## visualise

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This Rmarkdown contains all code necassary to re-produce plots and key statistics within the paper

## Load packages

```
library(dplyr)
library(ggplot2)
library(ggpubr)
library(tidyr)
library(maptools) ## For wrld_simpl
library(raster)
library(RColorBrewer)
library(ggtree)
library(ape)
library(phytools)
library(INLA)
library(brinla)
```

#### **Descriptives**

Overall summary, reporting on the number of populations, species and sites contained within the overall dataset

```
compiled_data = readRDS("../data/derived_data/compiled_data.rds")
length(unique(compiled_data$site_spec))
```

```
## [1] 30730
```

```
length(unique(compiled_data$species))
## [1] 3141
length(unique(compiled data$site))
## [1] 5991
Summaries per dataset Here we report the spatial, taxonomic and temporal extent of each dataset. This information feeds directly into Table S1.
analysis_list = readRDS("../data/derived_data/analysis_list.rds")
descriptives df = NULL
for(a in names(analysis_list)){
  tmp df = analysis list[[a]][[1]]
  tmp df = subset(tmp df, !is.na(log abundance))
  tmp descriptives df = data.frame(
    Dataset = a,
    Observations = nrow(tmp df),
    Populations = length(unique(tmp df$site spec code)),
    Species = length(unique(tmp_df$tips_code)),
    Sites = length(unique(tmp_df$site_code)),
    Timeframe = pasteO(min(tmp_df$date), " - ", max(tmp_df$date)),
   Latitude = paste0(round(min(tmp_df$latitude),1), " - ", round(max(tmp_df$latitude),1)),
   Longitude = paste0(round(min(tmp_df$longitude),1), " - ", round(max(tmp_df$longitude),1))
  descriptives_df = rbind(descriptives_df, tmp_descriptives_df)
print(descriptives_df)
               Dataset Observations Populations Species Sites
##
                                                                 Timeframe
## 1
               BioTIME
                              243993
                                           12065
                                                    1233
                                                           438 1933 - 2018
## 2
             CaPTrends
                                2670
                                             279
                                                           165 1880 - 2019
## 3
               EA NFPD
                                2999
                                             359
                                                      16 179 1984 - 2019
              FishGlob
                                            2286
                                                           229 1977 - 2020
## 4
                               67908
                                                     152
                                                    1333 1244 1950 - 2020
## 5
                   LPI
                              77773
                                            3613
```

```
## 6
               {\tt NAm\_BBS}
                             164317
                                            8718
                                                           584 1966 - 2019
## 7
               Pilotto
                              11353
                                            582
                                                            67 1974 - 2018
                                                     356
    ReSurvey Germany
                               4954
                                             356
                                                      93
                                                             7 1966 - 2016
## 8
               RivFish
                                            2386
                                                           197 1975 - 2019
## 9
                              40834
                                                     191
## 10
              TimeFISH
                                262
                                              86
                                                      52
                                                            12 2008 - 2022
##
          Latitude
                         Longitude
       -77.6 - 68.6 -179.8 - 179.2
## 1
## 2
        -40 - 71.6
                      -158 - 99.2
## 3
        50.4 - 55.4
                       -3.9 - 0.5
## 4
            26 - 62
                        -178 - 21
## 5
      -77.8 - 78.9 -179.6 - 180
        25.9 - 67 -165.3 - -55.4
## 6
## 7
       40.1 - 67.8
                       -8.9 - 29.6
       49.8 - 53.1
                        8.6 - 13.9
## 8
      -28.3 - 67.9 -122.4 - 153.4
## 10 -27.7 - -27.1 -48.5 - -48.3
```

Model fit per dataset Here we explore variance partiioning within each model

```
model_sum = read.csv("../outputs/model_summary.csv")
model compare df = NULL
for(a in seq(1,30,3)){
 df_tmp = model_sum[c(a:(a+2)),]
  df tmp2 = data.frame(
    code = df tmp$code[1],
   phy h = df tmp$tip h[3]/(df tmp$tip[3] + df tmp$tip h[3] + df tmp$gen[3]), #Contribution of phylogenetic covariance, relative to all
    spa h = df tmp$sit h[3]/(df tmp$sit[3] + df tmp$sit h[3] + df tmp$squ[3]), #Contribution of spatial covariance, relative to all spatial
   temp = df_tmp$obv_auto[3]/(df_tmp$obv[3] + df_tmp$obv_auto[3]),
    temp_phi = df_tmp$phi[3], #Rho correlation between abundances
   temp_v = sum(c(df_tmp$obv[3], df_tmp$obv_auto[3]),na.rm = T)/sum(c( #Proportion of variance captured by temporal model components
     df_tmp$fix[3],
     df_tmp$obv[3],
     df_tmp$obv_auto[3],
     df_tmp$tip[3],
     df_tmp$tip_h[3],
      df_tmp$gen[3],
     df_tmp$sit[3],
      df_tmp$sit_h[3],
     df tmp$squ[3],
```

```
df_tmp$sig[3]
),na.rm = T),
sit_v = sum(c(df_tmp\sit[3], df_tmp\sit_h[3], df_tmp\squ[3]),na.rm = T)/sum(c( #Proportion of variance captured by spatial model components
  df_tmp$fix[3],
  df_tmp$obv[3],
  df_tmp$obv_auto[3],
  df_tmp$tip[3],
  df_tmp$tip_h[3],
  df_tmp$gen[3],
  df_tmp$sit[3],
  df_tmp$sit_h[3],
  df_tmp$squ[3],
  df_tmp$sig[3]
),na.rm = T),
phy_v = sum(c(df_tmp$tip[3],df_tmp$tip_h[3], df_tmp$gen[3]),na.rm = T)/sum(c(#Proportion of variance captured by phylogentic model co
  df_tmp$fix[3],
  df_tmp$obv[3],
  df_tmp$obv_auto[3],
  df_tmp$tip[3],
  df_tmp$tip_h[3],
  df_tmp$gen[3],
  df_tmp$sit[3],
  df_tmp$sit_h[3],
  df_tmp$squ[3],
  df_tmp$sig[3]
),na.rm = T),
res_v = sum(c(df_tmp$sig[3]),na.rm = T)/sum(c( #Proportion of variance captured by residuals
  df_tmp$fix[3],
  df_tmp$obv[3],
  df_tmp$obv_auto[3],
  df_tmp$tip[3],
  df_tmp$tip_h[3],
  df_tmp$gen[3],
  df_tmp$sit[3],
  df_tmp$sit_h[3],
  df_tmp$squ[3],
  df_tmp$sig[3]
),na.rm = T),
```

```
fix_v = sum(c(df_tmp$fix[3]),na.rm = T)/sum(c( #Proportion of variance captured by fixed effect
    df_tmp$fix[3],
    df_tmp$obv[3],
    df_tmp$tip[3],
    df_tmp$tip_h[3],
    df_tmp$tip_h[3],
    df_tmp$stip_h[3],
    df_tmp$sit[3],
    df_tmp$sit[3],
    df_tmp$sit[3],
    df_tmp$sit_al,
    df_tmp$sit_al,
```

Proportion of variance captured by temporal componenents

## [1] 0.14

```
round(mean(model_compare_df$temp_v),2)

## [1] 0.24

round(sd(model_compare_df$temp_v),2)

## [1] 0.2

Proportion of variance captured by spatial componenents

round(mean(model_compare_df$sit_v, na.rm = T),2)
```

```
round(sd(model_compare_df$sit_v, na.rm = T),2)
## [1] 0.05
Proportion of variance captured by phylogenetic componenents
round(mean(model_compare_df$phy_v),2)
## [1] 0.08
round(sd(model_compare_df$phy_v),2)
## [1] 0.06
Proportion of variance captured by residual componenents
round(mean(model_compare_df$res_v),2)
## [1] 0.34
round(sd(model_compare_df$res_v),2)
## [1] 0.3
Proportion of variance captured by fixed effect componenents
round(mean(model_compare_df$fix_v),2)
## [1] 0.19
round(sd(model_compare_df$fix_v),2)
## [1] 0.15
```

Correlation between sequential abundance values

```
round(mean(model_compare_df$temp_phi),2)
## [1] 0.25
round(sd(model_compare_df$temp_phi),2)
## [1] 0.64
H2 of phylo component i.e. the amount of variance captured by the phylogeny relative to the combined variance captured by the phylogeny and
taxonomy
round(mean(model_compare_df$phy_h),2)
## [1] 0.32
round(sd(model_compare_df$phy_h),2)
## [1] 0.22
H2 of spatial componenests i.e. the amount of variance captured by space relative to the combined variance captured by the space and the hierarchical
space nesting
round(mean(model_compare_df$spa_h, na.rm = T),2)
## [1] 0.39
round(sd(model_compare_df$spa_h, na.rm = T),2)
## [1] 0.26
```

# Assumptions

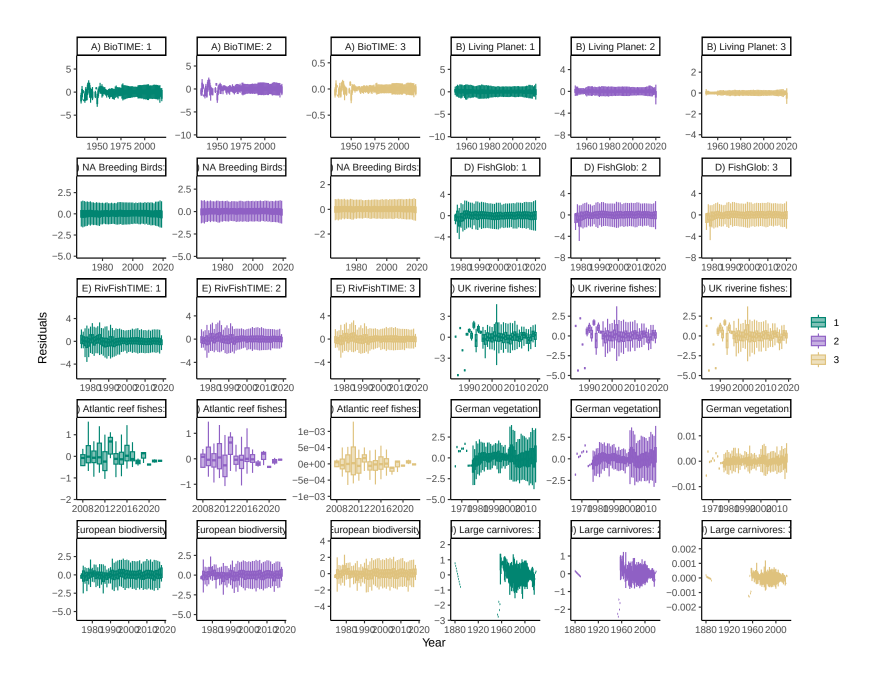
We calculate residuals by subtracting the true abundance values by the median predicted values. We also identify populations with particulary extreme residual values +- 2 standard deviations from the mean residual in each population.

```
analysis_list = readRDS("../data/derived_data/analysis_list.rds")
assumptions_df = NULL
for(a in names(analysis_list)[c(1:10)]){
  tmp_model = readRDS(paste0("../outputs/model_output_",a,".rds"))
  for(b in 1:3){
   if(b == 1){
     pred = tmp_model[[b]]$summary.fitted.values[,4]
     pred = head(pred,-100)
      true = analysis_list[[a]][[1]]$log_abundance
      true = head(true, -100)
      res = (true - pred)
   } else {
     pred = tmp_model[[b]]$summary.fitted.values[,4]
      pred = head(pred,-100)
     true = analysis_list[[a]][[1]]$cent_abundance
     true = head(true, -100)
      res = (true - pred)
    tmp_assumptions_df = data.frame(
      dataset = a,
      model = b,
     site_spec = head(analysis_list[[a]][[1]]$site_spec,-100),
     pred = pred,
      true = true,
     res = res,
     year = head(analysis_list[[a]][[1]]$date,-100))
    res_sum = tmp_assumptions_df %>%
      group_by(site_spec) %>%
      summarise(mean_res = mean(abs(res)), low_cut = mean(res) - (sd(res)*2), high_cut = mean(res) + (sd(res)*2))
    cut = quantile(res_sum$mean_res, probs = 0.5)
    res_sum$hq = ifelse((res_sum$mean_res < cut & res_sum$mean_res > -cut), "Keep", "Remove")
    tmp_assumptions_df = left_join(tmp_assumptions_df, res_sum)
    assumptions_df = rbind(assumptions_df, tmp_assumptions_df)
  }
}
dataset_rename = data.frame(
```

```
dataset = c("BioTIME", "CaPTrends", "EA_NFPD", "FishGlob", "LPI", "NAm_BBS", "Pilotto", "ReSurvey_Germany", "RivFish", "TimeFISH"),
  dataset_code = c("A) BioTIME", "J) Large carnivores", "F) UK riverine fishes", "D) FishGlob", "B) Living Planet", "C) NA Breeding Birds'
)
assumptions_df = left_join(assumptions_df, dataset_rename)
assumptions_df$model = as.character(assumptions_df$model)
assumptions_df$plot_code = pasteO(assumptions_df$dataset_code, ": ", assumptions_df$model)
```

