076614888
##
        fr cs
## 903
        fr bg
                  -0.032815537
                  0.032734709
## 1231 fr fi
## 1366 fr no
                  -0.066181331
## 1436 fr sv
                  0.048704798
## 2401 fr ru
                  -0.017702025
```

This may be because French 9 ("neuf") can mean '9' or "new". We used the North Euralex dictionary to find numbers which have homophones. See the file NumberHomophones.csv. This also includes the Hungarian '7' discussed below.

h[,c("1","number","otherMeanings")]

The plot below shows the distribution of non-homophones, with homophones drawn as dots. For 7 and 9, these fall outside of the general distribution, but there are several other cases where homophones look similar to the rest of the distribution:

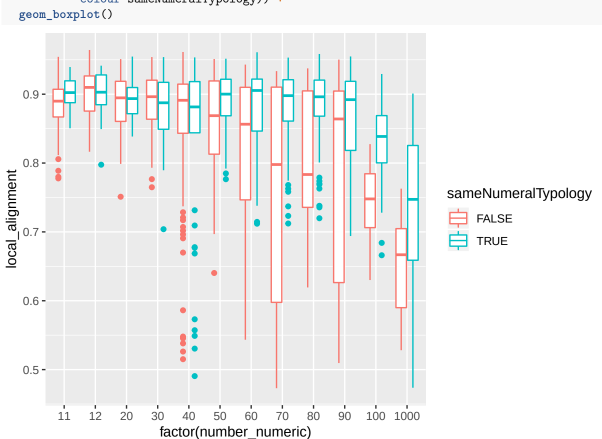


Number line

We can look at the importance of the different ways that numeral systems are compased. We use data from Calude & Verkerk (2016), and identify whether two languages have the same system for forming a particular numeral. Calude & Verkerk call this the 'number line'.

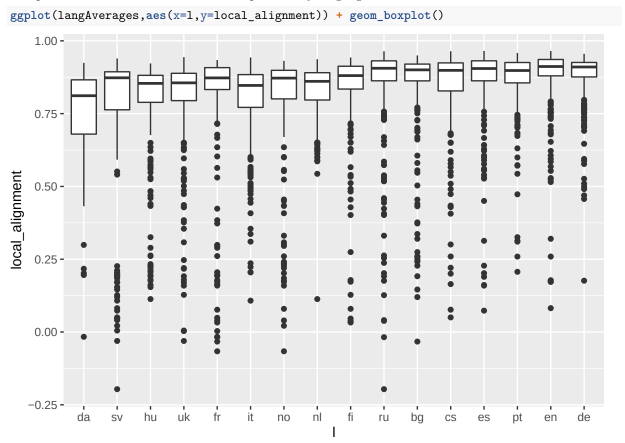
In our sample, all numbers below 10 are atoms, so there are only differences above 10. For example, English uses a unique atom to represent 12 (/twelv/), while Bulgarian uses a form that is composed of "2+10" (2 = dve, 10 = deset, 12 = dvanadeset).

It looks like there's an interaction between numeral typoogy and numeric value:

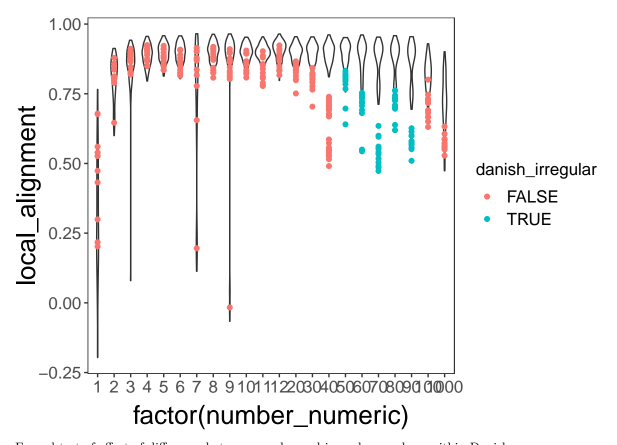


Variation by language

The plot below shows the overview of comparisons by language:



Danish seems to have a lower average. We note that some Danish 'crowns' are irregular (50,60,70,80,90). These pick out many outliers (dots are danish, blue dots are irregular, violin plots are the rest of the data):



Formal test of effect of difference between regular and irregular numbers within Danish:

```
summary(lm(local_alignment~irreg_in_danish, data=
 numbers[numbers$is_danish,]))
##
## Call:
## lm(formula = local_alignment ~ irreg_in_danish, data = numbers[numbers$is_danish,
##
       ])
##
## Residuals:
##
       Min
                  1Q
                       Median
                                    3Q
                                            Max
   -0.79901 -0.07008 0.04585 0.08937
##
## Coefficients:
##
                        Estimate Std. Error t value Pr(>|t|)
                                   0.008702 89.931 < 2e-16 ***
## (Intercept)
                        0.782552
                                   0.018609
                                            -6.442 4.5e-10 ***
## irreg_in_danishTRUE -0.119891
## ---
                  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
## Residual standard error: 0.1356 on 309 degrees of freedom
## Multiple R-squared: 0.1184, Adjusted R-squared: 0.1156
## F-statistic: 41.51 on 1 and 309 DF, p-value: 4.5e-10
```

Variation by frequency

Semantic alignment by frequency (brighter colours are higher numeric values):

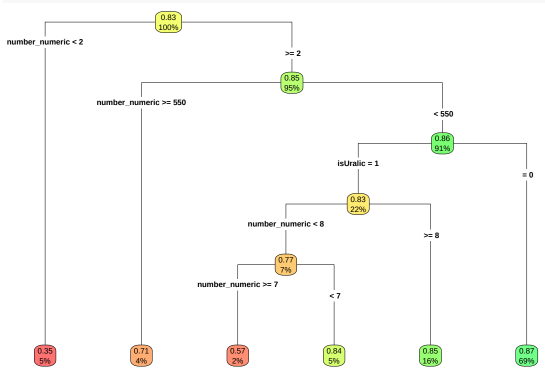
```
aes(x=freqDiff,y=local_alignment,colour=log10(number_numeric))) +
  geom_point() +
  stat_smooth() +
  scale_colour_gradient(name = "Numeral", trans = "log",
                        breaks = log10(c(1,5,10,30,100,1000)), labels =c(1,5,10,30,100,1000)) +
  myThemeBasic + xlab("Frequency difference") +
  ylab("Local alignment")
## Warning: Transformation introduced infinite values in discrete y-axis
## Warning: Transformation introduced infinite values in discrete y-axis
## `geom_smooth()` using method = 'gam' and formula 'y ~ s(x, bs = "cs")'
     1.00 -
     0.75
Local alignment
                                                                           Numeral
                                                                                1000
     0.50
                                                                                100
                                                                                30
                                                                                10
                                                                                5
     0.25
     0.00
   -0.25
                        0.5
                                     1.0
                                                   1.5
           0.0
                                                               2.0
                       Frequency difference
```

There appears to be a slight effect for larger frequency differences to be associated with lower alignment.

Decision tree

We use a decision tree to explore the data and find coherent clusters in the data. We try to predict local alignment based on various properties.

First we show that Uralic "7s" are a coherent cluster: A decision tree divides the data into categories that represent '1', '1000', and then combines divisions in the numeric value with 'isUralic' to find the cluster of Uralic 7s:



To help this, we include an explicit factor for seven. The final factors in the mode are:

- numeric value
- is a Uralic language
- is a Danish irregular
- is '7'
- has a homophone
- the 'n' variable: number of comparisons possible between language pairs in the whole corpus
- whether the numbers have the same underling compositional structure

```
data=numbers,
              MaxIterations=1000000)
rp.rt = tree(rt)
rp.rt$model = numbers
plot1 = rpart.plot(rp.rt, type=4, branch.lty=1, clip.facs = F, box.palette="RdYlGn")
cluster = factor(rp.rt$where,
                labels = c("One",
                            "Hungarian 7",
                            "French 9",
                            "Small\nhomophones",
                            "Large\nhomophones",
                            "100,1000",
                            "Danish\nirregulars",
                            "Two",
                            "3-90"))
plot2 = ggplot(numbers, aes(y=local_alignment,
                   x=cluster)) +
  geom_violin() + myThemeBasic +
  xlab("") + ylab("Semantic alginment")
pdf("rDecisionTree.pdf", width=12, height=8)
  layout(t(t(c(1,2))), heights=c(2.5,1))
  par(mar=c(1,10,1,1))
  rpart.plot(rp.rt, type=4, branch.lty=2,
             clip.facs = F, box.palette="RdYlGn",
             mar=c(1,4,1,1.5), cex = 1.2, split.yshift=1)
  plot.new()
  vps <- baseViewports()</pre>
  pushViewport(vps$figure)
  vp1 \leftarrow plotViewport(c(0,0,0,0))
  print(plot2, vp = vp1)
dev.off()
varimp = sort(rt$Tree$variable.importance)
varimp.plot = ggplot(data.frame(importance=varimp,
                  variable=factor(names(varimp),levels = names(varimp))),
       aes(y=importance,x=variable))+
  geom_col() + coord_flip()
pdf("../../results/numbers/VarImp.pdf")
varimp.plot
dev.off()
```

Run a GAM

Convert lang_pair to factor and scale variables:

```
numbers$lang_pair = factor(numbers$lang_pair)
numbers$number_numeric.log = log(numbers$number_numeric)
numbers$number_numeric.log.scaled = scale(numbers$number_numeric.log)
```

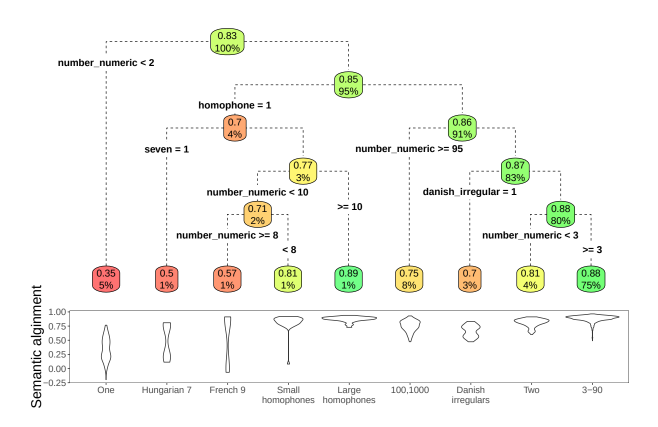


Figure 1:

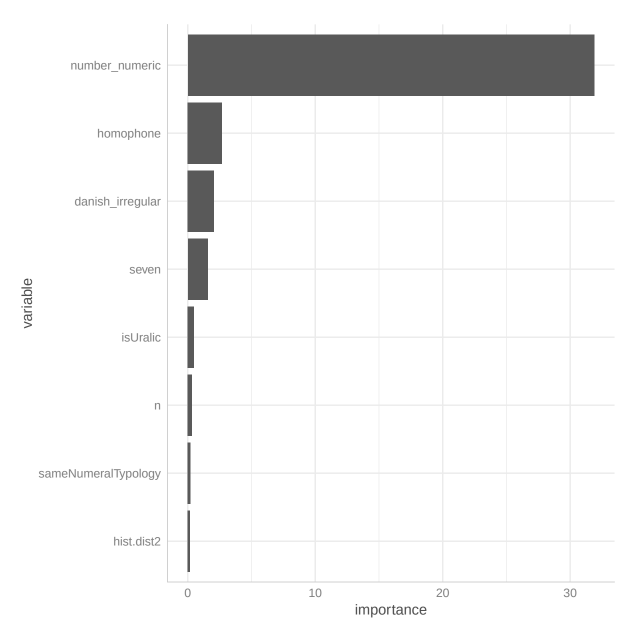


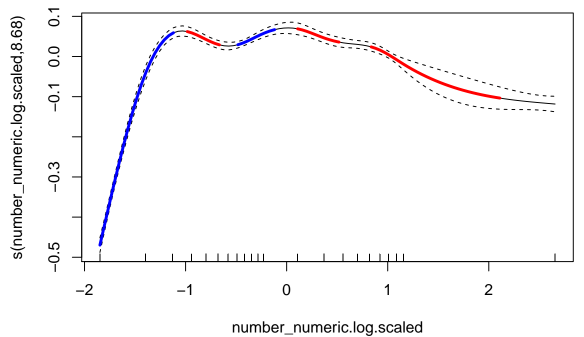
Figure 2:

