### GLLVMs A-Å

Some inputs into performing a full analysis using GLLVMs

Audun Rugstad, Ph.D. candidate

Department of Mathematical Sciences, NTNU

2025-04-01



Intro

### Outline of the session

#### Five-step approach:

- 1. Formulate your question in terms of a statistical model
- 2. Data exploration and preparation
- Model fitting
- 4. Model checking
- 5. Making inferences

### Outline of the session

We'll go through the full analysis of an ecological dataset

**Example:** Dou et al. (2022): Influence of environmental variables on macroinvertebrate community structure in Lianhuan Lake



### Formulating the question

# Formulating the question

Formulate your research question in terms of a statistical analysis

First question: What is the goal of the analysis?

- ▶ **Prediction:** Find the model that best predict future observations
- Explanation: Investigate causal relationships between explanatory variables and response(s)

#### Prediction

#### Prediction

- ► Find the model that best predict future values of one/more response variables (i.e. species distribution)
- Variable selection based on optimizing for prediction
- ▶ I.e. forward/backward selection using AIC, Root mean square error etc.

**Example:** A model that predicts future distributions of wood beetles under different climate change scenarios



ro Formulating the question Data preparation Model setup Model checking Making inferences References occoordooodo occoordo

### **Explanation**

#### Two types:

#### Confirmatory

- ► Test a **specific, clear hypothesis** with one/a few models
- Predictor variables selected based on a priori knowledge
- Ideally no variable selection and pre-registration



Formulating the question | Data preparation | Model setup | Model checking | Making inferences | References | References

### Explanation

#### Two types:

#### Confirmatory

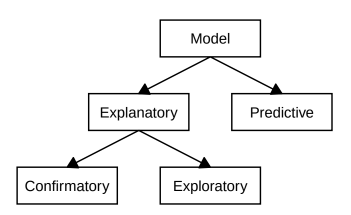
- ► Test a **specific, clear hypothesis** with one/a few models
- Predictor variables selected based on a priori knowledge
- Ideally no variable selection and pre-registration

#### Exploratory

- Use model(s) to explore possible causal relationships
- Should only be used to generate hypotheses
- Avoid automatic model selection



#### **Flowchart**





Exploratory "throwing everything at the wall" approaches perhaps most common

JSDMs more associated with prediction than other methods?

"True" confirmatory analyses are rare

▶ The line between the three can often be blurry

#### How can GLLVMs help?

- More streamlined and focused analysis
- Ideally, a single model of the community that is as complete as possible
- Can then be used to make inferences about different causal relationships etc.

**Q:** What type of analysis does a GLLVM with only **unconstrained** latent variables fit into?

**Q:** What type of analysis does a GLLVM with only **unconstrained** latent variables fit into?

**A:** Primarily exploratory, generating hypothesis about factors that structure the community.

*But:* if the goal is to test specific hypothesis about species co-occurrences, could potentially be used more confirmatory



**Q:** What type of analysis does a GLLVM with **constrained** latent variables fit into?



**Q:** What type of analysis does a GLLVM with **constrained** latent variables fit into?

A: All three, depending on approach?



Dou et al. (2022): Macroinvertebrates in 13 lakes in Lihuan, CN:

From the intro: The community composition of macro-invertebrate assemblages and their relationships with environmental variables were investigated

What type of modeling approach makes sense here?



Dou et al. (2022): Macroinvertebrates in 13 lakes in Lihuan, CN:

From the intro: The community composition of macro-invertebrate assemblages and their relationships with environmental variables were investigated

What type of modeling approach makes sense here?

**A:** Seems very clearly **exploratory** (which variables are potential drivers of community change?)



What should the model look like?

What should the model look like?

A concurrent ordination might be a good fit

- Allows for assessing the relative effect of many predictor variables at once, on a small number of latent variables
- Also allows us to account for potential residual variation in the L.V.s (i.e. variation not explained by the predictors)



### Example 2

Haven't yet found a good dataset/study that is not exploratory in the same way:(



Data preparation

### Important to always visually inspect key properties of the data before modeling

- Ensure the data meets the assumptions of the model
- Act as a "sanity check" for modeling output

Standardize and scale variables as needed

As with most other models, we recommend "common sense" practices and guidelines such as Zuur, leno, and Elphick (2010)



#### Some factors that can mess up a GLLVM:

- Predictor variables that are too co-linear
- Several species with very little information on abundances (i.e. zero-inflation)
- Sites with very little information on abundances
- Highly unbalanced sampling

# Model philosophy

The underlying philosophy of GLLVMs is:

- Data is ideally gathered with a spesific model or analysis in mind
- If not the case: tailor a model to the data properties
- ► The model should account for different properties of the data, as far as possible

The model should suit the data, not the other way around.