Plot median residuals and quantiles against year (the independent variable) to assess heteroscedasaticy

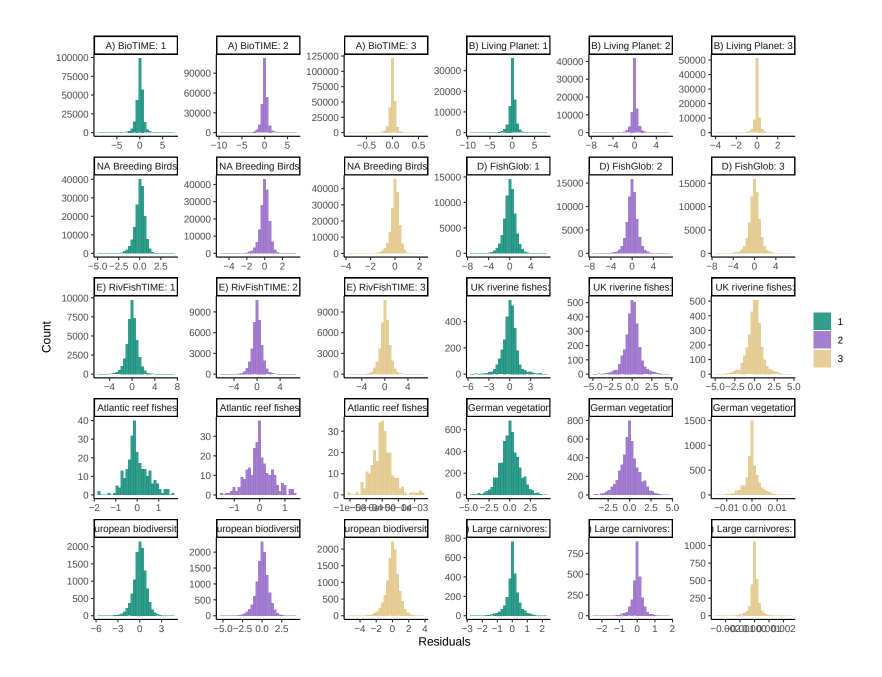
```
ggplot(data = assumptions_df) +
  geom_boxplot(aes(x = year, y = res, fill = model, colour = model, group = year), alpha = 0.5, outlier.shape = NA) +
  scale_fill_manual(values = c("#018571", "#8f60c4", "#dfc27d"), name = NULL) +
  scale_colour_manual(values = c("#018571", "#8f60c4", "#dfc27d"), name = NULL) +
  facet_wrap(plot_code~., scales = "free") +
  labs(x = "Year", y = "Residuals") +
  theme_classic()
```



```
ggsave("../outputs/figures/assumption_hetero.png", width = 13, height = 7, units = "in", device = "png")
```

Histogram of residuals for each dataset and model combinations to check for normality

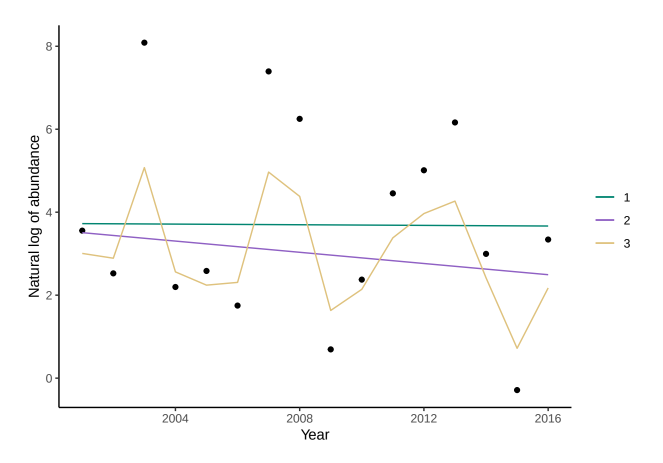
```
ggplot(data = assumptions_df) +
  geom_histogram(aes(x = res, fill = model), alpha = 0.8, position = "identity") +
  scale_fill_manual(values = c("#018571", "#8f60c4", "#dfc27d"), name = NULL) +
  facet_wrap(plot_code~., scales = "free") +
  labs(x = "Residuals", y = "Count") +
  theme_classic()
```



```
ggsave("../outputs/figures/assumption_normality1.png", width = 13, height = 7, units = "in", device = "png")
```

In some datasets, we see patterns of very heavy tails. These heavy tails tend to happen under two circumstances. 1) Example of population where the model poorly fits to cyclical and highly variable abudnance data

```
ggplot() +
  geom_point(data = assumptions_df[which(
    assumptions_df$site_spec == "Chen_2018_Yehliu power plant (site 2a)_25.209615_121.661948Siganus fuscescens_" & assumptions_df$model ==
  geom_line(data = assumptions_df[which(
    assumptions_df$site_spec == "Chen_2018_Yehliu power plant (site 2a)_25.209615_121.661948Siganus fuscescens_" & assumptions_df$model ==
  geom_line(data = assumptions_df[which(
    assumptions_df$site_spec == "Chen_2018_Yehliu power plant (site 2a)_25.209615_121.661948Siganus fuscescens_" & assumptions_df$model ==
  geom_line(data = assumptions_df[which(
    assumptions_df$site_spec == "Chen_2018_Yehliu power plant (site 2a)_25.209615_121.661948Siganus fuscescens_" & assumptions_df$model ==
  scale_colour_manual(values = c("#018571", "#8f60c4", "#dfc27d"), name = NULL) +
  labs(x = "Year", y = "Natural log of abundance") +
  theme_classic()
```

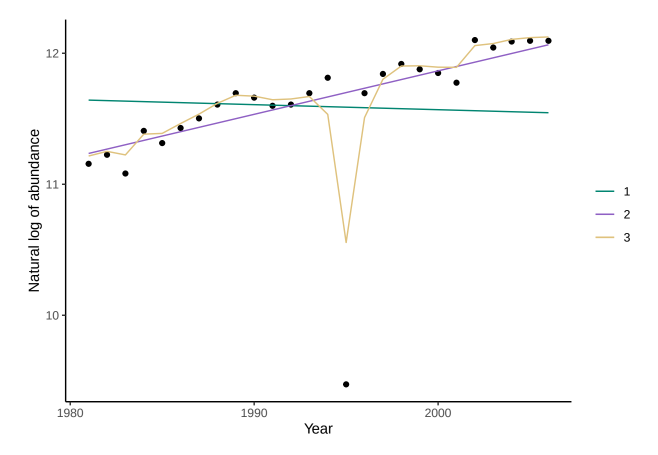


```
ggsave("../outputs/figures/assumption_poorfit1.png", width = 4, height = 4, units = "in", device = "png")
```

2) Example of population with generally well fitting data, but just one extreme abundance

```
ggplot() +
   geom_point(data = assumptions_df[which(
     assumptions_df$site_spec == "Rolley_2006_Western Forest, Wisconsin, USA_44.77778_-91.994720docoileus virginianus_" & assumptions_df$mc
   geom_line(data = assumptions_df[which(
     assumptions_df$site_spec == "Rolley_2006_Western Forest, Wisconsin, USA_44.77778_-91.994720docoileus virginianus_" & assumptions_df$mc
   geom_line(data = assumptions_df[which(
```

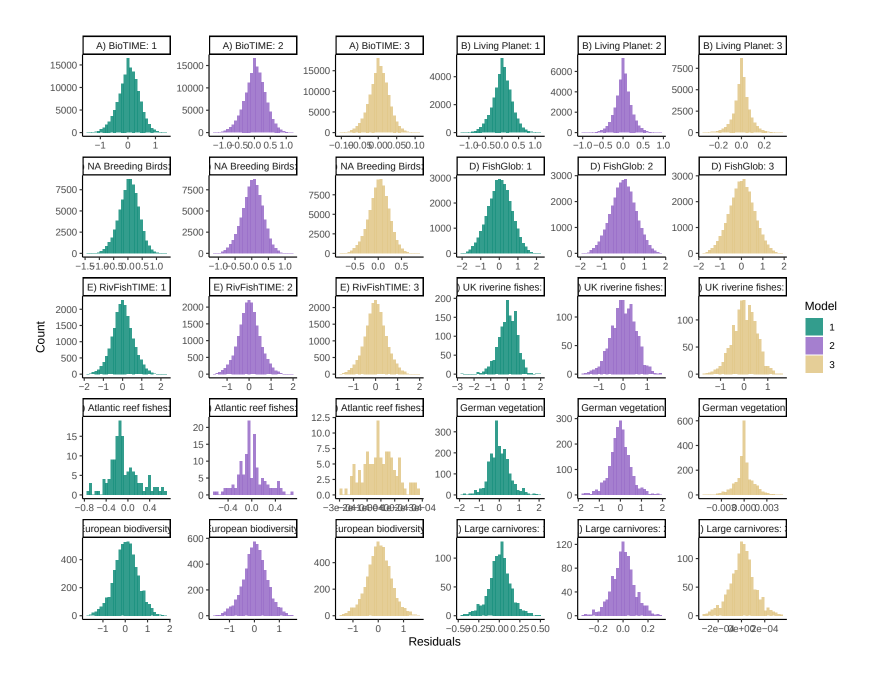
```
assumptions_df$site_spec == "Rolley_2006_Western Forest, Wisconsin, USA_44.77778_-91.994720docoileus virginianus_" & assumptions_df$mcgeom_line(data = assumptions_df[which(
    assumptions_df$site_spec == "Rolley_2006_Western Forest, Wisconsin, USA_44.77778_-91.994720docoileus virginianus_" & assumptions_df$mcscale_colour_manual(values = c("#018571", "#8f60c4", "#dfc27d"), name = NULL) +
labs(x = "Year", y = "Natural log of abundance") +
theme_classic()
```



ggsave("../outputs/figures/assumption\_poorfit2.png", width = 4, height = 4, units = "in", device = "png")

When we check the residuals after removing poorly fitting populations and extreme values, the residuals are looking far more normal

```
ggplot(data = assumptions_df[which(assumptions_df$hq == "Keep" & assumptions_df$res > assumptions_df$low_cut & assumptions_df$res < assumptions_df$res < assumptions_df$low_cut & assumptions_df$res < assumptions_df$low_cut & assumptions_df$res < assumptions_df$res < assumptions_df$low_cut & assumptions_df$flow_cut & assumptions_df$res < assumptions_df$low_cut & assumptions_df$flow_cut & assum
```



```
ggsave("../outputs/figures/assumption_normality2.png", width = 13, height = 7, units = "in", device = "png")
```

To ensure our inference is robust to heavy tails, we remove extrme values from the dataset with the heaviest tails (LPI) and re-model

```
lpi_trim = subset(assumptions_df, dataset == "LPI" & model == "3")
lpi_sub = head(analysis_list[["LPI"]][[1]],-100)
lpi_sub = cbind(lpi_sub, lpi_trim[,c(6,9,10,11)])
lpi_sub$log_abundance = ifelse(lpi_sub$res > lpi_sub$high_cut, NA, lpi_sub$log_abundance)
lpi_sub$log_abundance = ifelse(lpi_sub$res < lpi_sub$low_cut, NA, lpi_sub$log_abundance)
lpi_sub$log_abundance = ifelse(lpi_sub$hq == "Remove", NA, lpi_sub$log_abundance)
lpi_sub$cent_abundance = ifelse(lpi_sub$res > lpi_sub$high_cut, NA, lpi_sub$cent_abundance)
lpi_sub$cent_abundance = ifelse(lpi_sub$res < lpi_sub$low_cut, NA, lpi_sub$cent_abundance)
lpi_sub$cent_abundance = ifelse(lpi_sub$hq == "Remove", NA, lpi_sub$cent_abundance)
analysis_list_lpi = list(lpi_sub, analysis_list[["LPI"]][[2]], analysis_list[["LPI"]][[3]])
#source("modelling_lpi_assumption.R")</pre>
```

We detect the same pattern of an increasing standard deviation (from m1 to m2 to m3) around the collective trend. Inference appears robust

```
lpi_models = readRDS("../outputs/model_output_lpi_assumption.rds")
data.frame(
  model = c(1,2,3),
  uncertainty = c(
    lpi_models[[1]]$summary.fixed[2,2],
    lpi_models[[2]]$summary.fixed[2,2],
    lpi_models[[3]]$summary.fixed[2,2]))
```

```
## model uncertainty
## 1 1 0.0001956938
## 2 2 0.0048411846
## 3 3 0.0398701712
```

#### Figure 2

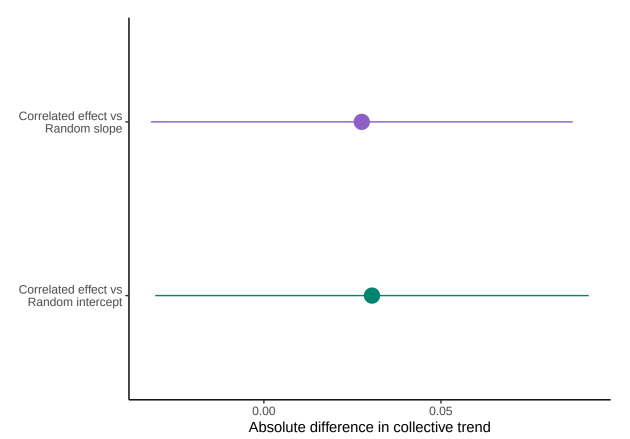
Load data from 'modelling core'