```
numbers$local_alignment.center = scale(numbers$local_alignment,scale=F)

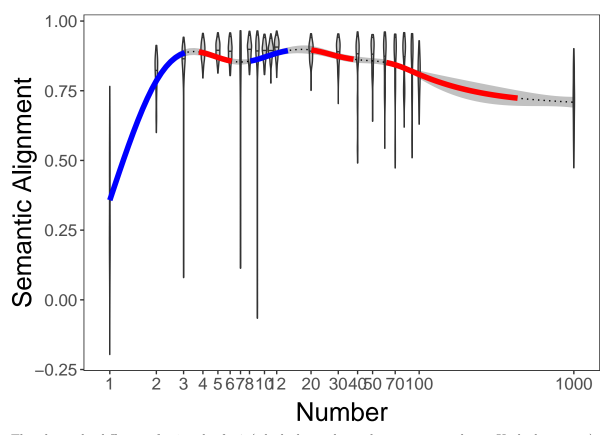
numbers$isUralic = factor(numbers$family_l1 == "Uralic" | numbers$family_l2 == "Uralic")
numbers$differentNumeralTypology = factor(!numbers$sameNumeralTypology)
numbers$seven = factor(numbers$seven)
numbers$danish_irregular = factor(numbers$danish_irregular)
numbers$homophone = factor(numbers$homophone)
```

We start by looking at a simple model that has a random effect for language pair and a main smooth term for number. Note that since this is a non-linear model, a random effect for numeric value is very similar to a fully articulated smooth slope, so we just model numeric as a fixed effect.

We plot the fit of the model below:



Warning: Removed 61 rows containing missing values (geom_path).
Warning: Removed 31 rows containing missing values (geom_path).



This shows the difference for 1, a dip for 7 (which the analyses above suggest is due to Uralic languages) and a decrease for 1000.

We now fit a full model with many other predictors:

- Number typology
- Interaction between numeric value and typology
- Historical distance (assuming Uralic is maximum distance)
- Whether the comparison is with a Uralic language
- Whether the number is 7
- Interaction between Uralic and 7
- Whether the number is a Danish irregular
- Whether the numbers word has a frequent homophone
- The frequency difference between the forms

Compare models. Adding the extra factors makes a difference.

```
lrtest(m0,m1)
## Likelihood ratio test
## Model 1: local_alignment.center ~ s(lang_pair, bs = "re") + s(number_numeric.log.scaled)
## Model 2: local_alignment.center ~ s(number_numeric.log.scaled, by = differentNumeralTypology) +
##
       s(hist.dist2) + s(lang_pair, bs = "re") + isUralic * seven +
##
       danish_irregular + differentNumeralTypology + homophone +
##
       s(hist.dist2) + s(freqDiff)
##
        #Df LogLik
                      Df Chisq Pr(>Chisq)
## 1 109.15 2199.7
## 2 117.38 2518.2 8.2261 637.08 < 2.2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
summary(m1)
##
## Family: gaussian
## Link function: identity
## Formula:
## local_alignment.center ~ s(number_numeric.log.scaled, by = differentNumeralTypology) +
       s(hist.dist2) + s(lang_pair, bs = "re") + isUralic * seven +
##
##
       danish_irregular + differentNumeralTypology + homophone +
##
       s(hist.dist2) + s(freqDiff)
##
## Parametric coefficients:
##
                                 Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                 0.017116 0.004897
                                                      3.495 0.000483 ***
                                 0.001801 0.009772
## isUralicTRUE
                                                     0.184 0.853816
                                          0.013742 -2.117 0.034367 *
## sevenTRUE
                                -0.029093
## danish_irregularTRUE
                                -0.144641
                                           0.013097 -11.044 < 2e-16 ***
## differentNumeralTypologyTRUE 0.024844
                                                      0.776 0.437883
                                           0.032019
## homophoneTRUE
                                -0.112477
                                           0.008755 -12.847 < 2e-16 ***
## isUralicTRUE:sevenTRUE
                                -0.191216
                                           0.020627 -9.270 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Approximate significance of smooth terms:
                                                                     Ref.df
                                                                 edf
## s(number_numeric.log.scaled):differentNumeralTypologyFALSE 8.598
                                                                       8.944
## s(number_numeric.log.scaled):differentNumeralTypologyTRUE
                                                               3.986
                                                                       4.514
## s(hist.dist2)
                                                               1.000
                                                                       1.000
## s(lang_pair)
                                                              92.082 117.000
## s(freqDiff)
                                                               1.000
                                                                       1.000
                                                                    F
## s(number_numeric.log.scaled):differentNumeralTypologyFALSE 305.027
## s(number_numeric.log.scaled):differentNumeralTypologyTRUE
                                                               45.087
## s(hist.dist2)
                                                                0.671
                                                                3.532
## s(lang_pair)
## s(freqDiff)
                                                               24,254
                                                               p-value
                                                              < 2e-16 ***
## s(number_numeric.log.scaled):differentNumeralTypologyFALSE
## s(number_numeric.log.scaled):differentNumeralTypologyTRUE
                                                               < 2e-16 ***
```

```
## s(hist.dist2)
## s(lang_pair)
## s(freqDiff)
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## R-sq.(adj) = 0.699 Deviance explained = 71.3%
## fREML = -2301.9 Scale est. = 0.0076339 n = 2415
```

Summary of main model

The final model explains 71.33% of the deviance, compared to 62.67% in the baseline model.

Parametric effects:

- Being an Uralic language is not strong predictor.
- The number 7 has lower alignment in general ($\beta = -0.0291$, p = 0.0344)
- Danish irregulars have lower alignment ($\beta = -0.145$, p = < 0.001))
- Overall, there is no difference for comparisons between numbers with different numeral typologies (β = 0.0248 , p = 0.4379)
- Alignment is lower for comparisons between words were at least one has a homophone ($\beta=$ -0.112 , p = < 0.001)
- Comparisons with Uralic sevens are significantly lower in alignment ($\beta = -0.191$, p = < 0.001)

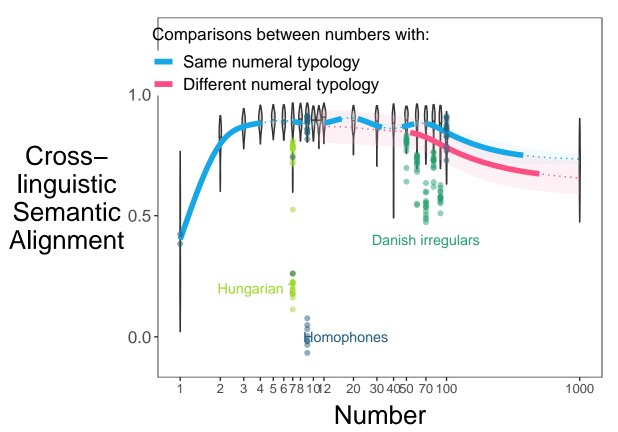
Smooth effects:

- The EDF for numeral is high, indicating strong non-linearities.
- There is an interaction between numeral value and numeral typology. The graph below shows that alignment is lower for comparisons between numbers with different typologies for numbers above 50. Also note that there is a slight increase in the semantic alignment for number 20 if the two languages have the same numeral typology.
- There was no effect of historical distance.
- There were significant random effects for particular language pairs.
- There was a significant linear effect of frequency difference: Alignment was lower when the frequency difference was larger.

The hidden code below (see the Rmd file) detects significantly steep slopes in the GAM curve. Thick line segments indicate significant rises or decreases. We see that 1 and 2 have lower alignment, then numbers 3-20 are fairly constant. Beyond 20, there is a decrease in alignment, especially for numbers with different numeral typologies. Various outliers captured by the model are drawn on top. Note that there are no numbers below 10 that have different numeral typologies, so we have truncated the curve accordingly.

gamPlot

```
## Warning: Removed 34 rows containing missing values (geom_path).
## Warning: Removed 141 rows containing missing values (geom_path).
## Warning: Removed 109 rows containing missing values (geom_path).
```



pdf("../../results/numbers/FinalGamModel.pdf", width=7.5,height=4.5)
gamPlot

```
## Warning: Removed 34 rows containing missing values (geom_path).
## Warning: Removed 141 rows containing missing values (geom_path).
## Warning: Removed 109 rows containing missing values (geom_path).
dev.off()
```

pdf ## 2

Controlling for linguistic history

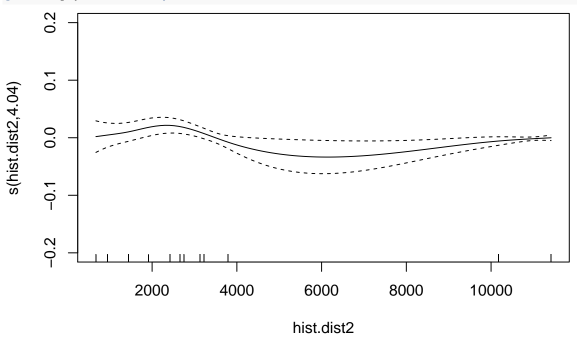
In the model above, the effects of historical distance are minimal: the fit is lienar and not significant. This might be because there are random effects for language pairs which are taking up the variance. In the section below, we use only historical distance:

```
m1.phylo = bam(local_alignment.center~
           s(number_numeric.log.scaled, by=differentNumeralTypology) +
           s(hist.dist2) +
           seven + danish_irregular +
           homophone +
           differentNumeralTypology +
           s(freqDiff),
         data = numbers[!is.na(numbers$hist.dist),])
# Model with numeric
summary(m1.phylo)
##
## Family: gaussian
## Link function: identity
##
## Formula:
## local_alignment.center ~ s(number_numeric.log.scaled, by = differentNumeralTypology) +
       s(hist.dist2) + seven + danish_irregular + homophone + differentNumeralTypology +
##
       s(freqDiff)
##
## Parametric coefficients:
##
                                 Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                                 0.017078 0.002922 5.845
                                                                6e-09 ***
## sevenTRUE
                                -0.036295
                                           0.014328 - 2.533
                                                             0.0114 *
## danish_irregularTRUE
                                -0.189497
                                           0.013823 -13.709
                                                             <2e-16 ***
## homophoneTRUE
                                -0.112117
                                            0.010836 -10.347
                                                               <2e-16 ***
## differentNumeralTypologyTRUE   0.048179
                                           0.072356
                                                      0.666
                                                              0.5056
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Approximate significance of smooth terms:
                                                                edf Ref.df
## s(number numeric.log.scaled):differentNumeralTypologyFALSE 8.535 8.927
## s(number_numeric.log.scaled):differentNumeralTypologyTRUE 4.697 5.104
## s(hist.dist2)
                                                              4.042 4.587
                                                              1.000 1.000
## s(freqDiff)
##
                                                                    F
## s(number_numeric.log.scaled):differentNumeralTypologyFALSE 215.732
## s(number_numeric.log.scaled):differentNumeralTypologyTRUE
## s(hist.dist2)
                                                                2.498
## s(freqDiff)
                                                               62.586
##
                                                               p-value
## s(number_numeric.log.scaled):differentNumeralTypologyFALSE
                                                               < 2e-16 ***
## s(number_numeric.log.scaled):differentNumeralTypologyTRUE
                                                               < 2e-16 ***
## s(hist.dist2)
                                                                0.0271 *
## s(freqDiff)
                                                              4.31e-15 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
```

```
## R-sq.(adj) = 0.674 Deviance explained = 67.8% ## fREML = -1731.5 Scale est. = 0.0081138 n = 1819
```

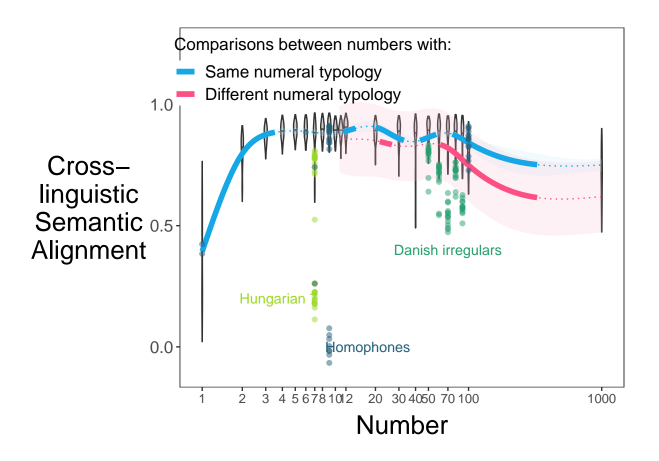
The effects are very similar. The effect of history is significant, but not large. Semantic alignment is lower for more distantly related languages:

plot(m1.phylo,select=3,ylim=c(-0.2,0.2))



Plot the main effects, which look much like the main graph above.

- ## Warning: Removed 34 rows containing missing values (geom_path).
- ## Warning: Removed 141 rows containing missing values (geom_path).
- ## Warning: Removed 119 rows containing missing values (geom_path).



Conclusion

Semantic alignment for number words is generally high, though there are some differences that can be explained (the model explained 71.33% of the deviance). 1 and 2 have lower alignment due to often being grammaticalised as indefinite or dual marker (Givon, 1981). Numbers 3-12 generally have high alignment (mean local alignment = 0.87), and higher numbers decline in alignment up to 1000. There are also language-specific differences due to how number lines are constructed (e.g. base, combination rules, see Calude & Verkerk, 2016), or for irregular forms (e.g. 50, 60, 70, 80 and 90 in Danish). Some number words have alternative associations due to homophones (e.g. the Hungarian 7 is used directly to mean 'week', and 'neuf' in French means '9' or 'new'). The historical distance between languages did not predict much of the variation.

The differences in semantic alignment may appear either because (A) the semantic associations are different for different languages, or (B) as a side-effect of numbers appearing with different words skewing the alignment metric. Effects that support (B) could include:

- Difference in frequency. Lower frequency terms will appear to have narrower semantic relations.
- Homophones. Different meanings will contribute different semantic relations.

However, it is more difficult to explain why different number line typologies would lead to semantic differences, unless the way numbers are constructed affects the way people think about the numbers.

References

Calude, A. S., & Verkerk, A. (2016). The typology and diachrony of higher numerals in Indo-European: a phylogenetic comparative study. Journal of Language Evolution, 1(2), 91-108.

Dehaene, S., & Mehler, J. (1992). Cross-linguistic regularities in the frequency of number words. Cognition, 43(1), 1-29.

Givón, T. (1981). On the development of the numeral 'one'as an indefinite marker. Folia Linguistica Historica, 15(Historica vol. 2, 1), 35-54.

4.3 Neighbour net visualisation of semantic alignment

The semantic alignment between languages was converted into a semantic distance and was visualised as a Neighbour Net using Splitstree (Huson & Bryant, 2006). This was done on the data with both the language filter and the concept filter. The code for building the Neighbour Net and the resulting nexus file are here:

In Neighbour Nets, distances are reflected in journeys along the lines, not in 2D space. Conflicting signals show up as parallel lines. Figure 2.3.1 shows the semantic distances for all languages in the sample (Delta score = 0.3844, Q-residual score = 0.0008379). Languages which are joined by short lines have higher semantic alignment (translation equivalents are closer in meaning). Languages are roughly grouped by language family, with the Indo-European group being most clear, but also Uralic, Turkic and Afro-Asiatic groups visible. The figure also reflects geographic distances, with Persian and Armenian showing up on the "Eastern" side. There is some clear conflicting signal for English between the Romance and Germanic, which reflects its mixed history (see e.g. McWhorter, 2008). This figure with languages from many different language families is somewhat misleading, since many languages that have high distances appear "together" in space (e.g. Basque and Korean), when in reality the distances along the lines are much further than for e.g. most Indo-European languages. Therefore, we also produced a Neighbour Net just for Indo-European languages.

Figure 2.3.2 shows the semantic distances for Indo-European languages visualised as a neighbournet (Delta score = 0.3062, Q-residual score = 0.003523). The semantic distances reflect established historical relationships, as shown by the labelling of the major sub-branches according to Glottolog (Hammarstrom et al.). Again, English shows conflicting signal between Germanic and Romance. There is a clear historical signal in the Slavic languages. Also, a split between 'eastern' and 'western' languages, with Romanian and Greek being halfway between the two. Hindi, Armenian, Lithuanian and Greek are more removed from the rest historically, and that's reflected in the fact that they're placed 'together', though actually the distance from Hindi to Armenian is much larger than for Russian to Ukrainian.

Some more speculative comments can also be made. The relationship between many languages reflects geographic proximity, such as the cline from Norwegian, Danish, German and Dutch. The proximity of Spanish and Catalan may also reflect contact and bilingualism. Romanian is an outlier in the romance languages and has a large amount of borrowed words from slavic languages (Schulte, 2009), so it's not suprising that it shows up on the edge of the Romance cluster. The proximity of Bulgarian and Greek may reflect geographical proximity and historical ties such as the Ottoman empire. The proximity of Armenian and Hindi is not predicted by linguistic family trees (e.g. Glottolog), but there is actually a history of contact, with trade leading to a historical Armenian population in Hindi-speaking places like Agra (e.g. Ferrier, 1973), and maybe more indirect borrowings through related languages (Pisowicz, 1995).

References

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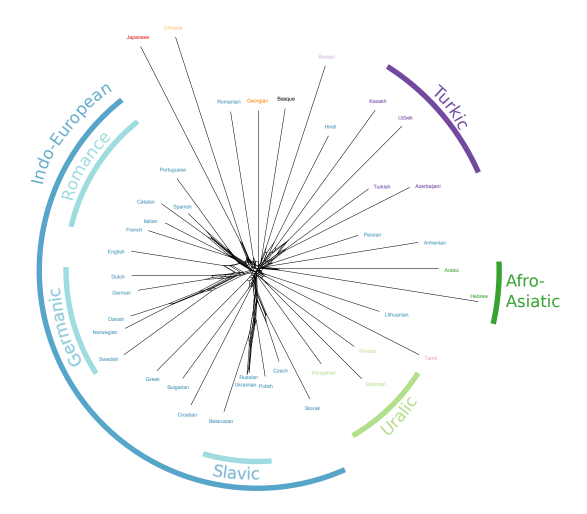
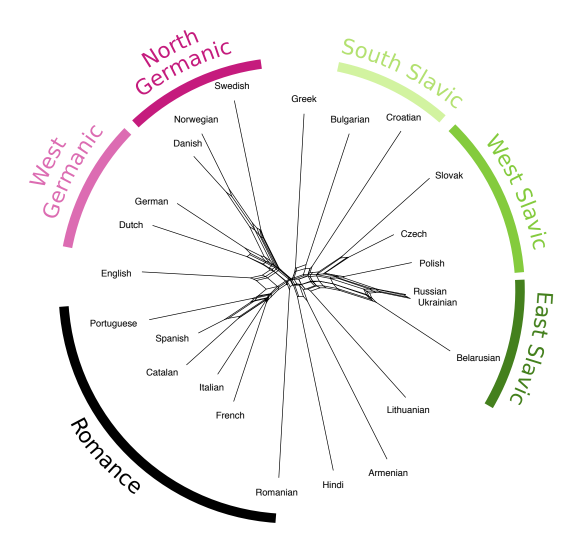


Figure 2.3.1: Neighbour Net reflecting semantic distance for all languages in the sample. Language labels are coloured by language family.



 $Figure \ 2.3.2: \ Neighbour \ Net \ reflecting \ semantic \ distance \ for \ Indo-European \ languages.$

4.4 Cross-cultural analysis (Common Crawl data)

Predicting semantic alignment by cultural similarity: Common crawl data

Bill Thompson, Seán Roberts & Gary Lupyan

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Introduction

This file replicates the tests for the main wikipedia data on the common crawl data.

Load libraries

```
library(ape)
library(ecodist)
library(lme4)
library(sjPlot)
library(ggplot2)
library(igraph)
library(lattice)
library(xtable)
```

Parameters (using data from Northuralex and common crawl, k=100, unfiltered):

```
datasetName = "cc"
datasetLabel = "Common Crawl"
lingDistancesFile = "../data/FAIR/nel-k100-cc-alignments-by-language-pair.csv"
lingDistancesFileNK = "../data/FAIR/nel-k100-cc-alignments-by-language-pair-without-kinsip.csv"
lingDistancesByDomainFile = "../results/EA_distances/nel-k100-cc_with_ling.csv"
# (generated by ../processing/combineCultAndLingDistances.R)
```

All domains

Load data

Read the cultural distances:

```
cult = read.csv("../results/EA_distances/CulturalDistances_Long.csv", stringsAsFactors = F)
names(cult) = c("11","12","cult.dist")

Add language family:

l = read.csv("../data/FAIR_langauges_glotto_xdid.csv", stringsAsFactors = F)

g = read.csv("../data/glottolog-languoid.csv/languoid.csv", stringsAsFactors = F)

l$family = g[match(1$glotto,g$id),]$family_pk

l$family = g[match(1$family,g$pk),]$name

Read the semantic distances

ling = read.csv(lingDistancesFile, stringsAsFactors = F)
```

There are very few possible comparisons for Slovenian and Northern Sami, so we'll remove these:

```
ling = ling[!(ling$11=="se" | ling$12 == "se"),]
ling = ling[!(ling$11=="s1" | ling$12 == "s1"),]
```

Combine the linguistic and cultural distances. Note that we flip the cultural measure from a distance measure to a similarity measure.