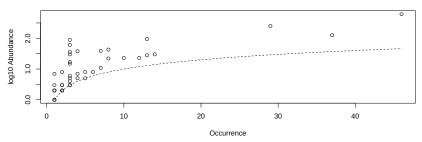


# Example: Lake invertebrates

```
# load data
lake_sp <- read.csv("../data/lake_inverts_sp.csv", sep = "\t")
lake_env <- read.csv("../data/lake_inverts_env.csv", sep = "\t")
# clean (according to paper)
lake_sp <- lake_sp |> select(-Site, -Group, -Season, -Lakes)
# reduce to actual counts!
lake_sp <- lake_sp / 16</pre>
```

### Example: Species

We can use a (Log-)abundance occupancy plot:



 $\mathbf{X} = \text{Number of plots a species occupies}, \mathbf{Y} = \text{average log-abundance}$ 

Can help identify outlier species + get a sense of the commonness of species in the dataset



# Species data

Table of total species abundances:

9 species with 1-2 occurrences in the dataset, and three species with very high abundances

# Filtering data

**Suggestion:** remove species occurring in only one plot, and plots containing only one species (remove 8 species and 9 plots).

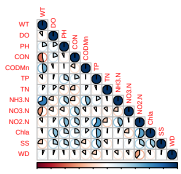
```
# remove rare species
lake_sp_filt <- lake_sp |>
    select_if(colSums(lake_sp !=0)>1)

# remove rare sites
lake_sp_filt <- lake_sp_filt[rowSums(lake_sp_filt != 0)>1,]

# remove same sites in env. dataframe
lake_env_filt <- lake_env[row.names(lake_sp_filt),]</pre>
```

### Environmental variables

#### Correlation of all predictors (lake chemical attributes):



\_10\_80\_60\_40 200 20 40 60 81



# Environmental variables

#### Why is this important?

- High co-linearity makes it hard for the model to converge (find estimates)
- Also makes the parameter uncertainties larger



### Environmental variables

#### What can be done?

- Omit predictors
  - ▶ Need to consider properties of the system and analysis
- "Model tricks"
  - Regularization ("reigning in" wild estimates)
  - ► Shrinkage (finding ways to make unimportant parameters -> 0)
  - Ex: Switching to random effects or putting constraints on the response distribution



# How high is too high?

Rule of thumb: no more than 0.8

Our max cor = 0.69

So we "just" standardize and scale to mean = 0 and variance = 1:

```
lake_{env}[,5:17] \leftarrow scale(lake_{env}[,5:17])
```

So all coefficient estimates are on the same magnitude



Model setup

# Model setup

Our suggestion: Concurrent ordination

What to include in the model?

- Our goal is to explore the impact of environmental variables
- As such, we can maybe argue for ignoring the study design here (differences between season/lake assumed to be manifested through the water properties)

### Model setup

#### Initial model setup:

- Two concurrent latent variables informed by the chemical properties of the water
- Random row effects (to account for site variation in overall abundance)
- Poisson distribution
- Could include quadratic effects, but might be overloading the model



### Model setup

### Model setup

A few (personal) tips for the gllvm() function:

- n.init: How many starting iterations you run
  - Depending on how long one run takes, maybe 5-50. Tells you when it has converged.
- trace = TRUE: tells you when each model run specified in n.init is complete (makes it less frustrating to fit)
- sd.errors = FALSE/TRUE: when diagnosing/comparing
  models, you can turn off estimation of standard errors to
  make each model fitting faster (and fit them retroactively
  using the se.gllvm() function)



### Model checking with GLLVMs

What to look at to determine whether the model is "good"?

- Diagnostic plots (plot() function) for model assumptions
  - Most important!
- AIC/BIC to compare model fit (e.g. number of LVs needed)
- Goodness of fit-tests (gllvm.goodnessOfFit())
- Very small/large parameter estimates (particularly sp. coefficients) for convergence



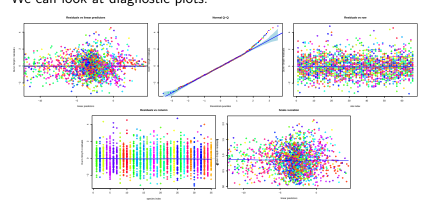
### Model checking with GLLVMs

### What to do to improve the fit?

- Change response distribution
- Run more iterations
- Change starting values
- Change from fixed to random effects
- Change number of LVs



# We can look at diagnostic plots:

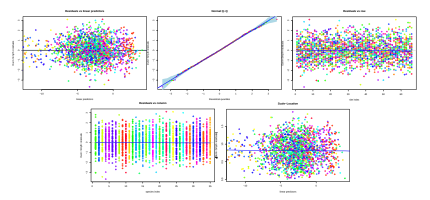


We see some fanning and potential overdispersion in the residual and QQ plots.