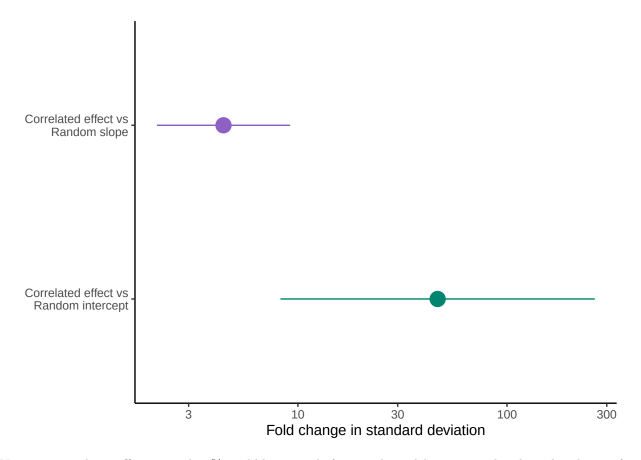
```
model_summary_core = read.csv("../outputs/model_summary.csv")
```

Assess average change in uncertainty and coefficient as you progress through the models

```
dataset_rename = data.frame(
     code = c("BioTIME", "CaPTrends", "EA_NFPD", "FishGlob", "LPI", "NAm_BBS", "Pilotto", "ReSurvey_Germany", "RivFish", "TimeFISH"),
     dataset code = c("A) BioTIME", "J) Large carnivores", "F) UK riverine fishes", "D) FishGlob", "B) Living Planet", "C) NA Breeding Birds'
model summary core = left join(model summary core, dataset rename)
stre = data.frame(
    name = c(rep("2",10), rep("1",10)),
    diff_rate = c((model_summary_core[which(model_summary_core$model == 1), "coef"]) - (model_summary_core[which(model_summary_core$model ==
                                        (model_summary_core[which(model_summary_core$model == 2), "coef"]) - (model_summary_core[which(model_summary_core] = 2), "coef"]) - (model_summary_core[which(model_summary_core] 
     diff_sd = c((model_summary_core[which(model_summary_core$model == 3), "coef_sd"])/(model_summary_core[which(model_summary_core$model == 1
                                   (model_summary_core[which(model_summary_core$model == 3),"coef_sd"])/(model_summary_core[which(model_summary_core$model == 2)
)
stre_sum = stre %>%
     dplyr::group by(name) %>%
     summarise(mn1 = mean(abs(diff rate)), sd1 = sd(abs(diff rate)), mn2 = mean(log10(diff sd)), sd2 = sd(log10(diff sd)))
m1 col = "#018571"
m2_{col} = "#8f60c4"
m3 col = "#dfc27d"
tmp_mod = lm(log(diff_sd) \sim 0 + name, data = stre)
exp(coef(tmp_mod))
##
                 name1
                                           name2
## 4.405152 46.575290
exp(confint(tmp_mod))
```

## 2.5 % 97.5 %





Next, we use the coefficients and 50% credible intervals from each model, against a baseline abundance of 100, and project abundance patterns for

each abundance datatset.

```
projections cmb = NULL
for(a in c(1:10)){
  cde = unique(model_summary_core$code)[a]
 df_tmp = subset(model_summary_core, code == cde)
 df_tmp2 = analysis_list[[cde]][[1]]
 run = length(min(df_tmp2$date, na.rm = T):max(df_tmp2$date, na.rm = T))-1
  y1 = c(100)
 v2 = c(100)
 y3 = c(100)
 for(tim in 1:(run)){
   y1 = c(y1,y1[tim]*(1 + df_tmp$coef[1]))
   y2 = c(y2,y2[tim]*(1 + df_tmp$coef[2]))
   y3 = c(y3, y3[tim]*(1 + df_tmp$coef[3]))
  lc1 = c(100)
  1c2 = c(100)
 1c3 = c(100)
  for(tim in 1:(run)){
   lc1 = c(lc1, lc1[tim]*(1 + df tmp$coef lc5[1]))
   1c2 = c(1c2,1c2[tim]*(1 + df_tmp$coef_1c5[2]))
   1c3 = c(1c3,1c3[tim]*(1 + df tmp$coef 1c5[3]))
  uc1 = c(100)
  uc2 = c(100)
  uc3 = c(100)
  for(tim in 1:(run)){
   uc1 = c(uc1, uc1[tim]*(1 + df_tmp$coef_uc5[1]))
   uc2 = c(uc2,uc2[tim]*(1 + df_tmp$coef_uc5[2]))
   uc3 = c(uc3,uc3[tim]*(1 + df_tmp$coef_uc5[3]))
  }
  projections_tmp = data.frame(
```

```
code = df_tmp$dataset_code[1],
    year = c(min(df_tmp2$date, na.rm = T):max(df_tmp2$date, na.rm = T)),
    y1 = y1, y2 = y2, y3 = y3,
   1c1 = 1c1, 1c2 = 1c2, 1c3 = 1c3,
    uc1 = uc1, uc2 = uc2, uc3 = uc3)
  projections cmb = rbind(projections cmb, projections tmp)
}
plt1 = ggplot(data = projections cmb[projections cmb$code %in% unique(projections cmb$code)[1],]) +
  geom line(aes(x = year, y1), colour = m1 col) +
  geom ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1 col) +
  geom_line(aes(x = year, y2), colour = m2_col) +
  geom_ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2_col) +
  geom line(aes(x = year, v3), colour = m3 col) +
  geom_ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3_col) +
  geom_hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
  scale_x_continuous(expand = c(0,0), breaks = c(1950,1980,2010)) +
  scale_y = c(0,0), breaks = c(10,100,1000) +
  coord_cartesian(xlim = c(1934, 2016), ylim = c(7, 2000)) +
  labs(x = " ", y = "Abundance") +
  theme classic() +
  theme(axis.text = element text(size=12))
plt2 = ggplot(data = projections_cmb[projections_cmb$code %in% unique(projections_cmb$code)[5],]) +
  geom line(aes(x = year, y1), colour = m1 col) +
  geom ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1 col) +
  geom line(aes(x = year, y2), colour = m2 col) +
  geom_ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2_col) +
  geom_line(aes(x = year, y3), colour = m3_col) +
  geom_ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3_col) +
  geom_hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
  scale_x_continuous(expand = c(0,0), breaks = c(1960,1985,2010)) +
  scale v log10(expand = c(0,0), breaks = c(20,100,500)) +
  coord_cartesian(ylim = c(10,600)) +
  labs(x = " ", y = "") +
```

```
theme classic() +
 theme(axis.title.y=element blank(),
       axis.text = element_text(size=12))
plt3 = ggplot(data = projections cmb[projections cmb$code %in% unique(projections cmb$code)[6],]) +
 geom line(aes(x = year, y1), colour = m1 col) +
 geom_ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1_col) +
 geom line(aes(x = year, y2), colour = m2 col) +
 geom ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2 col) +
 geom line(aes(x = year, y3), colour = m3 col) +
 geom ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3 col) +
 geom hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
 scale_x = c(1970, 1995, 2020), limits = c(1966, 2023) +
 scale v log10(expand = c(0,0), breaks = c(50,100,200)) +
 coord cartesian(vlim = c(30, 250)) +
 labs(x = " ", y = "") +
 theme classic() +
 theme(axis.title.y=element_blank(),
       axis.text = element text(size=12))
plt4 = ggplot(data = projections cmb[projections cmb$code %in% unique(projections cmb$code)[4],]) +
 geom line(aes(x = year, y1), colour = m1 col) +
 geom_ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1_col) +
 geom line(aes(x = year, y2), colour = m2 col) +
 geom ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2 col) +
 geom line(aes(x = year, y3), colour = m3 col) +
 geom ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3 col) +
 geom hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
 scale_x_continuous(expand = c(0,0), breaks = c(1980,1995,2010)) +
 scale v log10(expand = c(0,0), breaks = c(50,100,200)) +
 coord_cartesian(ylim = c(40,300)) +
 labs(x = " ", y = "") +
 theme classic() +
 theme(axis.title.y=element blank(),
       axis.text = element_text(size=12))
```

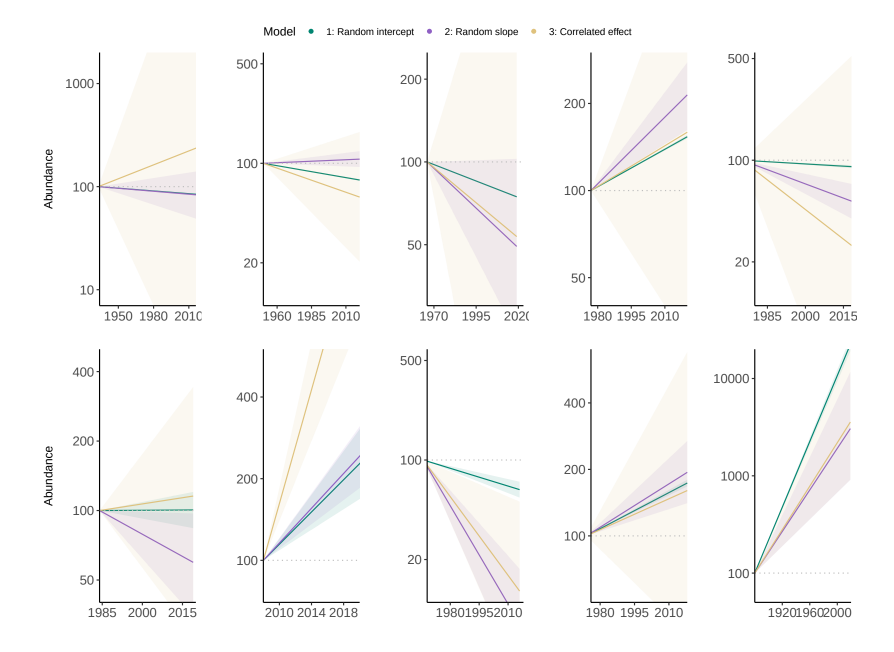
```
plt5 = ggplot(data = projections_cmb[projections_cmb$code %in% unique(projections cmb$code)[9].]) +
  geom line(aes(x = year, y1), colour = m1 col) +
  geom ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1 col) +
  geom line(aes(x = year, y2), colour = m2 col) +
  geom ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2 col) +
  geom line(aes(x = year, y3), colour = m3 col) +
  geom ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3 col) +
  geom hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
  scale x continuous(expand = c(0,0), breaks = c(1985,2000,2015)) +
  scale v log10(expand = c(0,0), breaks = c(20,100,500)) +
  coord_cartesian(xlim = c(1980, 2018), ylim = c(10, 550)) +
  labs(x = " ", y = "") +
  theme classic() +
  theme(axis.title.y=element blank(),
       axis.text = element_text(size=12))
plt6 = ggplot(data = projections cmb[projections cmb$code %in% unique(projections cmb$code)[3],]) +
  geom_line(aes(x = year, y1), colour = m1_col) +
  geom ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1 col) +
 geom_line(aes(x = year, y2), colour = m2_col) +
  geom ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2 col) +
 geom_line(aes(x = year, y3), colour = m3_col) +
  geom ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3 col) +
  geom hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
  scale x continuous(expand = c(0.0), breaks = c(1985,2000,2015)) +
  scale_y = c(0,0), breaks = c(50,100,200,400) +
  coord cartesian(xlim = c(1984,2020), ylim = c(40,500)) +
  labs(x = " ", y = "Abundance") +
  theme classic() +
  theme(axis.text = element text(size=12))
plt7 = ggplot(data = projections cmb[projections cmb$code %in% unique(projections cmb$code)[10],]) +
  geom_line(aes(x = year, y1), colour = m1_col) +
  geom ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1 col) +
```

```
geom_line(aes(x = year, y2), colour = m2_col) +
  geom ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2 col) +
  geom_line(aes(x = year, y3), colour = m3_col) +
  geom ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3 col) +
  geom_hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
  scale x continuous(expand = c(0,0), breaks = c(2010,2014,2018)) +
  scale y log10(expand = c(0,0), breaks = c(100,200,400)) +
  coord cartesian(xlim = c(2008, 2020), ylim = c(70, 600)) +
  labs(x = " ", y = "") +
  theme classic() +
  theme(axis.title.y=element blank(),
       axis.text = element text(size=12))
plt8 = ggplot(data = projections_cmb[projections_cmb$code %in% unique(projections_cmb$code)[8],]) +
  geom line(aes(x = year, y1), colour = m1 col) +
  geom_ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1_col) +
  geom_line(aes(x = year, y2), colour = m2_col) +
  geom_ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2_col) +
  geom_line(aes(x = year, y3), colour = m3_col) +
  geom_ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3_col) +
  geom_hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
  scale x continuous(expand = c(0,0), breaks = c(1980,1995,2010), limits = c(1968,2018)) +
  scale y log10(expand = c(0,0), breaks = c(20,100,500)) +
  coord cartesian(ylim = c(10,600)) +
  labs(x = " ", y = "") +
  theme classic() +
  theme(axis.title.y=element blank(),
        axis.text = element text(size=12))
plt9 = ggplot(data = projections_cmb[projections_cmb$code %in% unique(projections_cmb$code)[7],]) +
  geom_line(aes(x = year, y1), colour = m1_col) +
  geom_ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1_col) +
  geom_line(aes(x = year, y2), colour = m2_col) +
  geom_ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2_col) +
  geom_line(aes(x = year, y3), colour = m3_col) +
  geom_ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3_col) +
```

```
geom_hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
 scale x continuous(expand = c(0.0), breaks = c(1980.1995.2010), limits = c(1976.2018)) +
 scale_y_{log10}(expand = c(0,0), breaks = c(100,200,400)) +
 coord cartesian(vlim = c(50,700)) +
 labs(x = " ", y = "") +
 theme classic() +
 theme(axis.title.y=element blank(),
       axis.text = element text(size=12))
legend creator = data.frame(
 id = c("1","2","3"),
 x = 1000,
 y = 100
geom_line(aes(x = year, y1), colour = m1_col) +
 geom_ribbon(aes(x = year, ymin = lc1, ymax = uc1), alpha = 0.1, fill = m1_col) +
 geom_line(aes(x = year, y2), colour = m2_col) +
 geom_ribbon(aes(x = year, ymin = 1c2, ymax = uc2), alpha = 0.1, fill = m2_col) +
 geom_line(aes(x = year, y3), colour = m3_col) +
 geom_ribbon(aes(x = year, ymin = 1c3, ymax = uc3), alpha = 0.1, fill = m3_col) +
 geom_hline(aes(yintercept = 100), colour= "grey", linetype = "dotted") +
 geom point(data = legend creator, aes(x = x, y = y, colour = id)) +
 scale_x_continuous(expand = c(0,0), breaks = c(1920,1960,2000), limits = c(1880,2020)) +
 scale y log10(expand = c(0,0), breaks = c(100,1000,10000)) +
 coord cartesian(ylim = c(50, 20000)) +
 labs(x = " ", y = "") +
 theme classic() +
 scale colour manual(name = "Model", labels = c(
   "1: Random intercept",
   "2: Random slope",
   "3: Correlated effect"
 ), values = c(
   m1_col,
   m2_col,
   m3_col
 )) +
```

```
theme_classic() +
theme(axis.title.y=element_blank(),
    axis.text = element_text(size=12),
    legend.position = "bottom")

ggarrange(plt1, plt2, plt3, plt4, plt5, plt6, plt7, plt8, plt9, plt10, ncol = 5, nrow = 2, common.legend = T, legend = "top", align = "hv"
```