```
cult$11.iso2 = 1[match(cult$11,1$Language2),]$iso2
cult$12.iso2 = 1[match(cult$12,1$Language2),]$iso2
fairisos = unique(c(ling$11,ling$12))
cultisos = unique(c(cult$11.iso2, cult$12.iso2))
cult = cult[(cult$11.iso2 %in% fairisos) & (cult$12.iso2 %in% fairisos),]
ling = ling[(ling$11 %in% cultisos) & (ling$12 %in% cultisos),]
matches = sapply(1:nrow(ling), function(i){
 which(cult$11.iso2==ling$11[i] & cult$12.iso2==ling$12[i])
})
ling$cult.dist = cult[matches,]$cult.dist
# Flip
ling$cult.dist = 1 - ling$cult.dist
# Scale
ling$cult.dist.center = scale(ling$cult.dist)
cdc.s = attr(ling$cult.dist.center, "scaled:scale")
cdc.c = attr(ling$cult.dist.center, "scaled:center")
ling$cult.dist.center = as.numeric(ling$cult.dist.center)
ling$comparison_count.center =
  scale(ling$comparison_count)
ling$family1 = l[match(ling$11, l$iso2),]$family
ling$family2 = 1[match(ling$12, 1$iso2),]$family
1[1$Language=="Arabic",]$autotyp.area= "Greater Mesopotamia"
1[1$Language=="Persian",]$autotyp.area= "Greater Mesopotamia"
ling$area1 = l[match(ling$11, l$iso2),]$autotyp.area
```

```
ling$area2 = 1[match(ling$12, 1$iso2),]$autotyp.area
fgroup = cbind(ling$family1,ling$family2)
fgroup = apply(fgroup,1,sort)
ling$family.group = apply(fgroup,2,paste,collapse=":")
agroup = cbind(ling$area1,ling$area2)
agroup = apply(agroup,1,sort)
ling$area.group = apply(agroup,2,paste,collapse=":")
ling$rho.center = scale(ling$local_alignment)
Each observation is now assocaited with a language family pair:
head(ling[,c("11","12","local_alignment",'family.group')])
##
      11 12 local_alignment
                                     family.group
## 3 myv cv 0.05112061
                                    Turkic:Uralic
                 0.06646598 Indo-European:Turkic
## 5 la cv
## 6 cv sah
                 0.06823760
                                   Turkic:Turkic
## 13 ga cv
                0.07949409 Indo-European:Turkic
## 19 cv he
                0.08650789 Afro-Asiatic:Turkic
## 20 cv te
                 0.08761480
                                Dravidian:Turkic
And the same is true for area:
tail(ling[,c("11","12","local_alignment",'area.group')])
       11 12 local_alignment
                                          area.group
## 1112 uk ja 0.3804179 Inner Asia:N Coast Asia
## 1113 be ru
                 0.3821532 Inner Asia:Inner Asia
                  0.4022741 Inner Asia:Inner Asia
## 1115 uk be
                  0.4376378
## 1116 cs uk
                                   Europe: Inner Asia
                  0.4581089
## 1118 cs ru
                                   Europe: Inner Asia
                  0.5460480 Inner Asia:Inner Asia
## 1119 uk ru
Number of observations:
# Number of datapoints:
nrow(ling)
## [1] 308
# Number of unique languages:
length(unique(unlist(ling[,c("11","12")])))
## [1] 34
# Number of unique language families:
uniqueFamilies = unique(unlist(ling[,c("family1","family2")]))
length(uniqueFamilies)
## [1] 7
# Number of unique areas:
uniqueAreas = unique(unlist(ling[,c("area1", "area2")]))
length(uniqueAreas)
```

Cross-over between language famlies and areas:

Sino-Tibetan

Turkic ## Uralic

##

##

```
table(tx$fam,tx$area)
```

##							
##		Europe	${\tt Greater}$	Mesopotamia	${\tt Indic}$	Inner Asia N	N Coast Asia
##	Afro-Asiatic	0		1	0	0	0
##	Dravidian	0		0	3	0	0
##	Indo-European	10		2	1	5	0
##	Japonic	0		0	0	0	1
##	Sino-Tibetan	0		0	0	0	0
##	Turkic	0		1	0	5	0
##	Uralic	1		0	0	3	0
##							
##		Southea	ast Asia				
##	Afro-Asiatic		0				
##	Dravidian		0				
##	Indo-European		0				
##	Japonic		0				

1

0

LMER models

Mixed effects model, predicting Linguistic similaritys from cultural similarity, with random intercept for family and area and random slope for cultural similarity for family and area.

We start with a null model with random intercepts for family and area, and random slopes for cultural similarity by both. We add a fixed effect of the number of comparisons made for each datapoint (number of concepts that were available to compare). Then we add a fixed effect of cultural similarity

```
m0 = lmer(
  rho.center ~ 1 +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling
## boundary (singular) fit: see ?isSingular
m0.5 = lmer(
 rho.center ~ 1 +
    comparison_count.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling
)
## boundary (singular) fit: see ?isSingular
m1 = lmer(
 rho.center ~ 1 +
    comparison_count.center +
    cult.dist.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling
## boundary (singular) fit: see ?isSingular
an1 = anova(m0, m0.5, m1)
## refitting model(s) with ML (instead of REML)
an1
## Data: ling
## Models:
## m0: rho.center ~ 1 + (1 + cult.dist.center | family.group) + (1 +
           cult.dist.center | area.group)
## m0.5: rho.center ~ 1 + comparison_count.center + (1 + cult.dist.center |
## m0.5:
            family.group) + (1 + cult.dist.center | area.group)
## m1: rho.center ~ 1 + comparison_count.center + cult.dist.center +
           (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
## m1:
## m1:
           area.group)
##
       Df
             AIC
                    BIC logLik deviance
                                            Chisq Chi Df Pr(>Chisq)
        8 815.34 845.18 -399.67
## mO
                                 799.34
## m0.5 9 797.25 830.82 -389.63
                                   779.25 20.0908
                                                       1 7.385e-06 ***
## m1
       10 794.31 831.61 -387.15 774.31 4.9442
                                                       1
                                                            0.02618 *
## ---
```

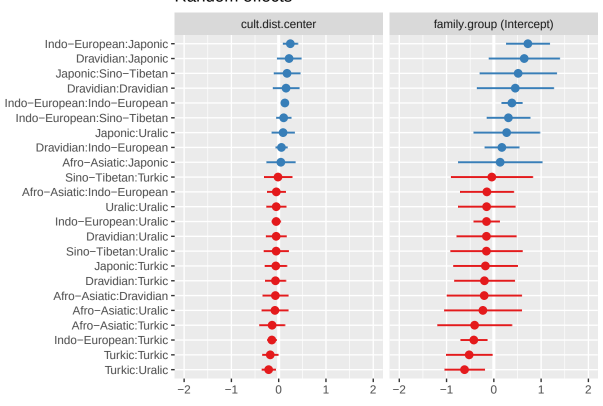
```
Cultural similarity is significantly correlated with Linguistic similarity. Here are the model estimates:
summary(m1)
## Linear mixed model fit by REML ['lmerMod']
## Formula: rho.center ~ 1 + comparison_count.center + cult.dist.center +
       (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
##
       area.group)
##
      Data: ling
##
## REML criterion at convergence: 784.1
##
## Scaled residuals:
       Min 1Q Median
                                3Q
                                        Max
## -1.9661 -0.8427 0.0488 0.7422 4.3734
##
## Random effects:
## Groups
                                   Variance Std.Dev. Corr
##
   family.group (Intercept)
                                  0.247132 0.49712
                 cult.dist.center 0.028994 0.17028
##
                                                     1.00
##
                 (Intercept)
                                  0.059174 0.24326
   area.group
##
                 cult.dist.center 0.006892 0.08302 1.00
                                   0.643045 0.80190
## Residual
## Number of obs: 308, groups: family.group, 23; area.group, 19
##
## Fixed effects:
##
                           Estimate Std. Error t value
## (Intercept)
                            0.01866 0.15042 0.124
## comparison_count.center 0.27254
                                        0.05689
                                                4.791
                                       0.07684
## cult.dist.center
                            0.18217
                                                  2.371
##
## Correlation of Fixed Effects:
##
              (Intr) cmpr_.
## cmprsn_cnt. 0.098
## clt.dst.cnt 0.669 -0.030
## convergence code: 0
## boundary (singular) fit: see ?isSingular
Plot the estimates, rescaling the variables back to the original units:
trans = function(X){
 X * attr(ling$rho.center,"scaled:scale") +
  attr(ling$rho.center,"scaled:center")
}
gx = plot_model(m1,'pred',terms='cult.dist.center')
gx$data$predicted = trans(gx$data$predicted)
gx$data$conf.low = trans(gx$data$conf.low)
gx$data$conf.high = trans(gx$data$conf.high)
gx$data$x = gx$data$x *
  cdc.s +cdc.c
gx = gx + \#coord\_cartesian(ylim=c(0,0.5),
                           xlim=c(0.15,0.85)) +
 xlab("Cultural similarity") +
```

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

```
ylab("Semantic alignment") +
  ggtitle("") +
  geom_point(data=ling,aes(x=cult.dist,y=local_alignment))
gx
   0.5 -
   0.4 -
Semantic alignment
   0.3 -
   0.2 -
   0.1 -
      0.2
                                                     0.5
                                                                                    0.7
                      0.3
                                     0.4
                                                                     0.6
                                            Cultural similarity
pdf(paste0("../results/stats/",datasetName,"/CulturalDistance_Rho_Graph.pdf"),
    height=2.5, width=2.5)
gx
dev.off()
## pdf
##
Plot the random effects:
plot_model(m1,'re', sort.est = "cult.dist.center")
```

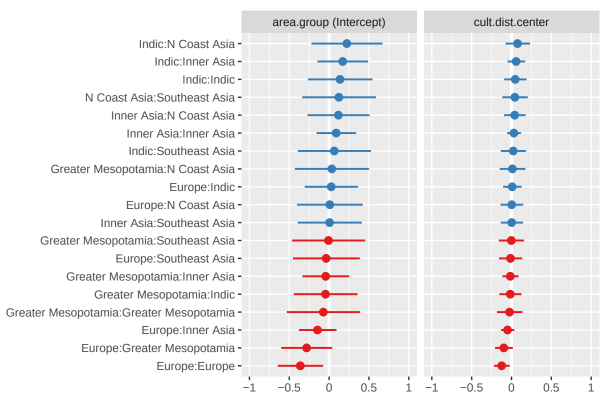
[[1]]

Random effects



[[2]]

Random effects



MRM

Use multiple regression on distance matrices to do the same test as above. The code below loads the data into a matrix format:

```
# Use graph method to make distance matrix
grph <- graph.data.frame(ling[,c("11",'12','local_alignment')], directed=FALSE)</pre>
# add value as a weight attribute
ling.m = get.adjacency(grph, attr="local_alignment", sparse=FALSE)
rownames(ling.m) = 1[match(rownames(ling.m), 1$iso2), ]$Language2
colnames(ling.m) = 1[match(colnames(ling.m),1$iso2),]$Language2
# Same for comparison_count.center
grph <- graph.data.frame(ling[,c("11",'12','comparison_count')], directed=FALSE)</pre>
# add value as a weight attribute
cc.m = get.adjacency(grph, attr="comparison_count", sparse=FALSE)
rownames(cc.m) = 1[match(rownames(cc.m),1$iso2),]$Language2
colnames(cc.m) = 1[match(colnames(cc.m),1$iso2),]$Language2
cult.m = read.csv("../results/EA_distances/CulturalDistances.csv", stringsAsFactors = F)
rownames(cult.m) = cult.m[,1]
cult.m = cult.m[,2:ncol(cult.m)]
cult.m = as.matrix(cult.m)
# Flip cultural value to distance
cult.m = 1-cult.m
mx = match(rownames(ling.m),rownames(cult.m))
cult.m = cult.m[mx,mx]
```

```
colnames(cult.m) = rownames(cult.m)
# Same/different matrix for language family
family.matrix = 1[match(rownames(ling.m), 1$Language), ]$family
family.matrix = outer(family.matrix,family.matrix,"!=") *1
# Load ASJP distances for second test
asjp = readRDS("../data/ASJP/asjp17-dists_FAIR.RData")
ling.m.glotto = 1[match(rownames(cult.m),1$Language2),]$glotto
ling.m.glotto = ling.m.glotto[ling.m.glotto %in% rownames(asjp)]
asjp.m = asjp[ling.m.glotto,ling.m.glotto]
asjp.lang.names = 1[match(rownames(asjp.m), 1$glotto), ]$Language2
# Matrices for second analysis with asjp
ling.m2 = ling.m[asjp.lang.names,asjp.lang.names]
cult.m2 = cult.m[asjp.lang.names,asjp.lang.names]
cc.m2 = cc.m[asjp.lang.names,asjp.lang.names]
# Load the geographic distances:
geoDist = read.csv("../data/GeographicDistances.csv",stringsAsFactors = F)
geoDist.m = as.matrix(geoDist)
geoDist.m = geoDist.m[!is.na(geoDist.m[,1]),!is.na(geoDist.m[1,])]
# Convert to log distance in thousand km
geoDist.m = log10(geoDist.m/1000)
geoDist.m[is.infinite(geoDist.m)] = 0
colnames(geoDist.m) = gsub("\\."," ",colnames(geoDist.m))
rownames(geoDist.m) = colnames(geoDist.m)
geoDist.m1 = geoDist.m[rownames(ling.m),rownames(ling.m)]
geoDist.m2 = geoDist.m[rownames(ling.m2),rownames(ling.m2)]
# center and scale values
ling.m = matrix(scale(as.vector(ling.m)),nrow=nrow(ling.m))
cc.m = matrix(scale(as.vector(cc.m)),nrow=nrow(cc.m))
cult.m = matrix(scale(as.vector(cult.m)),nrow=nrow(cult.m))
geoDist.m1 = matrix(scale(as.vector(geoDist.m1)),nrow=nrow(geoDist.m1))
asjp.m = matrix(scale(as.vector(asjp.m)),nrow=nrow(asjp.m))
ling.m2 = matrix(scale(as.vector(ling.m2)),nrow=nrow(ling.m2))
cc.m2 = matrix(scale(as.vector(cc.m2)),nrow=nrow(cc.m2))
cult.m2 = matrix(scale(as.vector(cult.m2)),nrow=nrow(cult.m2))
geoDist.m2 = matrix(scale(as.vector(geoDist.m2)),nrow=nrow(geoDist.m2))
```

Run the MRM model, predicting semantic alignment by cultural distance, controlling for family distance, geographic ditance, and the comparison count (number of observations). Here, the family distance between two languages is just whether they are part of the same family. Note that this does not take into account particular values for particular families, nor the random slopes within families.

```
set.seed(289)
MRM.fam = ecodist::MRM(as.dist(ling.m) ~
               as.dist(cult.m) +
               as.dist(family.matrix) +
               as.dist(geoDist.m1) +
               as.dist(cc.m),nperm = 10000)
MRM.asjp = ecodist::MRM(as.dist(ling.m2) ~
               as.dist(cult.m2) +
               as.dist(asjp.m) +
               as.dist(geoDist.m2) +
               as.dist(cc.m2),nperm = 10000)
rownames(MRM.fam$coef) = c("Intercept", "Cultural distance", "Language family",
                           "Geographic distance", "Comparison count")
colnames(MRM.fam$coef) = c("Estimate", "p-value")
statMRM.fam = xtable(MRM.fam$coef, digits = 3,display=c("s",'f',"fg"),
                     caption = paste0(
        "MRM analysis predicting semantic alignment (",
        datasetLabel,"), with family control. $R^2$=",
        signif(MRM.fam$r.squared[1],3)))
print(statMRM.fam, "latex",
    file="../results/stats/tex/MRM_family_CC.tex")
rownames(MRM.asjp$coef) = c("Intercept", "Cultural distance", "ASJP",
                           "Geographic distance", "Comparison count")
colnames(MRM.asjp$coef) = c("Estimate","p-value")
statMRM.fam = xtable(MRM.asjp$coef, digits = 3,display=c("s",'f',"fg"),
                     caption = paste0(
        "MRM analysis predicting semantic alignment (",
        datasetLabel,"), with ASJP control. $R^2$=",
        signif(MRM.asjp$r.squared[1],3)))
print(statMRM.fam, "latex",
    file="../results/stats/tex/MRM_ASJP_CC.tex")
```

Mantel tests

Read the historical distances for Indo-European, based on the phylogenetic distances.

Data prep

```
The geographic distances are loaded above (from "../data/GeographicDistances.csv").
```

Load historical distances:

```
hist = read.csv("../data/trees/IndoEuropean_historical_distances.csv", stringsAsFactors = F)
hist = hist[!duplicated(hist[,1]),!duplicated(hist[,1])]
rownames(hist) = hist[,1]
```

```
hist = hist[,2:ncol(hist)]
hist.m = as.matrix(hist)
colnames(hist.m) = rownames(hist.m)
hist.m = hist.m/max(hist.m)
```

Read the cultural distance as a matrix:

```
cult.m = read.csv("../results/EA_distances/CulturalDistances.csv", stringsAsFactors = F)
rownames(cult.m) = cult.m[,1]
cult.m = cult.m[,2:ncol(cult.m)]
```

Flip the cultural distance into a cultural similarity measure:

```
cult.m = 1-cult.m
```

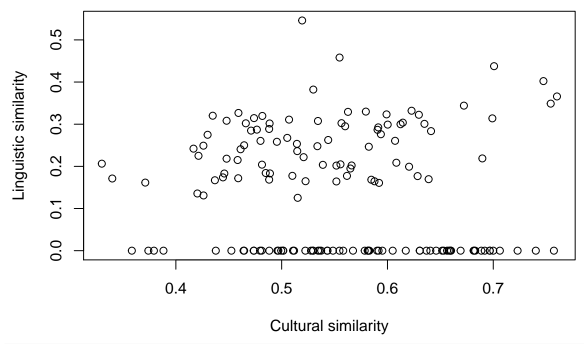
Convert the linguistic similarities to a matrix. This uses igraph to make an undirected graph from the long format with local_alignment as the edge weights, then output a matrix of adjacencies.

```
grph <- graph.data.frame(ling[,c("l1",'l2','local_alignment')], directed=FALSE)
# add value as a weight attribute
ling.m = get.adjacency(grph, attr="local_alignment", sparse=FALSE)
rownames(ling.m) = l[match(rownames(ling.m),l$iso2),]$Language2
colnames(ling.m) = l[match(colnames(ling.m),l$iso2),]$Language2</pre>
```

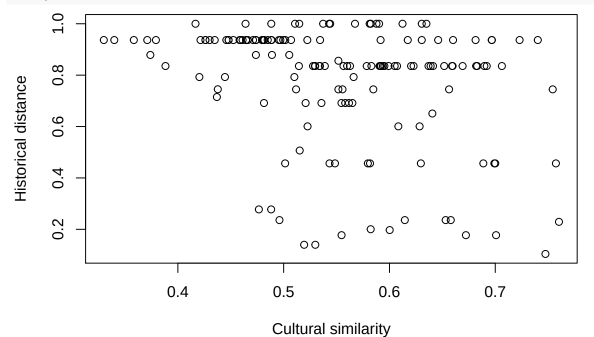
Match the distance matrices

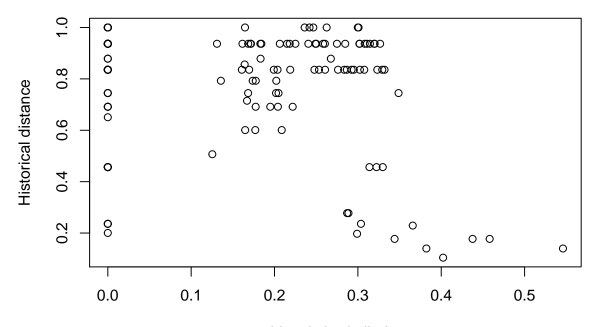
```
in.analysis = intersect(rownames(ling.m),rownames(cult.m))
in.analysis = intersect(in.analysis, rownames(hist.m))
cult.m2 = cult.m[in.analysis,in.analysis]
ling.m2 = ling.m[in.analysis,in.analysis]
hist.m2 = hist.m[in.analysis,in.analysis]
geo.m2 = geoDist.m[in.analysis,in.analysis]
```

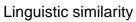
Note that there are only 18 languages with data on linguistic, cultural and historical distance.

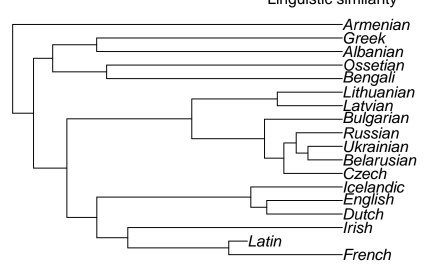












Tests

The results of the test list the following measures:

```
• mantelr: Mantel correlation coefficient.

• pval1: one-tailed p-value (null hypothesis: r <= 0).

• pval2: one-tailed p-value (null hypothesis: r >= 0).

• pval3: two-tailed p-value (null hypothesis: r = 0).

• llim: lower confidence limit for r.

• ulim: upper confidence limit for r.
```

set.seed(1498)

Run tests between each pair of measures.