### Model checking: Example

After re-running the model with family = "negative.binomial":



The residual plots indicate that the NB model meets the



## Model checking: Example

The difference in **AICc** values between model 1 and 2 also indicates that the negative binomial model is better.

[1] 2918.967 2476.823



# Model checking: Example

Looking at some **goodness-of-fit** measures to the original data, however...

	${\tt Measure}$	Poisson	NB
1	cor	0.98382312	0.02060082
2	RMSE	0.78925359	65.12833567
3	MAE	0.28928256	2.24318316
4	MARNE	0.05016865	0.15774418

Looking at some **goodness-of-fit** measures to the original data, however...

	Measure	Poisson	NB
1	cor	0.98382312	0.02060082
2	RMSE	0.78925359	65.12833567
3	MAE	0.28928256	2.24318316
4	MARNE	0.05016865	0.15774418

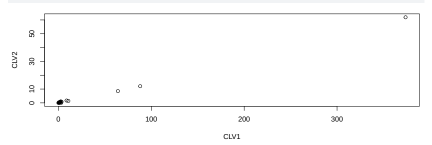
**Conclusion:** Our N.B. model, while meeting the model assumptions (a valid model), is a pretty bad fit to our data



# What has happened?

If we look at the plots of the standard errors for the species loadings, we also see that some are very large, indicating a poor convergence of the model:

plot(mod\_lakes\_nb\$sd\$theta) # theta = species coefficients



### What has happened?

If we look at the residual

[1] 3.23946e-07 1.56823e-08



### Improving the NB model

### We try two things:

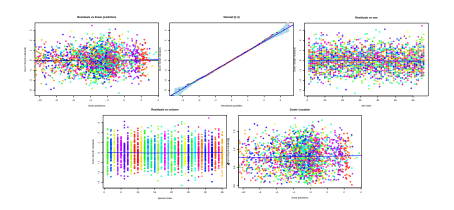
- Specify the LV-coefficients as random effects, drawn from a distribution unique for each coefficient
- ► Tell the model to estimate a common *dispersion* parameter for the negative binomial distribution for each species

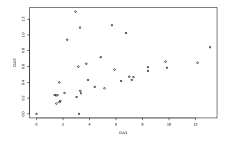
Both could help in steering the model away from "extreme" estimates and lessen overfitting (=regularizing)



# Improving the NB model

```
mod_lakes_nb_2 <- gllvm(lake_sp_filt, X = lake_env_filt,
                   lv.formula = ~WT + DO + PH + CON + CODMn +
                         TP + TN + NH3.N + NO3.N + NO2.N + Chla +
                         SS + WD.
                   row.eff = "random",
                   family = "negative.binomial",
                   ## LV coefficients are random effects:
                   randomB = "P",
                   ## Only one dispersion parameter estimated:
                   disp.formula = rep(1, ncol(lake_sp_filt)),
                   num.lv.c = 2
                   n.init = 20
```





```
NB1
                          NB2
Measure
         0.02060082 0.4658600
   RMSE 65.12833567 3.8379823
         2.24318316 0.8962227
         0.15774418 0.0670415
```

# Making inferences

What you look at depends on your question.

Typical visualizations:

- Model summary
- Biplots and triplots
- Variance explained
- Coefficient plots (caterpillar plots)



Two most important parts for us:

Residual standard deviation of LVs:

[1] 4.198412e-05 1.926389e-06

Very low, implies our predictors explain all the variance in the latent variables

We could go back and fit the model again without residual LV variation (as a "classic" constrained ordination)



# Model summary

### Coefficients LV predictors:

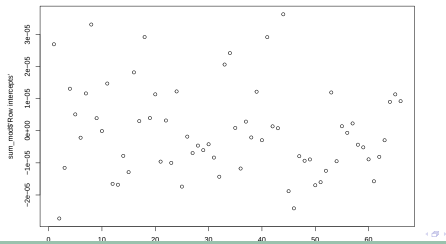
		204.2002.2		204120110212	
1	WT	0.2022	0.0409	0.1724	0.0
2	DO	0.0003	0.0000	0.0003	0.0
3	PH	0.2332	0.0544	0.1989	0.0
4	CON	0.0001	0.0000	0.0001	0.0
5	${\tt CODMn}$	0.0000	0.0000	0.0000	0.0
6	TP	0.1105	0.0122	0.0942	0.0
7	TN	0.0004	0.0000	0.0003	0.0
8	NH3.N	1.0835	1.1739	0.9239	0.8
9	NO3.N	0.0000	0.0000	0.0000	0.0
10	NO2.N	0.0005	0.0000	0.0004	0.0
11	Chla	1.7938	3.2179	1.5296	2.3
12	SS	0.0003	0.0000	0.0002	<b>4 △0 ≥</b> (

Name Std.Dev.CLV1 Variance.CLV1 Std.Dev.CLV2 Variance.