#### #Figure 3

Here we plot the abundance observations and assoicated predictions for a series of species in a specific site. We plot this data under each model to show how each model fits to the data

```
example dat = readRDS("../outputs/model output BioTIME.rds")
df = readRDS("../data/derived data/analysis list.rds")[[1]][[1]]
pop level = data.frame(id = rownames(example dat[[1]]$summary.fitted.values))
pop_level$m1_pred = example_dat[[1]]$summary.fitted.values$`0.5quant`
pop_level$m1_pred_lc = example_dat[[1]]$summary.fitted.values$`0.025quant`
pop_level$m1_pred_uc = example_dat[[1]]$summary.fitted.values$`0.975quant`
pop_level$m2_pred = example_dat[[2]]$summary.fitted.values$`0.5quant`
pop_level$m2_pred_lc = example_dat[[2]]$summary.fitted.values$`0.025quant`
pop_level$m2_pred_uc = example_dat[[2]]$summary.fitted.values$`0.975quant`
pop_level$m3_pred = example_dat[[3]]$summary.fitted.values$`0.5quant`
pop_level$m3_pred_lc = example_dat[[3]]$summary.fitted.values$`0.025quant`
pop level$m3 pred uc = example dat[[3]]$summary.fitted.values$`0.975quant`
pop level = cbind(df, pop level)
obs by site species = pop level %>%
  group_by(site_code, species) %>%
  summarise(N = n())
trends by site = pop level %>%
  group_by(site_code, latitude, longitude, species) %>%
  summarise(N = n())
site_freq = as.data.frame(table(unique(pop_level[,c("site_code","latitude", "longitude", "species")])$site_code))
site_freq = subset(site_freq, Freq == 3)
#Select site 716 as a good example
trends_by_site = trends_by_site[trends_by_site$site_code %in% site_freq$Var1, ]
plt m1 a = ggplot(data = pop level[which(pop level$site code == 336),]) +
  geom line(aes(x = date, y = (exp(m1 pred)), colour = species)) +
  geom ribbon(aes(x = date, ymin = (exp(m1 pred lc)), ymax = (exp(m1 pred uc)), fill = species), alpha = 0.1) +
  geom_point(aes(x = date, y = exp(log_abundance), colour = species)) +
  scale x continuous(expand = c(0,0), breaks = c(1985, 1995, 2005), limits = c(1980, 2008)) +
```

```
scale_y_continuous(expand = c(0,0)) +
  coord cartesian(vlim = c(1,4000), xlim = c(1980, 2008)) +
  scale colour manual(
   name = "".
   labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
   values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
  scale fill manual(
   name = "",
   labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
   values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
  labs(x = "", y = "Abundance", title = "Random intercept") +
  theme classic() +
  theme(plot.title = element text(size=10, face="italic"))
plt_m2_a = ggplot(data = pop_level[which(pop_level$site_code == 336),]) +
  geom_line(aes(x = date, y = (exp(m2_pred + mean_log)), colour = species)) +
  geom_ribbon(aes(x = date, ymin = (exp(m2_pred_lc + mean_log)), ymax = (exp(m2_pred_uc + mean_log)), fill = species), alpha = 0.1) +
  geom_point(aes(x = date, y = exp(cent_abundance + mean_log), colour = species)) +
  scale_x = continuous(expand = c(0,0), breaks = c(1985, 1995, 2005), limits = c(1980, 2008)) +
  scale v continuous(expand = c(0,0)) +
  coord_cartesian(ylim = c(1,4000), xlim = c(1980, 2008)) +
  scale colour manual(
   name = "",
   labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
   values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
  scale fill manual(
   name = "".
   labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
   values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
  labs(x = "", y = "Abundance", title = "Random slope") +
  theme classic() +
  theme(plot.title = element text(size=10, face="italic"))
plt_m3_a = ggplot(data = pop_level[which(pop_level$site_code == 336),]) +
  geom_line(aes(x = date, y = (exp(m3_pred + mean_log)), colour = species)) +
  geom_ribbon(aes(x = date, ymin = (exp(m3_pred_lc + mean_log)), ymax = (exp(m3_pred_uc + mean_log)), fill = species), alpha = 0.1) +
  geom_point(aes(x = date, y = exp(cent_abundance + mean_log), colour = species)) +
```

```
scale_x_continuous(expand = c(0,0), breaks = c(1985, 1995, 2005), limits = c(1980, 2008)) +
scale_y_continuous(expand = c(0,0)) +
coord_cartesian(ylim = c(1,4000), xlim = c(1980, 2008)) +
scale_colour_manual(
    name = "",
    labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
    values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
scale_fill_manual(
    name = "",
    labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
    values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
labs(x = "", y = "Abundance", title = "Correlated effect") +
theme_classic() +
theme(plot.title = element_text(size=10, face="italic"))
```

Under the same site described above, we plot the predicted site level trend

```
sit_level = readRDS("../outputs/model_output_predict_BioTIME.rds")
df2 = readRDS("../data/derived_data/analysis_list_predict.rds")
sit level = data.frame(
 year = df2[[1]][[1]]$year_centre[c((nrow(sit_level[[1]]$summary.fitted.values)-99):nrow(sit_level[[1]]$summary.fitted.values))],
  mn1 = sit level[[1]] summary.fitted.values[c((nrow(sit level[[1]] summary.fitted.values)-99):nrow(sit level[[1]] summary.fitted.values)
 lc1 = sit level[[1]] summary.fitted.values[c((nrow(sit level[[1]] summary.fitted.values)-99):nrow(sit level[[1]] summary.fitted.values)
  uc1 = sit level[[1]] summary.fitted.values[c((nrow(sit level[[1]] summary.fitted.values)-99):nrow(sit level[[1]] summary.fitted.values)
  mn2 = sit level[[2]] summary.fitted.values[c((nrow(sit level[[1]] summary.fitted.values)-99):nrow(sit level[[1]] summary.fitted.values)
 1c2 = sit level[[2]] summary.fitted.values[c((nrow(sit level[[1]] summary.fitted.values)-99):nrow(sit level[[1]] summary.fitted.values)
  uc2 = sit_level[[2]] summary.fitted.values[c((nrow(sit_level[[1]] summary.fitted.values)-99):nrow(sit_level[[1]] summary.fitted.values)
  mn3 = sit_level[[3]] summary.fitted.values[c((nrow(sit_level[[1]] summary.fitted.values)-99):nrow(sit_level[[1]] summary.fitted.values)
 lc3 = sit_level[[3]] summary.fitted.values[c((nrow(sit_level[[1]] summary.fitted.values)-99):nrow(sit_level[[1]] summary.fitted.values)
  uc3 = sit_level[[3]] summary.fitted.values[c((nrow(sit_level[[1]] summary.fitted.values)-99):nrow(sit_level[[1]] summary.fitted.values)
plt_m1_b = ggplot() +
  geom_line(data = pop_level[which(pop_level$site_code == 336),], aes(x = date, y = (exp(m1_pred)), colour = species), alpha = 0.5) +
  geom line(data = sit level, aes(x = year+1980, y = exp(mn1 + 2)), colour = "black") +
  geom_ribbon(data = sit_level, aes(x = year+1980, ymin = exp(lc1 + 2), ymax = exp(uc1 + 2)), alpha = 0.2, fill = "black") +
  scale x continuous(expand = c(0,0), breaks = c(1985, 1995, 2005)) +
```

```
scale_y_continuous(expand = c(0,0)) +
  coord_cartesian(ylim = c(1,4000), xlim = c(1980, 2008)) +
  scale colour manual(
   name = "",
   labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
   values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
  scale fill manual(
   name = "",
   labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
   values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
 labs(x = "", y = "", title = "") +
  theme classic() +
  theme(plot.title = element text(size=10, face="italic"))
plt_m2_b = ggplot(data = pop_level[which(pop_level$site_code == 336),]) +
  geom\_line(aes(x = date, y = (exp(m2\_pred + mean\_log)), colour = species), alpha = 0.5) +
  geom_line(data = sit_level, aes(x = year+1995, y = exp(mn2+6)), colour = "black") +
 geom_ribbon(data = sit_level, aes(x = year+1995, ymin = exp(lc2+6), ymax = exp(uc2+6)), alpha = 0.2, fill = "black") +
  scale_x_{ontinuous}(expand = c(0,0), breaks = c(1985, 1995, 2005)) +
  scale_y = continuous(expand = c(0,0)) +
  coord_cartesian(ylim = c(1,4000), xlim = c(1980, 2008)) +
  scale_colour_manual(
   name = "",
   labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
   values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
  scale_fill_manual(
   name = "".
   labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
   values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
 labs(x = "", y = "", title = "") +
  theme classic() +
  theme(plot.title = element_text(size=10, face="italic"))
plt_m3_b = ggplot() +
  geom_line(data = pop_level[which(pop_level$site_code == 336),],aes(x = date, y = (exp(m3_pred + mean_log)), colour = species), alpha = (
  geom_line(data = sit_level, aes(x = year+1995, y = exp(mn3+6)), colour = "black") +
```

```
geom_ribbon(data = sit_level, aes(x = year+1995, ymin = exp(lc3+6), ymax = exp(uc3+6)), alpha = 0.2, fill = "black") +
scale x continuous(expand = c(0,0), breaks = c(1985, 1995, 2005)) +
scale_y_continuous(expand = c(0,0)) +
coord cartesian(vlim = c(1,4000), xlim = c(1980, 2008)) +
scale_colour_manual(
  name = "",
 labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
  values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
scale fill manual(
  name = "",
 labels = c("A. opacum", "A. talpoideum", "G. carolinensis"),
  values = c("#9fbdbf", "#1885d9", "#1c50d4")) +
labs(x = "", y = "", title = "") +
labs(x = "", y = "", title = "") +
theme_classic() +
theme(plot.title = element text(size=10, face="italic"))
```