```
distms = list("Cultrual"= cult.m2,
              "Linguistic" = ling.m2,
              "Historical" = hist.m2,
              "Geographic" = geo.m2)
mantelRes1 = data.frame(
 Var1 = NA, Var2 = NA, r = NA,
 llim = NA, ulim = NA, p = NA,
  stringsAsFactors = F)
for(i in 1:3){
  for(j in (i+1):4){
   var1 = names(distms)[i]
    var2 = names(distms)[j]
    print(paste("Correlation between",
                var1, "and", var2))
    stat = ecodist::mantel(as.dist(distms[[i]]) ~
                as.dist(distms[[j]]),
                nperm = 100000)
    print(stat)
    mantelRes1 = rbind(mantelRes1,
        c(var1, var2, stat[1], stat[5], stat[6],
          min(c(stat[2],stat[3]))))
    stat = round(stat,2)
    stat2 = sprintf("$r$ = %s[%s,%s], one-tailed $p$ = %s",
      stat[1],
      stat[5],
      stat[6],
      min(c(stat[2],stat[3])))
    # TODO: output stats
    #cat(stat2,file=
           paste0("../results/stats/tex/Mantel",var1,"Vs",var2,"Distance_CC.tex"))
  }
```

```
## [1] "Correlation between Cultrual and Linguistic"
## mantelr pval1 pval2 pval3 llim.2.5%
## -0.0839735358 0.7232800000 0.2767300000 0.5379200000 -0.2009313168
## ulim.97.5%
## 0.0008846432
## [1] "Correlation between Cultrual and Historical"
```

```
pval2
##
     mantelr
                 pval1
                                      pval3 llim.2.5% ulim.97.5%
## [1] "Correlation between Cultrual and Geographic"
                                      pval3 llim.2.5% ulim.97.5%
     mantelr
                 pval1
                            pval2
## -0.4608256 0.9970600 0.0029500 0.0029500 -0.5860424 -0.3118663
## [1] "Correlation between Linguistic and Historical"
     mantelr
                 pval1
                            pval2
                                      pval3 llim.2.5% ulim.97.5%
## -0.2496921 0.9850200 0.0149900 0.0220500 -0.3570435 -0.1142948
## [1] "Correlation between Linguistic and Geographic"
                                                  llim.2.5% ulim.97.5%
##
                                          pval3
                   pval1
                               pval2
  0.13621101 0.12091000 0.87910000 0.25937000 -0.01342846 0.23456752
##
## [1] "Correlation between Historical and Geographic"
                 pval1
##
     mantelr
                            pval2
                                      pval3 llim.2.5% ulim.97.5%
   0.4052690 \quad 0.0010100 \quad 0.9990000 \quad 0.0010100 \quad 0.3098220 \quad 0.5192832
mantelRes1= mantelRes1[2:nrow(mantelRes1),]
mantelRes1[,3:6] = apply(mantelRes1[,3:6],2,function(X){
 signif(as.numeric(X),3)
```

Run a mantel test comparing the Linguistic alignment to the cultural similarity, controlling for the historical distance between languages:

```
## mantelr pval1 pval2 pval3 llim.2.5% ulim.97.5% ## -0.17690227 0.91913000 0.08088000 0.17506000 -0.24784616 -0.08286161
```

Main Test: Run a mantel test comparing the Linguistic alignment to the cultural similarity, controlling for the historical distance and geographic distance between languages:

```
mainMantel = ecodist::mantel(as.dist(ling.m2)~
                  as.dist(cult.m2) +
                  as.dist(hist.m2) +
                  as.dist(geo.m2),
                nperm = 100000)
mainMantel = signif(mainMantel,3)
mantelRes1 = rbind(mantelRes1,
  c("Linguistic", "Cultural **",
    mainMantel[1],mainMantel[5],mainMantel[6],
    min(mainMantel[2:3])))
mantelRes1Text = xtable(mantelRes1,
   caption = paste0(
    "Mantel tests (",
    datasetLabel,
    "). ** = partial Mantel test, controlling for historical and geographical distance."))
print(mantelRes1Text,
   file="../results/stats/tex/Mantel CC.tex")
```

MRM

Perform the main test, but using multiple regression on distance matrices (MRM).

Lichstein, J. W. (2007). Multiple regression on distance matrices: a multivariate spatial analysis tool. Plant Ecology, 188(2), 117-131.

```
mainMRM = ecodist::MRM(as.dist(ling.m2)~
                  as.dist(cult.m2) +
                  as.dist(hist.m2) +
                  as.dist(geo.m2), nperm=10000)
mainMRM
## $coef
##
                    as.dist(ling.m2)
                                       pval
## Int
                          0.3645355 0.0352
                         -0.1327268 0.5157
## as.dist(cult.m2)
## as.dist(hist.m2)
                         -0.2254171 0.0010
## as.dist(geo.m2)
                         0.1030124 0.0503
##
## $r.squared
##
        R2
                pval
## 0.135791 0.017100
##
## $F.test
##
       F F.pval
## 7.8040 0.0171
mainMRM2 = sprintf("$\\beta= $%s, $p=$%s",
                   round(mainMRM$coef[2,1],2),
                   round(mainMRM$coef[2,2],2))
cat(mainMRM2,
   file="../results/stats/tex/MRMCultrualVsLinguisticDistance_Partial_CC.tex")
```

4.5 Cross-cultural analysis (Subtitles data)

Predicting semantic alignment by cultural similarity: Subtitles data

Bill Thompson, Seán Roberts & Gary Lupyan

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Introduction

This file replicates the tests for the main wikipedia data on the subtitles data.

Load libraries

```
library(ape)
library(ecodist)
library(lme4)
library(sjPlot)
library(ggplot2)
library(igraph)
library(lattice)
library(xtable)
```

Parameters (using data from Northuralex and common crawl, k=100, unfiltered):

```
datasetName = "subs"
datasetLabel = "Subtitles"
lingDistancesFile = "../data/FAIR/nel-k100-subs-alignments-by-language-pair.csv"
lingDistancesByDomainFile = "../results/EA_distances/nel-k100-subs_with_ling.csv"
# (generated by ../processing/combineCultAndLingDistances.R)
```

All domains

Load data

Read the cultural distances:

```
cult = read.csv("../results/EA_distances/CulturalDistances_Long.csv", stringsAsFactors = F)
names(cult) = c("11","12","cult.dist")

Add language family:

l = read.csv("../data/FAIR_languages_glotto_xdid.csv", stringsAsFactors = F)

g = read.csv("../data/glottolog-languoid.csv/languoid.csv", stringsAsFactors = F)

l$family = g[match(1$glotto,g$id),]$family_pk

l$family = g[match(1$family,g$pk),]$name

Read the semantic distances

ling = read.csv(lingDistancesFile, stringsAsFactors = F)
```

There are very few possible comparisons for Slovenian and Northern Sami, so we'll remove these:

```
ling = ling[!(ling$11=="se" | ling$12 == "se"),]
ling = ling[!(ling$11=="sl" | ling$12 == "sl"),]
```

Combine the linguistic and cultural distances. Note that we flip the cultural measure from a distance measure to a similarity measure.

```
cult$11.iso2 = 1[match(cult$11,1$Language2),]$iso2
cult$12.iso2 = 1[match(cult$12,1$Language2),]$iso2
fairisos = unique(c(ling$11,ling$12))
cultisos = unique(c(cult$11.iso2, cult$12.iso2))
cult = cult[(cult$11.iso2 %in% fairisos) & (cult$12.iso2 %in% fairisos),]
ling = ling[(ling$11 %in% cultisos) & (ling$12 %in% cultisos),]
matches = sapply(1:nrow(ling), function(i){
 which(cult$11.iso2==ling$11[i] & cult$12.iso2==ling$12[i])
})
ling$cult.dist = cult[matches,]$cult.dist
# Flip
ling$cult.dist = 1 - ling$cult.dist
# Scale
ling$cult.dist.center = scale(ling$cult.dist)
cdc.s = attr(ling$cult.dist.center, "scaled:scale")
cdc.c = attr(ling$cult.dist.center, "scaled:center")
ling$cult.dist.center = as.numeric(ling$cult.dist.center)
ling$comparison_count.center =
  scale(ling$comparison_count)
ling$family1 = l[match(ling$11, l$iso2),]$family
ling$family2 = 1[match(ling$12, 1$iso2),]$family
1[1$Language=="Arabic",]$autotyp.area= "Greater Mesopotamia"
1[1$Language=="Persian",]$autotyp.area= "Greater Mesopotamia"
ling$area1 = l[match(ling$11, l$iso2),]$autotyp.area
```

```
ling$area2 = 1[match(ling$12, 1$iso2),]$autotyp.area
fgroup = cbind(ling$family1,ling$family2)
fgroup = apply(fgroup,1,sort)
ling$family.group = apply(fgroup,2,paste,collapse=":")
agroup = cbind(ling$area1,ling$area2)
agroup = apply(agroup,1,sort)
ling$area.group = apply(agroup,2,paste,collapse=":")
ling$rho.center = scale(ling$local_alignment)
Each observation is now assocaited with a language family pair:
head(ling[,c("11","12","local_alignment",'family.group')])
##
     11 12 local_alignment
                                           family.group
## 18 hy sq 0.01014219 Indo-European: Indo-European
## 23 hy ko
                0.01862729
                                Indo-European:Koreanic
## 32 hy is
                0.03000878 Indo-European: Indo-European
## 48 hy ja
               0.05326118
                                 Indo-European: Japonic
## 49 et hy
                0.05535150
                                  Indo-European:Uralic
## 52 hy nl
                0.05835286 Indo-European:Indo-European
And the same is true for area:
tail(ling[,c("11","12","local_alignment",'area.group')])
      11 12 local_alignment
                                    area.group
## 651 cs es
             0.3339303
                                Europe: Europe
                 0.3353707
## 653 bg cs
                                Europe: Europe
                 0.3378855
## 656 el bg
                                Europe: Europe
                 0.3446584 Europe:Inner Asia
## 659 el ru
                 0.3625137
## 661 el cs
                                Europe: Europe
## 664 cs ru
                  0.3790251 Europe:Inner Asia
Number of observations:
# Number of datapoints:
nrow(ling)
## [1] 190
# Number of unique languages:
length(unique(unlist(ling[,c("11","12")])))
## [1] 20
# Number of unique language families:
uniqueFamilies = unique(unlist(ling[,c("family1","family2")]))
length(uniqueFamilies)
## [1] 6
# Number of unique areas:
uniqueAreas = unique(unlist(ling[,c("area1", "area2")]))
length(uniqueAreas)
```

Cross-over between language famlies and areas:

##								
##		Europe	Greater	Mesopotamia	Inner	Asia	N Coast	Asia
##	Afro-Asiatic	0		1		0		0
##	Indo-European	9		1		4		0
##	Japonic	0		0		0		1
##	Koreanic	0		0		0		1
##	Turkic	0		1		0		0
##	Uralic	1		0		1		0

LMER models

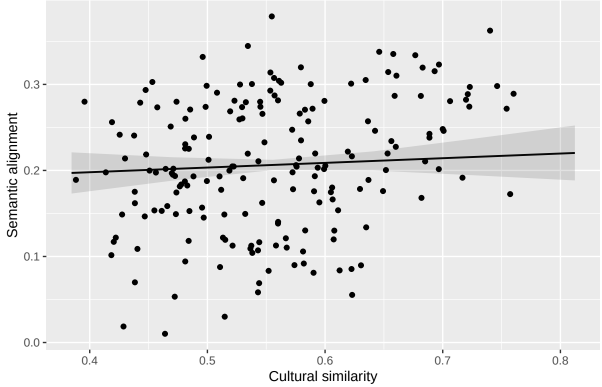
Mixed effects model, predicting Linguistic similaritys from cultural similarity, with random intercept for family and area and random slope for cultural similarity for family and area.

We start with a null model with random intercepts for family and area, and random slopes for cultural similarity by both. We add a fixed effect of the number of comparisons made for each datapoint (number of concepts that were available to compare). Then we add a fixed effect of cultural similarity