Next we plot all site level trends to depict heterogeniety in spatial trend variation, and then display the global trend over this site level variability, indicating that random intercept and random slope models are a poor representation of this heterogeniety

```
site_level = example_dat[[1]]$summary.random$site_code[,c(1,4,5,6)]
site_level = cbind(site_level, example_dat[[2]]$summary.random$site_code[,c(4,5,6)])
site_level = cbind(site_level, example_dat[[3]]$summary.random$site_code2[,c(4,5,6)] + example_dat[[3]]$summary.random$site_code2[,c(4,5,6)]
colnames(site_level) = c("site_code", "m1_lc_a", "m1_med_a", "m2_uc_a", "m2_med_a", "m2_med_a", "m2_uc_a", "m3_lc_a", "m3_med_a", "m3_uc_a"
region_level = example_dat[[1]]$summary.random$region_code[,c(1,5)]
region_level = cbind(region_level, example_dat[[2]]$summary.random$region_code[,c(5)])
region_level = cbind(region_level, example_dat[[3]]$summary.random$region_code[,c(5)])
colnames(region_level) = c("region_code", "m1_med_b", "m2_med_b", "m3_med_b")
region_site_link = unique(df[,c("region_code", "site_code")])
region_site_link = left_join(region_site_link, region_level, by = "region_code")
region_site_link$m1_lc = region_site_link$m1_lc_a + region_site_link$m1_med_b
region_site_link$m1_med = region_site_link$m1_med_a + region_site_link$m1_med_b
region_site_link$m1_uc = region_site_link$m1_uc_a + region_site_link$m1_med_b
region_site_link$m2_lc = region_site_link$m2_lc_a + region_site_link$m1_med_b
```

```
region_site_link$m2_med = region_site_link$m2_med_a + region_site_link$m2_med_b
region_site_link$m2_uc = region_site_link$m2_uc_a + region_site_link$m2_med_b
region_site_link$m3_lc = region_site_link$m3_lc_a + region_site_link$m3_med_b
region site link$m3 med = region site link$m3 med a + region site link$m3 med b
region_site_link$m3_uc = region_site_link$m3_uc_a + region_site_link$m3_med_b
expand_reg_sit_link = unique(df[,c("site_code", "mean_log", "mean_year")])
cmb_df = NULL
for(a in 1:nrow(region_site_link)){
 mn_lg = mean(subset(expand_reg_sit_link, site_code == region_site_link$site_code[a])$mean_log)[1]
 tmp_df = data.frame(
    site_code = subset(expand_reg_sit_link, site_code == region_site_link$site_code[a])$site_code[1],
   year_centre = seq(-20, 20, length.out = 30),
   year_adj = seq(-20,20,length.out = 30) + subset(expand_reg_sit_link, site_code == region_site_link$site_code[a])$mean_year[1],
   mn lg = mn lg,
   v_1 = 2.847 + -0.002*c(seq(-20,20,length.out = 30)) + region site link$m1 med[a],
   y2 = 0 + (-0.0009 + \text{region site link} \text{m2 med}[a]) *c(seq(-20,20,length.out = 30)),
   v3 = 0 + (0.011 + region site link$m3 med[a])*c(seq(-20,20,length.out = 30)))
  cmb df = rbind(cmb df, tmp df)
glob_level = data.frame(
  year = df2[[1]][[1]]$year_centre[c((nrow(example_dat[[1])$summary.fitted.values)-99):nrow(example_dat[[1]]$summary.fitted.values))],
  mn1 = example_dat[[1]]$summary.fitted.values[c((nrow(example_dat[[1]]$summary.fitted.values)-99):nrow(example_dat[[1]]$summary.fitted.values)
 lc1 = example_dat[[1]] summary.fitted.values[c((nrow(example_dat[[1]] summary.fitted.values)-99):nrow(example_dat[[1]] summary.fitted.values)
 uc1 = example_dat[[1]] summary.fitted.values[c((nrow(example_dat[[1]] summary.fitted.values)-99):nrow(example_dat[[1]] summary.fitted.values)
  mn2 = example_dat[[2]] summary.fitted.values[c((nrow(example_dat[[1]] summary.fitted.values)-99):nrow(example_dat[[1]] summary.fitted.values)
 lc2 = example_dat[[2]] summary.fitted.values[c((nrow(example_dat[[1]] summary.fitted.values)-99):nrow(example_dat[[1]] summary.fitted.values)
 uc2 = example_dat[[2]]$summary.fitted.values[c((nrow(example_dat[[1]]$summary.fitted.values)-99):nrow(example_dat[[1]]$summary.fitted.values)
  mn3 = example dat[[3]] summary.fitted.values[c((nrow(example dat[[1]] summary.fitted.values)-99):nrow(example dat[[1]] summary.fitted.values)
```

```
lc3 = example_dat[[3]] summary.fitted.values[c((nrow(example_dat[[1]] summary.fitted.values)-99):nrow(example_dat[[1]] summary.fitted.values)
 uc3 = example_dat[[3]] summary.fitted.values[c((nrow(example_dat[[1]] summary.fitted.values)-99):nrow(example_dat[[1]] summary.fitted.values)
m1_col = "#018571"
m2 col = "#8f60c4"
m3_{col} = "#dfc27d"
plt_m1_c = ggplot() +
  geom line(data = cmb df,
              aes(x = year_centre+1980, y = exp(y1), group = site_code), alpha = 0.35, colour = "grey") +
  geom_line(data = glob_level, aes(x = year+1980, y = exp(mn1)), colour = "#018571", size = 1.5) +
  geom_ribbon(data = glob_level, aes(x = year+1980, ymin = exp(lc1), ymax = exp(uc1)), alpha = 0.4, fill = "#018571") +
  scale_x_continuous(expand = c(0,0), breaks = c(1965, 1980, 1995)) +
  scale_y_continuous(expand = c(0,0)) +
  coord_cartesian(ylim = c(1,500), xlim = c(1963, 1999)) +
  labs(x = "", y = "", title = "") +
  theme classic()
plt_m2_c = ggplot() +
  geom line(data = cmb df,
            aes(x = year_centre+1980, y = exp(y3+2.919), group = site_code), alpha = 0.35, colour = "grey") +
  geom line(data = glob level, aes(x = year+1980, y = exp(mn2+2.919)), colour = "#8f60c4", size = 1.5) +
  geom_ribbon(data = glob_level, aes(x = year+1980, ymin = exp(lc2+2.919), ymax = exp(uc2+2.919)), alpha = 0.4, fill = "#8f60c4") +
  scale x continuous(expand = c(0,0), breaks = c(1965, 1980, 1995)) +
  scale y continuous(expand = c(0,0)) +
  coord cartesian(ylim = c(1,500), xlim = c(1963, 1999)) +
  labs(x = "", y = "", title = "") +
  theme_classic()
plt_m3_c = ggplot() +
  geom line(data = cmb df,
            aes(x = year_centre+1980, y = exp(y3+2.919), group = site_code), alpha = 0.35, colour = "grey") +
  geom_line(data = glob_level, aes(x = year+1980, y = exp(mn3+2.919)), colour = "#dfc27d", size = 1.5) +
```

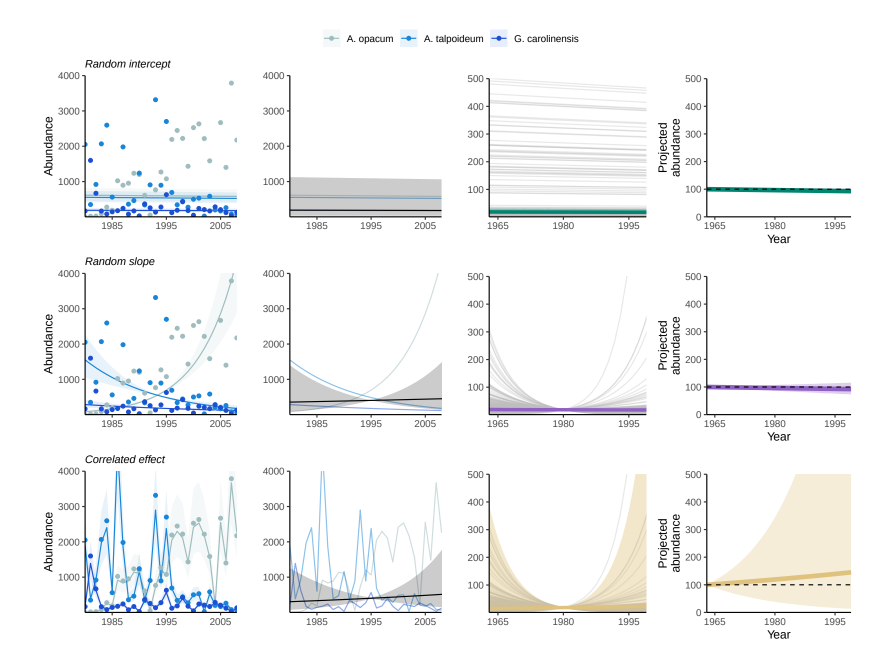
```
geom_ribbon(data = glob_level, aes(x = year+1980, ymin = exp(lc3+2.919), ymax = exp(uc3+2.919)), alpha = 0.4, fill = "#dfc27d") +
scale_x_continuous(expand = c(0,0), breaks = c(1965, 1980, 1995)) +
scale_y_continuous(expand = c(0,0)) +
coord_cartesian(ylim = c(1,500), xlim = c(1963, 1999)) +
labs(x = "", y = "", title = "") +
theme_classic()
```

We then take global level trend coefficients and 50% credible intervals to project abundance trends over time

```
ab 1c = c(100)
ab_mn = c(100)
ab uc = c(100)
for(a in c(1:36)){
 ab_lc = c(ab_lc,ab_lc[a]*(1 + inla.hpdmarginal(0.5, example_dat[[1])$marginals.fixed[[2]])[1]))
 ab_mn = c(ab_mn,ab_mn[a]*(1 + example_dat[[1]]$summary.fixed$`0.5quant`[2]))
 ab\_uc = c(ab\_uc, ab\_uc[a]*(1 + inla.hpdmarginal(0.5, example\_dat[[1]]\$marginals.fixed[[2]])[2]))
example dat proj m1 = data.frame(
 year = c(1963:1999),
 ab lc = ab lc,
 ab_mn = ab_mn,
  ab uc = ab uc
plt_m1_d = ggplot(example_dat_proj_m1) +
  geom\_line(aes(x = year, y = ab_mn), colour = "#018571", size = 2) +
  geom_ribbon(aes(x = year, ymin = ab_lc, ymax = ab_uc), alpha = 0.3, fill = "#018571") +
  geom_hline(aes(yintercept = 100), linetype = "dashed") +
  scale_y_continuous(expand = c(0,0)) +
  scale_x_{ontinuous}(expand = c(0,0), breaks = c(1965, 1980, 1995)) +
  coord_cartesian(ylim = c(0,500)) +
  theme_classic() +
  labs(x = "Year", y = "Projected\nabundance", title = "")
ab 1c = c(100)
```

```
ab mn = c(100)
ab uc = c(100)
for(a in c(1:36)){
  ab_lc = c(ab_lc,ab_lc[a]*(1 + inla.hpdmarginal(0.5, example_dat[[2]])*marginals.fixed([2]])[1]))
  ab_mn = c(ab_mn,ab_mn[a]*(1 + example_dat[[2]]$summary.fixed$`0.5quant`[2]))
  ab uc = c(ab uc,ab uc[a]*(1 + inla.hpdmarginal(0.5, example dat[[2]])*marginals.fixed([2]])(2)))
  #ab lc = c(ab \ lc, ab \ lc[a]*(1 + 0.05))
  \#ab \ mn = c(ab \ mn, ab \ mn[a]*(1 + 0))
  \#ab\ uc = c(ab\ uc, ab\ uc[a]*(1 + -0.05))
example_dat_proj_m2 = data.frame(
 year = c(1963:1999),
 ab_lc = ab_lc,
 ab_mn = ab_mn,
 ab_uc = ab_uc
plt_m2_d = ggplot(example_dat_proj_m2) +
  geom\_line(aes(x = year, y = ab_mn), colour = "#8f60c4", size = 2) +
  geom_ribbon(aes(x = year, ymin = ab_lc, ymax = ab_uc), alpha = 0.3, fill = "#8f60c4") +
  geom_hline(aes(yintercept = 100), linetype = "dashed") +
  scale v continuous(expand = c(0,0)) +
  scale x continuous(expand = c(0,0), breaks = c(1965, 1980, 1995)) +
  coord cartesian(ylim = c(0,500)) +
  theme classic() +
 labs(x = "Year", y = "Projected\nabundance", title = "")
ab 1c = c(100)
ab_mn = c(100)
ab uc = c(100)
for(a in c(1:36)){
  ab_lc = c(ab_lc,ab_lc[a]*(1 + inla.hpdmarginal(0.5, example_dat[[3])$ marginals.fixed([2]))(1)))
 ab_mn = c(ab_mn,ab_mn[a]*(1 + example_dat[[3])$summary.fixed$`0.5quant`[2]))
  ab\_uc = c(ab\_uc, ab\_uc[a]*(1 + inla.hpdmarginal(0.5, example_dat[[3])$marginals.fixed([2]))(2)))
  #ab_lc = c(ab_lc, ab_lc[a]*(1 + 0.05))
  \#ab \ mn = c(ab \ mn, ab \ mn[a]*(1 + 0))
```

```
\#ab\_uc = c(ab\_uc, ab\_uc[a]*(1 + -0.05))
example_dat_proj_m3 = data.frame(
 year = c(1963:1999),
  ab_lc = ab_lc,
  ab_mn = ab_mn,
  ab_uc = ab_uc
plt_m3_d = ggplot(example_dat_proj_m3) +
  geom\_line(aes(x = year, y = ab_mn), colour = "#dfc27d", size = 2) +
 geom_ribbon(aes(x = year, ymin = ab_lc, ymax = ab_uc), alpha = 0.3, fill = "#dfc27d") +
 geom_hline(aes(yintercept = 100), linetype = "dashed") +
  scale_y_continuous(expand = c(0,0)) +
  scale_x_{continuous}(expand = c(0,0), breaks = c(1965, 1980, 1995)) +
 coord_cartesian(ylim = c(0,500)) +
  theme_classic() +
 labs(x = "Year", y = "Projected\nabundance", title = "")
ggarrange(
 plt_m1_a,
 plt_m1_b,
 plt_m1_c,
 plt_m1_d,
 plt_m2_a,
 plt_m2_b,
 plt_m2_c,
 plt_m2_d,
 plt_m3_a,
  plt_m3_b,
  plt_m3_c,
  plt_m3_d,
 ncol =4, nrow = 3, common.legend = T)
```



#### Figure 4

Here we plot estimates of abundance change over space for one high profile species - American Robin. To calculate this abundance change in a given location, we sum: population trend + species trend + genus trend + phylogeny trend + site trend + region trend + spatial trend + overall trend. We then take uncertainty around spatial trends to estimate whether a trend would be significant at a selection of thresholds. This model is based on BioTIME data.

```
m3 = readRDS("../outputs/model output predict2 BioTIME.rds")
m3 = m3[[3]]
tmp_df = readRDS("../data/derived_data/analysis_list_predic2.rds")
tmp_df = tmp_df[[1]][[1]]
link_df = unique(tmp_df[,c("site_spec","site_spec_code","site_code","region_code","tips_code","genus_code", "lat_round", "lon_round", "spec_code","site_spec_code", "site_spec_code", "site_spec
link df = link df [c(12066:14976),]
tips code med = m3$summary.random$tips code[,c(1,2)]
colnames(tips code med) = c("ID", "val tips code mn")
link df = left join(link df, tips code med, by = c("tips code" = "ID"))
tips_code2_med = m3$summary.random$tips_code2[,c(1,2)]
colnames(tips code2 med) = c("ID", "val tips code2 mn")
link df = left join(link df, tips code2 med, by = c("tips code" = "ID"))
genus_code_med = m3$summary.random$genus_code[,c(1,2)]
colnames(genus_code_med) = c("ID", "val_genus_code_mn")
link_df = left_join(link_df, genus_code_med, by = c("genus_code" = "ID"))
site_code_med = m3$summary.random$site_code[,c(1,2)]
colnames(site code med) = c("ID", "val site code mn")
link_df = left_join(link_df, site_code_med, by = c("site_code" = "ID"))
site code2 med = m3$summary.random$site code2[,c(1,2)]
colnames(site code2 med) = c("ID", "val site code2 mn")
link df = left join(link df, site code2 med, by = c("site code" = "ID"))
site_spec_code_med = m3$summary.random$site_spec_code[,c(1,2,3)]
colnames(site spec code med) = c("ID", "val site spec code mn", "val site spec code sd")
site_spec_code_med$site_spec_qt400 = site_spec_code_med$val_site_spec_code_mn - (0.25*site_spec_code_med$val_site_spec_code_sd) #80%
site spec code med$site spec qt600 = site spec code med$val site spec code mn + (0.25*site spec code med$val site spec code sd)
```

```
site_spec_code_med$site_spec_qt300 = site_spec_code_med$val_site_spec_code_mn - (0.52*site_spec_code_med$val_site_spec_code_sd) #80%
site spec code med$site spec qt700 = site spec code med$val site spec code mn + (0.52*site spec code med$val site spec code sd)
site_spec_code_med$site_spec_qt200 = site_spec_code_med$val_site_spec_code_mn - (0.84*site_spec_code_med$val_site_spec_code_sd) #80%
site_spec_code_med\site_spec_qt800 = site_spec_code_med\sval_site_spec_code_mn + (0.84*site_spec_code_med\sval_site_spec_code_sd)
site_spec_code_med$site_spec_qt100 = site_spec_code_med$val_site_spec_code_mn - (1.28*site_spec_code_med$val_site_spec_code_sd) #80%
site spec code med$site spec qt900 = site spec code med$val site spec code mn + (1.28*site spec code med$val site spec code sd)
link df = left join(link df, site spec code med, by = c("site spec code" = "ID"))
region code med = m3$summary.random$region code[,c(1,2)]
colnames(region code med) = c("ID", "val region code mn")
link df = left join(link df, region code med, by = c("region code" = "ID"))
link df$pop trend sit mn =
  link_df$val_tips_code_mn +
  link_df$val_tips_code2_mn +
  link df$val genus code mn +
  link_df$val_site_code_mn +
  link_df$val_site_code2_mn +
  link_df$val_region_code_mn +
  m3\$summary.fixed\$mean[2]
link_df$pop_trend_sit_100 =
  link_df$val_tips_code_mn +
  link df$val tips code2 mn +
  link_df$val_genus_code_mn +
  link df$val site code mn +
  link_df$val_site_code2_mn +
  link df$site spec qt100 +
  link df$val region code mn +
  m3\$summary.fixed\$mean[2]
link_df$pop_trend_sit_200 =
  link_df$val_tips_code_mn +
  link_df$val_tips_code2_mn +
  link_df$val_genus_code_mn +
  link_df$val_site_code_mn +
  link_df$val_site_code2_mn +
  link_df$site_spec_qt200 +
```

```
link_df$val_region_code_mn +
  m3$summary.fixed$mean[2]
link_df$pop_trend_sit_300 =
  link_df$val_tips_code_mn +
  link_df$val_tips_code2_mn +
  link_df$val_genus_code_mn +
  link df$val site code mn +
  link_df$val_site_code2_mn +
  link_df$site_spec_qt300 +
  link_df$val_region_code_mn +
  m3$summary.fixed$mean[2]
link_df$pop_trend_sit_400 =
  link_df$val_tips_code_mn +
  link_df$val_tips_code2_mn +
  link_df$val_genus_code_mn +
  link_df$val_site_code_mn +
  link_df$val_site_code2_mn +
  link_df$site_spec_qt400 +
  link_df$val_region_code_mn +
  m3\$summary.fixed\$mean[2]
link_df$pop_trend_sit_600 =
  link_df$val_tips_code_mn +
  link_df$val_tips_code2_mn +
  link_df$val_genus_code_mn +
  link_df$val_site_code_mn +
  link_df$val_site_code2_mn +
  link_df$site_spec_qt600 +
  link_df$val_region_code_mn +
  m3$summary.fixed$mean[2]
link_df$pop_trend_sit_700 =
  link_df$val_tips_code_mn +
  link_df$val_tips_code2_mn +
  link_df$val_genus_code_mn +
  link_df$val_site_code_mn +
```

```
link_df$val_site_code2_mn +
  link df$site spec qt700 +
  link_df$val_region_code_mn +
  m3\$summary.fixed\$mean[2]
link df$pop trend sit 800 =
  link df$val tips code mn +
  link df$val tips code2 mn +
  link df$val genus code mn +
  link df$val site code mn +
  link_df$val_site_code2_mn +
  link_df$site_spec_qt800 +
  link_df$val_region_code_mn +
  m3\$summary.fixed\$mean[2]
link_df$pop_trend_sit_900 =
  link_df$val_tips_code_mn +
  link_df$val_tips_code2_mn +
  link_df$val_genus_code_mn +
  link_df$val_site_code_mn +
  link_df$val_site_code2_mn +
  link df$site spec qt900 +
  link df$val region code mn +
  m3\$summary.fixed\$mean[2]
link_df_sum = link_df
link_df_sum$cat = NA
link_df_sum$cat = ifelse(link_df_sum$pop_trend_sit_400 > 0, "+20%", link_df_sum$cat)
link_df_sum$cat = ifelse(link_df_sum$pop_trend_sit_300 > 0, "+40%", link_df_sum$cat)
link_df_sum$cat = ifelse(link_df_sum$pop_trend_sit_200 > 0, "+60%", link_df_sum$cat)
link_df_sum$cat = ifelse(link_df_sum$pop_trend_sit_100 > 0, "+80%", link_df_sum$cat)
link_df_sum$cat = ifelse(link_df_sum$pop_trend_sit_600 < 0, "-20%", link_df_sum$cat)</pre>
link_df_sum$cat = ifelse(link_df_sum$pop_trend_sit_700 < 0, "-40%", link_df sum$cat)</pre>
link_df_sum$cat = ifelse(link_df_sum$pop_trend_sit_800 < 0, "-60%", link_df_sum$cat)</pre>
link df sum$cat = ifelse(link df sum$pop trend sit 900 < 0, "-80%", link df sum$cat)
```

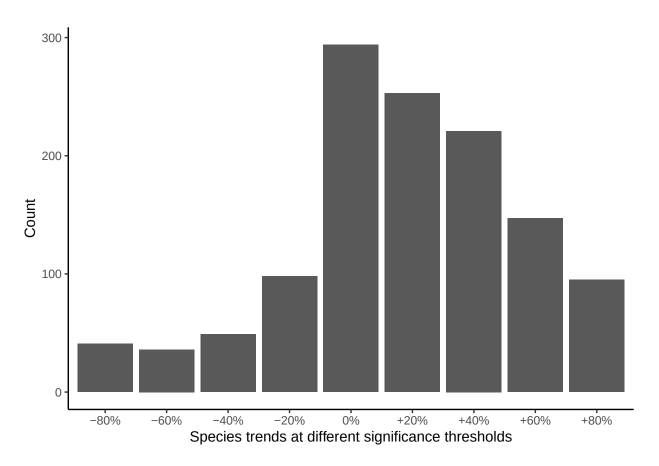
```
link_df_sum$cat = ifelse(is.na(link_df_sum$cat), "0%", link_df_sum$cat)
link_df_sum$cat = factor(link_df_sum$cat, levels = c("-80%", "-60%", "-40%", "-20%", "0%", "+20%", "+40%", "+60%", "+80%"))
link_df_sum$cat_num = as.numeric(link_df_sum$cat)
## Example SpatialPolygonsDataFrame
data(wrld simpl)
SPDF <- subset(wrld simpl, ISO3=="USA" | ISO3=="CAN" | ISO3=="MEX")
## Example RasterLayer
rst <- rasterFromXYZ(link df sum[,c("lon round", "lat round", "cat num")], crs=proj4string(SPDF))
## crop and mask
rst <- crop(rst, extent(SPDF))</pre>
rst <- mask(rst, SPDF)</pre>
rst_df <- as(rst, "SpatialPixelsDataFrame")</pre>
rst df <- as.data.frame(rst df)</pre>
colnames(rst_df) <- c("value", "x", "y")</pre>
world <- map_data("world")</pre>
plt_map = ggplot() +
  geom_tile(data=rst_df, aes(x=x, y=y, fill=as.factor(value)), alpha=0.8) +
  geom map(data = world, map = world, aes(long, lat, map id = region), fill = NA, colour = "black", alpha = 0.4) +
  #geom_polygon(data=SPDF, aes(x=long, y=lat, group=group),
                fill=NA, color="black", size=1) +
  coord_sf(xlim = c(-130, -60), ylim = c(20, 60)) +
  scale x continuous(expand = c(0,0)) +
  scale y continuous(expand = c(0,0)) +
  scale fill brewer(palette = "RdBu", labels = c("-80", "-60", "-40", "-20", "0", "+20", "+40", "+60", "+80"), name = "Confidence interval
  labs (x = "Longitude", y = "Latitude") +
  theme_classic()
```