```
m0 = lmer(
  rho.center ~ 1 +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling
## boundary (singular) fit: see ?isSingular
m0.5 = lmer(
 rho.center ~ 1 +
    comparison_count.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling
)
## Warning in checkConv(attr(opt, "derivs"), opt$par, ctrl =
## control$checkConv, : Model failed to converge with max|grad| = 0.0073357
## (tol = 0.002, component 1)
m1 = lmer(
 rho.center ~ 1 +
    comparison_count.center +
    cult.dist.center +
    (1 + cult.dist.center | family.group) +
    (1 + cult.dist.center | area.group),
  data = ling
## boundary (singular) fit: see ?isSingular
an1 = anova(m0, m0.5, m1)
## refitting model(s) with ML (instead of REML)
an1
## Data: ling
## Models:
## m0: rho.center ~ 1 + (1 + cult.dist.center | family.group) + (1 +
          cult.dist.center | area.group)
## m0.5: rho.center ~ 1 + comparison_count.center + (1 + cult.dist.center |
            family.group) + (1 + cult.dist.center | area.group)
## m0.5:
## m1: rho.center ~ 1 + comparison_count.center + cult.dist.center +
           (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
## m1:
## m1:
           area.group)
##
       Df
             AIC
                     BIC logLik deviance
                                             Chisq Chi Df Pr(>Chisq)
## mO
         8 409.01 434.99 -196.51
                                   393.01
## m0.5 9 288.62 317.84 -135.31
                                   270.62 122.3948
                                                        1
                                                               <2e-16 ***
```

```
10 289.88 322.35 -134.94
## m1
                                   269.88
                                             0.7417
                                                                0.3891
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Cultural similarity is significantly correlated with Linguistic similarity. Here are the model estimates:
summary(m1)
## Linear mixed model fit by REML ['lmerMod']
## Formula: rho.center ~ 1 + comparison_count.center + cult.dist.center +
       (1 + cult.dist.center | family.group) + (1 + cult.dist.center |
##
##
       area.group)
##
      Data: ling
##
## REML criterion at convergence: 282.8
##
## Scaled residuals:
##
       Min
             1Q Median
                                 30
                                        Max
## -1.8968 -0.7471 -0.0734 0.6181 3.6760
##
## Random effects:
##
   Groups
                 Name
                                   Variance Std.Dev. Corr
##
   family.group (Intercept)
                                  0.0002465 0.01570
##
                 cult.dist.center 0.0056335 0.07506
                                                      -1.00
##
   area.group
                 (Intercept)
                                  0.0015278 0.03909
##
                 cult.dist.center 0.0211421 0.14540 -1.00
##
  Residual
                                   0.2322780 0.48195
## Number of obs: 190, groups: family.group, 17; area.group, 10
##
## Fixed effects:
##
                           Estimate Std. Error t value
                           -0.01139
                                       0.03975 -0.286
## (Intercept)
                                        0.03807 21.346
## comparison_count.center 0.81265
## cult.dist.center
                            0.06060
                                        0.07272
                                                 0.833
##
## Correlation of Fixed Effects:
##
               (Intr) cmpr_.
## cmprsn_cnt. 0.013
## clt.dst.cnt -0.231 -0.136
## convergence code: 0
## boundary (singular) fit: see ?isSingular
Plot the estimates, rescaling the variables back to the original units:
trans = function(X){
 X * attr(ling$rho.center, "scaled:scale") +
  attr(ling$rho.center, "scaled:center")
}
gx = plot_model(m1,'pred',terms='cult.dist.center')
gx$data$predicted = trans(gx$data$predicted)
gx$data$conf.low = trans(gx$data$conf.low)
gx$data$conf.high = trans(gx$data$conf.high)
gx$data$x = gx$data$x *
  cdc.s +cdc.c
gx = gx + \#coord\_cartesian(ylim=c(0,0.5),
```

```
# xlim=c(0.15,0.85)) +
xlab("Cultural similarity") +
ylab("Semantic alignment") +
ggtitle("") +
geom_point(data=ling,aes(x=cult.dist,y=local_alignment))
gx
```

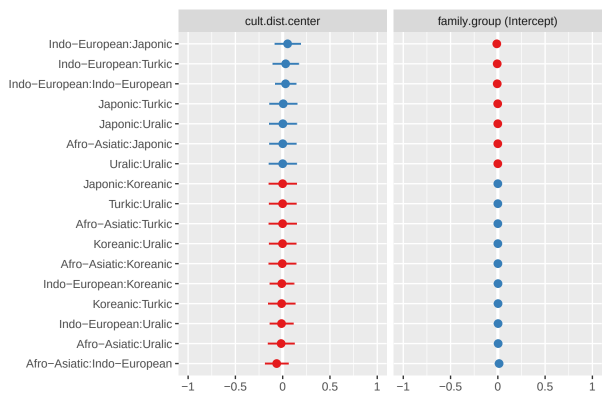


```
pdf(paste0("../results/stats/",datasetName,"/CulturalDistance_Rho_Graph.pdf"),
    height=2.5, width=2.5)
gx
dev.off()

## pdf
## 2
Plot the random effects:
plot_model(m1,'re', sort.est = "cult.dist.center")
```

[[1]]

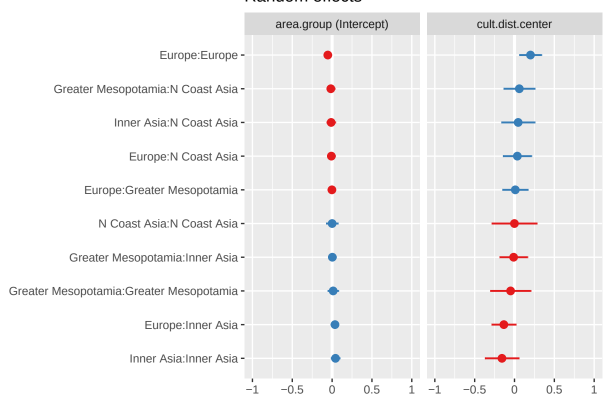
Random effects



##

[[2]]

Random effects



MRM

Use multiple regression on distance matrices to do the same test as above. The code below loads the data into a matrix format:

```
# Use graph method to make distance matrix
grph <- graph.data.frame(ling[,c("11",'12','local_alignment')], directed=FALSE)</pre>
# add value as a weight attribute
ling.m = get.adjacency(grph, attr="local_alignment", sparse=FALSE)
rownames(ling.m) = 1[match(rownames(ling.m), 1$iso2), ]$Language2
colnames(ling.m) = 1[match(colnames(ling.m),1$iso2),]$Language2
# Same for comparison_count.center
grph <- graph.data.frame(ling[,c("11",'12','comparison_count')], directed=FALSE)</pre>
# add value as a weight attribute
cc.m = get.adjacency(grph, attr="comparison_count", sparse=FALSE)
rownames(cc.m) = 1[match(rownames(cc.m),1$iso2),]$Language2
colnames(cc.m) = 1[match(colnames(cc.m),1$iso2),]$Language2
cult.m = read.csv("../results/EA_distances/CulturalDistances.csv", stringsAsFactors = F)
rownames(cult.m) = cult.m[,1]
cult.m = cult.m[,2:ncol(cult.m)]
cult.m = as.matrix(cult.m)
# Flip cultural value to distance
cult.m = 1-cult.m
mx = match(rownames(ling.m),rownames(cult.m))
cult.m = cult.m[mx,mx]
```

```
colnames(cult.m) = rownames(cult.m)
# Same/different matrix for language family
family.matrix = 1[match(rownames(ling.m), 1$Language), ]$family
family.matrix = outer(family.matrix,family.matrix,"!=") *1
# Load ASJP distances for second test
asjp = readRDS("../data/ASJP/asjp17-dists_FAIR.RData")
ling.m.glotto = 1[match(rownames(cult.m),1$Language2),]$glotto
ling.m.glotto = ling.m.glotto[ling.m.glotto %in% rownames(asjp)]
asjp.m = asjp[ling.m.glotto,ling.m.glotto]
asjp.lang.names = 1[match(rownames(asjp.m), 1$glotto), ]$Language2
# Matrices for second analysis with asjp
ling.m2 = ling.m[asjp.lang.names,asjp.lang.names]
cult.m2 = cult.m[asjp.lang.names,asjp.lang.names]
cc.m2 = cc.m[asjp.lang.names,asjp.lang.names]
# Load the geographic distances:
geoDist = read.csv("../data/GeographicDistances.csv",stringsAsFactors = F)
geoDist.m = as.matrix(geoDist)
geoDist.m = geoDist.m[!is.na(geoDist.m[,1]),!is.na(geoDist.m[1,])]
# Convert to log distance in thousand km
geoDist.m = log10(geoDist.m/1000)
geoDist.m[is.infinite(geoDist.m)] = 0
colnames(geoDist.m) = gsub("\\."," ",colnames(geoDist.m))
rownames(geoDist.m) = colnames(geoDist.m)
geoDist.m1 = geoDist.m[rownames(ling.m),rownames(ling.m)]
geoDist.m2 = geoDist.m[rownames(ling.m2),rownames(ling.m2)]
# center and scale values
ling.m = matrix(scale(as.vector(ling.m)),nrow=nrow(ling.m))
cc.m = matrix(scale(as.vector(cc.m)),nrow=nrow(cc.m))
cult.m = matrix(scale(as.vector(cult.m)),nrow=nrow(cult.m))
geoDist.m1 = matrix(scale(as.vector(geoDist.m1)),nrow=nrow(geoDist.m1))
asjp.m = matrix(scale(as.vector(asjp.m)),nrow=nrow(asjp.m))
ling.m2 = matrix(scale(as.vector(ling.m2)),nrow=nrow(ling.m2))
cc.m2 = matrix(scale(as.vector(cc.m2)),nrow=nrow(cc.m2))
cult.m2 = matrix(scale(as.vector(cult.m2)),nrow=nrow(cult.m2))
geoDist.m2 = matrix(scale(as.vector(geoDist.m2)),nrow=nrow(geoDist.m2))
```

Run the MRM model, predicting semantic alignment by cultural distance, controlling for family distance, geographic ditance, and the comparison count (number of observations). Here, the family distance between two languages is just whether they are part of the same family. Note that this does not take into account particular values for particular families, nor the random slopes within families.