Here we plot estimates of abundance change over a phylogeny by taking the sum of: species trend + genus trend + phylogeny trend + overall trend. We then take uncertainty around phylogeny trends to estimate whether a trend would be significant at a selection of thresholds. This model is based on BioTIME data

```
m3 = readRDS("../outputs/model_output_BioTIME.rds")
m3 = m3[[3]]
tmp_df = readRDS("../data/derived_data/analysis_list.rds")
tmp df = tmp df[[1]][[1]]
link_df = unique(tmp_df[,c("tips_code","genus_code","species")])
tips code med = m3$summary.random$tips code[,c(1,2)]
colnames(tips code med) = c("ID", "val tips code mn")
link_df = left_join(link_df, tips_code_med, by = c("tips_code" = "ID"))
tips code2 med = m3$summary.random$tips code2[,c(1,2,3,4,6)]
colnames(tips_code2_med) = c("ID","val_tips_code2_mn", "val_tips_code2_sd","lc","uc")
tips_code2_med$tip_qt400 = tips_code2_med$val_tips_code2_mn - (0.25*tips_code2_med$val_tips_code2_sd) #80%
tips_code2_med$tip_qt600 = tips_code2_med$val_tips_code2_mn + (0.25*tips_code2_med$val_tips_code2_sd)
tips_code2_med$tip_qt300 = tips_code2_med$val_tips_code2_mn - (0.52*tips_code2_med$val_tips_code2_sd) #80%
tips_code2_med$tip_qt700 = tips_code2_med$val_tips_code2_mn + (0.52*tips_code2_med$val_tips_code2_sd)
tips_code2_med$tip_qt200 = tips_code2_med$val_tips_code2_mn - (0.84*tips_code2_med$val_tips_code2_sd) #80%
tips_code2_med$tip_qt800 = tips_code2_med$val_tips_code2_mn + (0.84*tips_code2_med$val_tips_code2_sd)
tips_code2_med$tip_qt100 = tips_code2_med$val_tips_code2_mn - (1.28*tips_code2_med$val_tips_code2_sd) #80%
tips_code2_med$tip_qt900 = tips_code2_med$val_tips_code2_mn + (1.28*tips_code2_med$val_tips_code2_sd)
link df = left join(link df, tips code2 med, by = c("tips code" = "ID"))
genus code med = m3$summary.random$genus code[,c(1,2)]
colnames(genus code med) = c("ID", "val genus code mn")
link_df = left_join(link_df, genus_code_med, by = c("genus_code" = "ID"))
link df$pop trend tip mn =
  link df$val tips code mn +
  link df$val tips code2 mn +
  link_df$val_genus_code_mn +
  m3\$summary.fixed\$mean[2]
link_df$pop_trend_tip_100 =
  link_df$val_tips_code_mn +
  link df$tip qt100 +
  link_df$val_genus_code_mn +
  m3\$summary.fixed\$mean[2]
```

```
link_df$pop_trend_tip_200 =
 link_df$val_tips_code_mn +
  link_df$tip_qt200 +
 link_df$val_genus_code_mn +
 m3$summary.fixed$mean[2]
link df$pop trend tip 300 =
 link_df$val_tips_code_mn +
 link df$tip qt300 +
 link_df$val_genus_code_mn +
 m3$summary.fixed$mean[2]
link_df$pop_trend_tip_400 =
 link_df$val_tips_code_mn +
 link_df$tip_qt400 +
 link_df$val_genus_code_mn +
 m3$summary.fixed$mean[2]
link_df$pop_trend_tip_600 =
 link_df$val_tips_code_mn +
 link_df$tip_qt600 +
 link_df$val_genus_code_mn +
 m3$summary.fixed$mean[2]
link_df$pop_trend_tip_700 =
 link_df$val_tips_code_mn +
  link_df$tip_qt700 +
  link_df$val_genus_code_mn +
 m3$summary.fixed$mean[2]
link_df$pop_trend_tip_800 =
 link_df$val_tips_code_mn +
 link_df$tip_qt800 +
  link_df$val_genus_code_mn +
 m3$summary.fixed$mean[2]
link_df$pop_trend_tip_900 =
  link_df$val_tips_code_mn +
```

```
link_df$tip_qt900 +
  link df$val genus code mn +
  m3$summary.fixed$mean[2]
link df sum = link df
link df sum$cat = NA
link_df_sum$cat = ifelse(link_df_sum$pop_trend_tip_400 > 0, "+20%", link_df_sum$cat)
link df sum$cat = ifelse(link df sum$pop trend tip 300 > 0, "+40%", link df sum$cat)
link_df_sum$cat = ifelse(link_df_sum$pop_trend_tip_200 > 0, "+60%", link_df_sum$cat)
link df sum$cat = ifelse(link df sum$pop trend tip 100 > 0, "+80%", link df sum$cat)
link_df_sum$cat = ifelse(link_df_sum$pop_trend_tip_600 < 0, "-20%", link_df_sum$cat)</pre>
link_df_sum$cat = ifelse(link_df_sum$pop_trend_tip_700 < 0, "-40%", link_df_sum$cat)
link_df_sum$cat = ifelse(link_df_sum$pop_trend_tip_800 < 0, "-60%", link_df_sum$cat)</pre>
link_df_sum$cat = ifelse(link_df_sum$pop_trend_tip_900 < 0, "-80%", link_df_sum$cat)</pre>
link_df_sum$cat = ifelse(is.na(link_df_sum$cat), "0%", link_df_sum$cat)
link_df_sum$cat = factor(link_df_sum$cat, levels = c("-80%", "-60%", "-40%", "-20%", "0%", "+20%", "+40%", "+60%", "+80%"))
ggplot(link_df_sum) +
  geom_bar(aes(cat)) +
  labs(x = "Species trends at different significance thresholds", y = "Count") +
  theme classic()
```



```
spec_link = unique(tmp_df[,c("tips_code", "tips_chr")])
spec_link = subset(spec_link, !is.na(tips_chr))
link_df_sum2 = left_join(link_df_sum[,c("tips_code", "cat")], spec_link, by = "tips_code")
link_df_sum2 = subset(link_df_sum2, !is.na(link_df_sum2$tips_chr))
keep = unique(link_df_sum2$tips_chr)

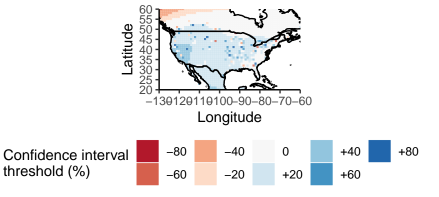
tr = readRDS("../data/derived_data/trees.rds")[[1]][[3]]
tr2 = keep.tip(tr, keep)
```

```
tr2$edge.length = log(tr2$edge.length) - min(log(tr2$edge.length)) + 0.01
tr2 = force.ultrametric(tr2)
## ***********************
## *
                          Note:
       force.ultrametric does not include a formal method to
## *
       ultrametricize a tree & should only be used to coerce
## *
      a phylogeny that fails is.ultramtric due to rounding --
       not as a substitute for formal rate-smoothing methods.
p = ggtree(tr2, layout = "circular")
plt_trends = data.frame(
  cat = link_df_sum2$cat
plt_trends$cat = as.factor(plt_trends$cat)
rownames(plt_trends) = link_df_sum2$tips_chr
plt clad = gheatmap(p, plt trends, width = 0.2, colnames = F) +
   scale fill brewer(palette = "RdBu", labels = c("-80", "-60", "-40", "-20", "0", "+20", "+40", "+60", "+80"), name = "Confidence interv
ggarrange(plt clad, plt map, ncol = 1, labels = c("A", "B"), common.legend = T, heights = c(1.2,0.65), legend = "bottom")
```

Α







#### Phylogeny - sensitivity analysis

Load data and develop a dataset summarising the three phylogeny models (A: Full - OTL, B: Restricted - OTL, C: Restricted - TimeTree). The OTL models use a phylogeny lacking branch lengths from the Open Tree of Life. The TimeTree phylogeny contains branch lengths, but has a 'restricted' taxonomic extent. In this restricted dataset, we only kept species occurring in both the Open Tree of Life and TimeTree.

```
model_summary_core = read.csv("../outputs/model_summary.csv")
model_summary_phylo = read.csv("../outputs/model_summary_phylo.csv")
```

```
model_summary_core = subset(model_summary_core, model == 3)
df_tmp = rbind(model_summary_core, model_summary_phylo)
df_tmp$phy_h = df_tmp$tip_h/(df_tmp$tip + df_tmp$tip_h + df_tmp$gen)
df_tmp$spa_h = df_tmp$sit_h/(df_tmp$sit + df_tmp$sit_h + df_tmp$squ)
df_tmp$temp = df_tmp$obv_auto/(df_tmp$obv + df_tmp$obv_auto)
df_tmp$temp_v = (df_tmp$obv + df_tmp$obv_auto)/(
  df_tmp$fix +
  df tmp$obv +
  df_tmp$obv_auto +
 df_tmp$gen +
  df_tmp$sit +
  df_tmp$sit_h +
 df_tmp$squ +
  df_tmp$sig)
df_tmp$sit_v = (df_tmp$sit + df_tmp$sit_h + df_tmp$squ)/(
  df_tmp$fix +
 df_tmp$obv +
 df_tmp$obv_auto +
 df_tmp$gen +
  df_tmp$sit +
  df_tmp$sit_h +
 df_tmp$squ +
  df_tmp$sig)
df_tmp$phy_v = (df_tmp$tip + df_tmp$tip_h + df_tmp$gen)/(
 df_tmp$fix +
  df tmp$obv +
  df_tmp$obv_auto +
  df tmp$gen +
 df_tmp$sit +
  df_tmp$sit_h +
 df_tmp$squ +
 df_tmp$sig)
df_tmp$res_v = (df_tmp$sig)/(
  df_tmp$fix +
 df_tmp$obv +
  df_tmp$obv_auto +
```

```
df_tmp$tip +
  df tmp$tip h +
  df_tmp$gen +
  df tmp$sit +
  df_tmp\sit_h +
  df tmp$squ +
  df_tmp$sig)
df_{tmp}fix_v = (df_{tmp}fix)/(
  df_tmp$fix +
  df tmp$obv +
  df_tmp$obv_auto +
  df_tmp$gen +
  df_tmp$sit +
  df_tmp$sit_h +
  df_tmp$squ +
  df_tmp$sig)
dataset rename = data.frame(
  code = c("BioTIME", "CaPTrends", "EA_NFPD", "FishGlob", "LPI", "NAm_BBS", "Pilotto", "ReSurvey_Germany", "RivFish", "TimeFISH"),
 dataset code = c("A) BioTIME", "J) Large carnivores", "F) UK riverine fishes", "D) FishGlob", "B) Living Planet", "C) NA Breeding Birds'
df tmp = left join(df tmp, dataset rename)
```

Collective trends and estimates of uncertainty for each of the model and dataset combinations

```
ggplot(data = df_tmp) +
    geom_pointrange(aes(x = model, ymin = coef_lc, y = coef, ymax = coef_uc, colour = model)) +
    scale_colour_manual(values = c("grey10", "grey50", "grey80"), labels = c("Full: OTL", "Restricted: OTL", "Restricted: TimeTree"), name = Nt
    scale_x_discrete(labels = c("A", "B", "C")) +
    labs(x = "", y = expression(beta)) +
    #coord_flip() +
    facet_wrap(dataset_code~., scales = "free") +
    theme_classic() +
    theme(legend.position = "bottom")
```

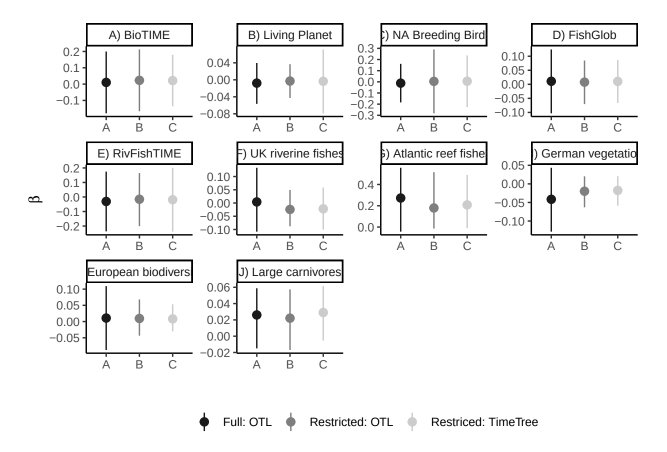
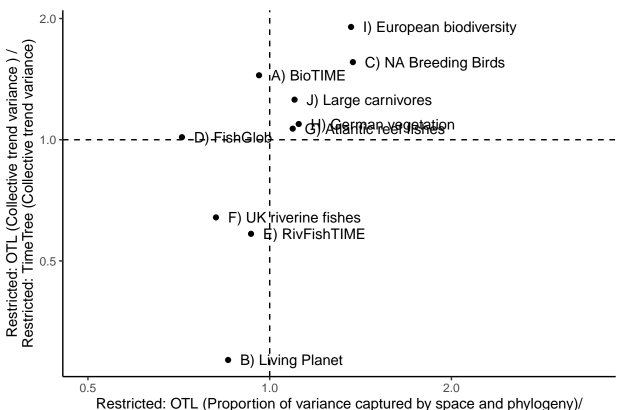


Figure describing how the porportion of variance captures by space and time and the driving forces behind the degree of uncertainty in the collective trend.

```
sd_div = df_rotl_trim$coef_sd^2/df_timetree_trim$coef_sd^2,
var = (df_rotl_trim$sit_v + df_rotl_trim$phy_v)/
    (df_timetree_trim$sit_v + df_timetree_trim$phy_v))

ggplot(data = df_compare) +
geom_point(aes(x = var, y = sd_div)) +
geom_text(aes(x = var, y = sd_div, label = name), hjust = 0, nudge_x = 0.02) +
scale_y_log10(breaks = c(0.5,1,2,4,8)) +
scale_x_log10(breaks = c(0.5,1,2,4,8), limits = c(0.5,3.4)) +
geom_hline(aes(yintercept = 1), linetype = "dashed") +
geom_vline(aes(xintercept = 1), linetype = "dashed") +
labs(x = "Restricted: OTL (Proportion of variance captured by space and phylogeny)/\nRestricted: TimeTree (Proportion of variance capture)
theme_classic()
```



# Restricted: OTE (Proportion of variance captured by space and phylogeny) Restricted: TimeTree (Proportion of variance captured by space and phylogeny)

### Structure - sensitivity analysis

Load data

```
model_summary_core = read.csv("../outputs/model_summary.csv")
model_summary_struc = read.csv("../outputs/summary_sensitivity_to_structure.csv")
```

Manipulate data. Renaming columns. Calculating the collective trend variance difference between the random slope model and all other models

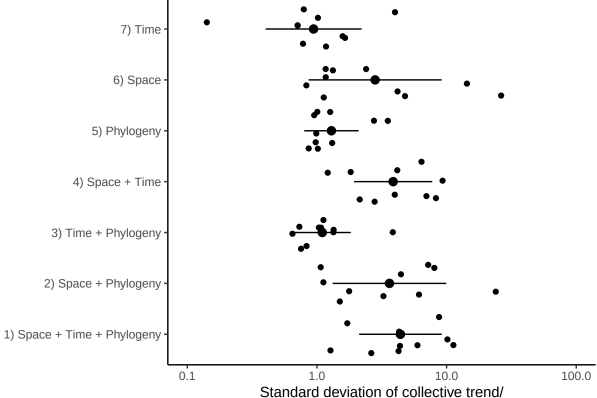
```
model_summary_core = subset(model_summary_core, model != 1, select = c(
  "X", "code", "model", "coef", "coef sd", "coef lc", "coef uc"
))
colnames(model summary core)[3] = "model type"
df tmp = rbind(model summary core, model summary struc)
df tmp = df tmp %>% mutate(model type=recode(model type,
                                    '3'='1) Space + Time + Phylogeny',
                                    'Time absent'='2) Space + Phylogeny',
                                    'Space absent'='3) Time + Phylogeny',
                                    'Phylogeny absent'='4) Space + Time',
                                    'Time and space absent'='5) Phylogeny',
                                    'Time and phylogeny absent'='6) Space',
                                    'Space and phylogeny absent'='7) Time',
                                    '2'='8) Random slope model'
                                    ))
df_{tmp2} = spread(df_{tmp}[,c(2,3,5)], model_{type}, coef_{sd})
df_tmp2$`1) Space + Time + Phylogeny` = df_tmp2$`1) Space + Time + Phylogeny`/ df_tmp2$`8) Random slope model`
df_tmp2$`2) Space + Phylogeny` = df_tmp2$`2) Space + Phylogeny`/ df_tmp2$`8) Random slope model`
df_tmp2$`3) Time + Phylogeny` = df_tmp2$`3) Time + Phylogeny`/ df_tmp2$`8) Random slope model`
df tmp2$`4) Space + Time` = df tmp2$`4) Space + Time`/ df tmp2$`8) Random slope model`
df tmp2$`5) Phylogeny` = df tmp2$`5) Phylogeny' / df tmp2$`8) Random slope model`
df tmp2$ 6) Space = df tmp2$ 6) Space / df tmp2$ 8) Random slope model
df tmp2$`7) Time` = df tmp2$`7) Time`/ df tmp2$`8) Random slope model`
df tmp3 = gather(df tmp2, model, sd div, `1) Space + Time + Phylogeny`: `7) Time`, factor key=TRUE)
```

Calucalte average variance difference and standard deviation

```
df_tmp4 = df_tmp3 %>%
  group_by(model) %>%
  summarise(mn = mean(log(sd_div)), sd = sd(log(sd_div)))
```

Plot variance differences

```
ggplot() +
    geom_jitter(data = df_tmp3, aes(y = model, x = sd_div)) +
    geom_pointrange(data = df_tmp4, aes(y = model, x = exp(mn), xmin = exp(mn-sd), xmax = exp(mn+sd))) +
    coord_cartesian(xlim = c(0.1,100)) +
    labs(x = "Standard deviation of collective trend/\n Standard deviation of the collective trend in the random slope model", y = "") +
    scale_x_log10() +
    theme_classic()
```



Standard deviation of collective trend/
Standard deviation of the collective trend in the random slope model

#### Prediction - trends

Here we take the predictions of missing trends in each dataset and estimate the median absolute error in preciction. This predictive error is then summarised across datasets, and then compared across models

```
mae full = NULL
for(a in names(analysis_list)[c(1:10)]){
  tmp_df = readRDS(paste0("../outputs/model_summary_miss_trends_", a,".rds"))
  tmp_mae_full = data.frame(
    code = a,
    mae2 = median(abs(abs(tmp_df$true - tmp_df$m2)/tmp_df$m2)*100),
    mae3 = median(abs(abs(tmp_df$true - tmp_df$m3)/tmp_df$m3)*100)
  mae_full = rbind(mae_full, tmp_mae_full)
mean(mae_full$mae2)
## [1] 90.50804
sd(mae_full$mae2)
## [1] 235.9675
median(mae_full$mae2)
## [1] 12.66181
mean(mae_full$mae3)
## [1] 16.29376
sd(mae_full$mae3)
## [1] 13.24278
```

```
median(mae_full$mae3)

## [1] 10.45509

mean(mae_full$mae2)/mean(mae_full$mae3)

## [1] 5.554767
```

#### Prediction - abundances

Here we take the predictions of missing abundance in each dataset and estimate the median absolute error in preciction. This predictive error is then summarised across datasets, and then compared across models

```
cmb_preds = NULL
for(a in names(analysis_list)[c(1:10)]){
    print(a)
    tmp_df_predict = readRDS(paste0("../outputs/model_output_miss_abundance_", a,".rds"))

tmp_preds = data.frame(
    code = a,
    log_abundance = tmp_df_predict[[4]]$log_abundance,
    mean_log = tmp_df_predict[[4]]$mean_log,
    miss = ifelse(is.na(tmp_df_predict[[4]]$log_abundance_trim),"Missing", "Complete"),
    pred1 = tmp_df_predict[[1]]$summary.fitted.values$mean,
    pred2 = tmp_df_predict[[2]]$summary.fitted.values$mean,
    pred3 = tmp_df_predict[[3]]$summary.fitted.values$mean
)
    cmb_preds = rbind(cmb_preds, tmp_preds)
}
```

```
## [1] "BioTIME"

## [1] "CaPTrends"

## [1] "EA_NFPD"

## [1] "FishGlob"

## [1] "LPI"
```

```
## [1] "NAm BBS"
## [1] "Pilotto"
## [1] "ReSurvey_Germany"
## [1] "RivFish"
## [1] "TimeFISH"
cmb_preds = subset(cmb_preds, miss == "Missing")
cmb_preds$pred2 = cmb_preds$pred2 + cmb_preds$mean_log
cmb_preds$pred3 = cmb_preds$pred3 + cmb_preds$mean_log
cmb_preds$diff1 = abs(abs(cmb_preds$log_abundance - cmb_preds$pred1)/cmb_preds$pred1)*100
cmb_preds$diff2 = abs(abs(cmb_preds$log_abundance - cmb_preds$pred2)/cmb_preds$pred2)*100
cmb_preds$diff3 = abs(abs(cmb_preds$log_abundance - cmb_preds$pred3)/cmb_preds$pred3)*100
sum_preds = cmb_preds %>%
  group by(code) %>%
 summarise(mn1 = median(diff1), mn2 = median(diff2), mn3 = median(diff3))
sum_preds$mn1 = signif(sum_preds$mn1, 3)
sum_preds$mn2 = signif(sum_preds$mn2, 3)
sum_preds$mn3 = signif(sum_preds$mn3, 3)
mean(sum_preds$mn1)
## [1] 30.295
sd(sum_preds$mn1)
## [1] 28.71189
median(sum_preds$mn1)
## [1] 20.35
mean(sum_preds$mn2)
## [1] 19.867
```

```
sd(sum_preds$mn2)
## [1] 12.93638
median(sum_preds$mn2)
## [1] 17.65
mean(sum_preds$mn3)
## [1] 16.508
sd(sum_preds$mn3)
## [1] 8.16333
median(sum_preds$mn3)
## [1] 17.35
mean(sum_preds$mn1)/mean(sum_preds$mn3)
## [1] 1.835171
mean(sum_preds$mn2)/mean(sum_preds$mn3)
## [1] 1.203477
```

## Driver of uncertainty

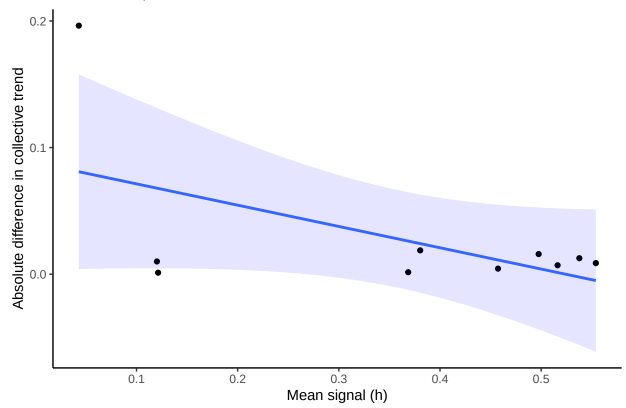
We now explore how the variance captured by spatial and phylogentic covariaince terms (relative to combined variance of covariance + hierarchical terms) impacts our inference

```
model_summary_core = read.csv("../outputs/model_summary.csv")
stre2 = data.frame(
  diff_rate = c((model_summary_core[which(model_summary_core$model == 2), "coef"]) - (model_summary_core[which(model_summary_core$model ==
  diff_sd = c((model_summary_core[which(model_summary_core$model == 3), "coef_sd"])/(model_summary_core[which(model_summary_core$model == 2
  var tip = rowSums((model summary core[which(model summary core$model == 3),c("tip", "gen", "tip h")])),
  var_phy = (model_summary_core[which(model_summary_core$model == 3),c("tip_h")]),
 var loc = rowSums((model summary core[which(model summary core$model == 3),c("sit", "squ", "sit h")])),
 var spa = (model summary core[which(model summary core$model == 3),c("sit h")])
stre2$phy_h = stre2$var_phy/stre2$var_tip
stre2$spa h = stre2$var spa/stre2$var loc
stre2$mean_h = rowMeans(stre2[,c("phy_h", "spa_h")])
summary(lm(abs(diff_rate) ~ mean_h, data = stre2))
##
## Call:
## lm(formula = abs(diff rate) ~ mean h, data = stre2)
## Residuals:
       Min
                  1Q Median
## -0.06658 -0.02016  0.00018  0.01321  0.11539
##
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.08814
                          0.03675
                                    2.399 0.0433 *
## mean h
              -0.16822
                          0.09099 -1.849 0.1017
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.05286 on 8 degrees of freedom
## Multiple R-squared: 0.2993, Adjusted R-squared: 0.2118
## F-statistic: 3.418 on 1 and 8 DF, p-value: 0.1017
confint(lm(abs(diff_rate) ~ mean_h, data = stre2))
                      2.5 %
                               97.5 %
##
```

```
## (Intercept) 0.003402222 0.17287770
## mean_h -0.378052728 0.04160454
```

```
ggplot(stre2) +
geom_point(aes(x = mean_h, y = abs(diff_rate))) +
geom_smooth(aes(x = mean_h, y = abs(diff_rate)), method = "lm", fill = "blue", alpha = 0.1) +
labs(x = "Mean signal (h)", y = "Absolute difference in collective trend", title = "Beta: -0.16; 95% CI: -0.38 - 0.04") +
theme_classic()
```

Beta: -0.16; 95% CI: -0.38 - 0.04



```
summary(lm(log10(diff_sd) ~ mean_h, data = stre2))
##
## Call:
## lm(formula = log10(diff_sd) ~ mean_h, data = stre2)
## Residuals:
       Min
                  1Q Median
                                           Max
## -0.25078 -0.12197 -0.04336 0.11665 0.38488
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.1906
                           0.1510 1.262 0.24247
## mean h
                1.2607
                           0.3739 3.371 0.00977 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.2172 on 8 degrees of freedom
## Multiple R-squared: 0.5869, Adjusted R-squared: 0.5353
## F-statistic: 11.37 on 1 and 8 DF, p-value: 0.009766
confint(lm(log10(diff sd) ~ mean h, data = stre2))
                    2.5 %
                            97.5 %
## (Intercept) -0.1576455 0.5388161
## mean h
                0.3983662 2.1229531
ggplot(stre2) +
  geom_point(aes(x = mean_h, y = (diff_sd))) +
  geom_smooth(aes(x = mean_h, y = (diff_sd)), method = "lm", fill = "blue", alpha = 0.1) +
  scale_y_log10() +
  labs(x = "Mean signal (h)", y = "Fold change in standard deviation", title = "Beta: 1.26; 95% CI: 0.39 - 2.12") +
  theme classic()
```