```
set.seed(289)
MRM.fam = ecodist::MRM(as.dist(ling.m) ~
               as.dist(cult.m) +
               as.dist(family.matrix) +
               as.dist(geoDist.m1) +
               as.dist(cc.m),nperm = 10000)
MRM.asjp = ecodist::MRM(as.dist(ling.m2) ~
               as.dist(cult.m2) +
               as.dist(asjp.m) +
               as.dist(geoDist.m2) +
               as.dist(cc.m2),nperm = 10000)
rownames(MRM.fam$coef) = c("Intercept", "Cultural distance", "Language family",
                           "Geographic distance", "Comparison count")
colnames(MRM.fam$coef) = c("Estimate", "p-value")
statMRM.fam = xtable(MRM.fam$coef, digits = 3,display=c("s",'f',"fg"),
                     caption = paste0(
        "MRM analysis predicting semantic alignment (",
        datasetLabel,"), with family control. $R^2$=",
        signif(MRM.fam$r.squared[1],3)))
print(statMRM.fam, "latex",
    file="../results/stats/tex/MRM_family_SUBS.tex")
rownames(MRM.asjp$coef) = c("Intercept", "Cultural distance", "ASJP",
                           "Geographic distance", "Comparison count")
colnames(MRM.asjp$coef) = c("Estimate","p-value")
statMRM.fam = xtable(MRM.asjp$coef, digits = 3,display=c("s",'f',"fg"),
                     caption = paste0(
        "MRM analysis predicting semantic alignment (",
        datasetLabel,"), with ASJP control. $R^2$=",
        signif(MRM.asjp$r.squared[1],3)))
print(statMRM.fam, "latex",
    file="../results/stats/tex/MRM_ASJP_SUBS.tex")
```

Mantel tests

Read the historical distances for Indo-European, based on the phylogenetic distances.

Data prep

```
The geographic distances are loaded above (from "../data/GeographicDistances.csv").
```

Load historical distances:

```
hist = read.csv("../data/trees/IndoEuropean_historical_distances.csv", stringsAsFactors = F)
hist = hist[!duplicated(hist[,1]),!duplicated(hist[,1])]
rownames(hist) = hist[,1]
```

```
hist = hist[,2:ncol(hist)]
hist.m = as.matrix(hist)
colnames(hist.m) = rownames(hist.m)
hist.m = hist.m/max(hist.m)
```

Read the cultural distance as a matrix:

```
cult.m = read.csv("../results/EA_distances/CulturalDistances.csv", stringsAsFactors = F)
rownames(cult.m) = cult.m[,1]
cult.m = cult.m[,2:ncol(cult.m)]
```

Flip the cultural distance into a cultural similarity measure:

```
cult.m = 1-cult.m
```

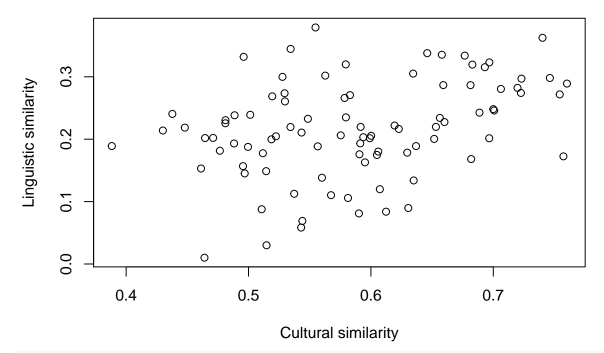
Convert the linguistic similarities to a matrix. This uses igraph to make an undirected graph from the long format with local_alignment as the edge weights, then output a matrix of adjacencies.

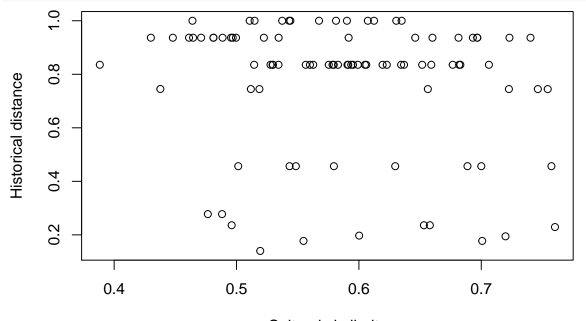
```
grph <- graph.data.frame(ling[,c("l1",'l2','local_alignment')], directed=FALSE)
# add value as a weight attribute
ling.m = get.adjacency(grph, attr="local_alignment", sparse=FALSE)
rownames(ling.m) = l[match(rownames(ling.m), 1$iso2),]$Language2
colnames(ling.m) = l[match(colnames(ling.m), 1$iso2),]$Language2</pre>
```

Match the distance matrices

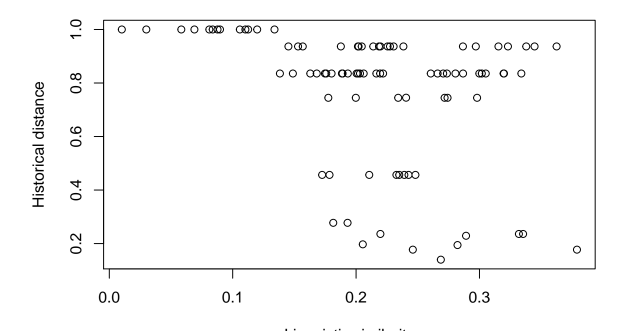
```
in.analysis = intersect(rownames(ling.m),rownames(cult.m))
in.analysis = intersect(in.analysis, rownames(hist.m))
cult.m2 = cult.m[in.analysis,in.analysis]
ling.m2 = ling.m[in.analysis,in.analysis]
hist.m2 = hist.m[in.analysis,in.analysis]
geo.m2 = geoDist.m[in.analysis,in.analysis]
```

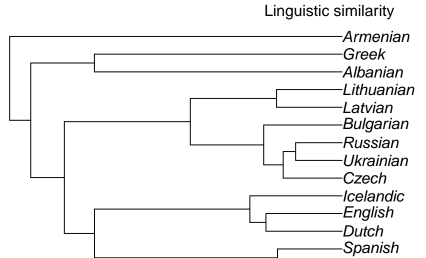
Note that there are only 14 languages with data on linguistic, cultural and historical distance.





```
Cultural similarity
```





-French

Tests

The results of the test list the following measures:

```
• mantelr: Mantel correlation coefficient.

• pval1: one-tailed p-value (null hypothesis: r <= 0).

• pval2: one-tailed p-value (null hypothesis: r >= 0).

• pval3: two-tailed p-value (null hypothesis: r = 0).

• llim: lower confidence limit for r.
```

• ulim: upper confidence limit for r.

```
set.seed(1498)
```

Run tests between each pair of measures.

```
distms = list("Cultrual"= cult.m2,
              "Linguistic" = ling.m2,
              "Historical" = hist.m2,
              "Geographic" = geo.m2)
mantelRes1 = data.frame(
 Var1 = NA, Var2 = NA, r = NA,
 llim = NA, ulim = NA, p = NA,
  stringsAsFactors = F)
for(i in 1:3){
  for(j in (i+1):4){
   var1 = names(distms)[i]
    var2 = names(distms)[j]
    print(paste("Correlation between",
                var1, "and", var2))
    stat = ecodist::mantel(as.dist(distms[[i]]) ~
                as.dist(distms[[j]]),
                nperm = 100000)
    print(stat)
    mantelRes1 = rbind(mantelRes1,
        c(var1, var2, stat[1], stat[5], stat[6],
          min(c(stat[2],stat[3]))))
    stat = round(stat,2)
    stat2 = sprintf("$r$ = %s[%s,%s], one-tailed $p$ = %s",
      stat[1],
      stat[5],
      stat[6],
      min(c(stat[2],stat[3])))
    # TODO: output stats
    #cat(stat2,file=
           paste0("../results/stats/tex/Mantel",var1,"Vs",var2,"Distance_SUBS.tex"))
  }
```

```
## [1] "Correlation between Cultrual and Linguistic"
## mantelr pval1 pval2 pval3 llim.2.5% ulim.97.5%
## 0.3509606 0.0883600 0.9116500 0.1382200 0.2413895 0.5377689
## [1] "Correlation between Cultrual and Historical"
## mantelr pval1 pval2 pval3 llim.2.5% ulim.97.5%
## -0.16966425 0.84537000 0.15464000 0.30858000 -0.28569297 -0.01526378
```

```
## [1] "Correlation between Cultrual and Geographic"
               pval1
                        pval2
##
                                 pval3 llim.2.5% ulim.97.5%
    mantelr
## -0.3397939 0.9723900 0.0276200 0.0308800 -0.5704463 -0.1718367
## [1] "Correlation between Linguistic and Historical"
##
     mantelr
                 pval1
                          pval2
                                    pval3
                                           llim.2.5% ulim.97.5%
## [1] "Correlation between Linguistic and Geographic"
                        pval2
                                 pval3 llim.2.5% ulim.97.5%
               pval1
## [1] "Correlation between Historical and Geographic"
##
                                 pval3 llim.2.5% ulim.97.5%
    mantelr
               pval1
                        pval2
   0.3457543 0.0100300 0.9899800 0.0100300 0.1811727 0.5217647
mantelRes1= mantelRes1[2:nrow(mantelRes1),]
mantelRes1[,3:6] = apply(mantelRes1[,3:6],2,function(X){
 signif(as.numeric(X),3)
})
```

Run a mantel test comparing the Linguistic alignment to the cultural similarity, controlling for the historical distance between languages:

```
## mantelr pval1 pval2 pval3 llim.2.5% ulim.97.5%
## 0.3157282 0.1119100 0.8881000 0.1894500 0.1495754 0.5004242
```

Main Test: Run a mantel test comparing the Linguistic alignment to the cultural similarity, controlling for the historical distance and geographic distance between languages:

```
mainMantel = ecodist::mantel(as.dist(ling.m2)~
                  as.dist(cult.m2) +
                  as.dist(hist.m2) +
                  as.dist(geo.m2),
                nperm = 100000)
mainMantel = signif(mainMantel,3)
mantelRes1 = rbind(mantelRes1,
  c("Linguistic", "Cultural **",
    mainMantel[1],mainMantel[5],mainMantel[6],
    min(mainMantel[2:3])))
mantelRes1Text = xtable(mantelRes1,
   caption = paste0(
    "Mantel tests (",
    datasetLabel,
    "). ** = partial Mantel test, controlling for historical and geographical distance."))
print(mantelRes1Text,
    file="../results/stats/tex/Mantel_SUBS.tex")
```

MRM

Perform the main test, but using multiple regression on distance matrices (MRM).

```
mainMRM = ecodist::MRM(as.dist(ling.m2)~
                 as.dist(cult.m2) +
                 as.dist(hist.m2) +
                 as.dist(geo.m2), nperm=10000)
mainMRM
## $coef
##
                   as.dist(ling.m2) pval
## Int
                        0.14290281 0.6955
## as.dist(cult.m2)
                        0.24395908 0.2398
                      -0.08531160 0.1408
## as.dist(hist.m2)
## as.dist(geo.m2)
                       -0.02225994 0.6608
##
## $r.squared
## R2
                 pval
## 0.2169477 0.1326000
##
## $F.test
## F F.pval
## 8.034563 0.132600
mainMRM2 = sprintf("$\\beta= $%s, $p=$%s",
                  round(mainMRM$coef[2,1],2),
                  round(mainMRM$coef[2,2],2))
cat(mainMRM2,
   file="../results/stats/tex/MRMCultrualVsLinguisticDistance_Partial_SUBS.tex")
```