Audun Rugstad, Ph.D. candidate

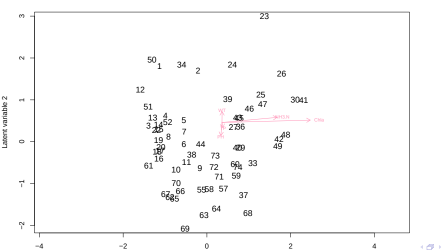
### Model summary

We can also look at the estimates for the random row effects:



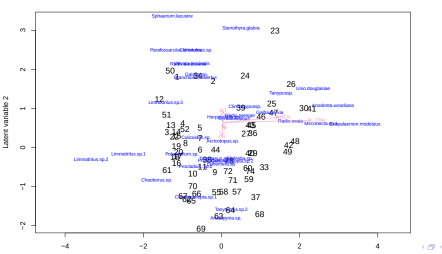
### Biplot and triplot

### Ordination (type='conditional')



### Biplot and triplot

#### Ordination (type='conditional')

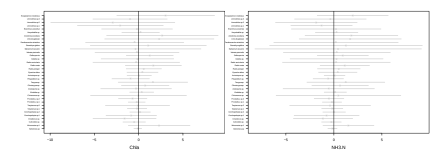


### Variance explained

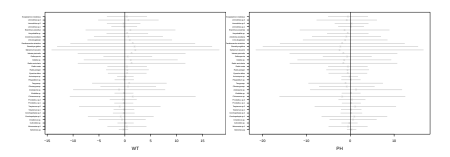
#### varPartitioning(mod\_lakes\_nb\_2)

```
Effect
                            Mean.explained.variance
CI.V · WT
                            20.1%
CI.V:DO
                             0.0%
CLV:PH
                            27.9%
CLV:CON
                             0.0%
CL.V: CODMn
                             0.0%
CLV:TP
                             5.2%
                             0.0%
CLV: TN
CLV: NH3.N
                            15.2%
CLV:NO3.N
                             0.0%
CLV:NO2.N
                             0.0%
CLV:Chla
                            31.5%
CLV:SS
                             0.0%
                             0.0%
CLV:WD
CI.V1
                             0.0%
CLV2
                             0.0%
Row random effect: sample
                             0.0%
```

### Coefficient plots



# Coefficient plots

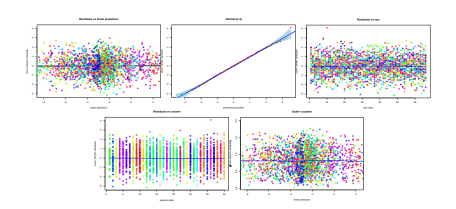


- While the model seems well behaved and reasonably fit to the data, estimates for the effect of predictors on the species are very high
- Suggests it it not very useful for explanatory purpose (only very weak evidence)

# Comparison to fully constrained model

To get a sense of the problem, we could compare it to the slightly more "shaved" model that the parameter estimates seem to suggest:

```
mod_lakes_nb_3 <- gllvm(lake_sp_filt, X = lake_env_filt,
                   lv.formula = ~WT + DO + PH + CON + CODMn +
                         TP + TN + NH3.N + NO3.N + NO2.N + Chla +
                         SS + WD.
                   ## Remove random row effects:
                   #row.eff = "random",
                   family = "negative.binomial",
                   ## LV coefficients are random effects:
                   randomB = "P",
                   ## Only one dispersion parameter estimated:
                   disp.formula = rep(1, ncol(lake_sp_filt)),
                   ## Fully constrained (reduced rank) coefs:
                   num.RR = 2.
                   n.init = 20)
```

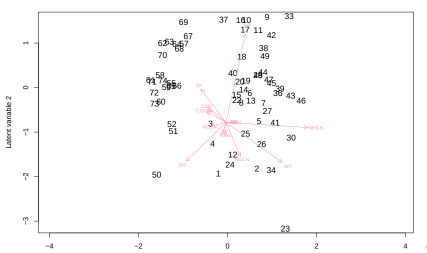




ro Formulating the question Data preparation Model setup Model checking **Making inferences** References

# Biplot (model 2)

### Ordination (type='marginal')



# Variance explained (model 2)

#### varPartitioning(mod\_lakes\_nb\_3)

```
Effect
          Mean.explained.variance
CLV:WT
          19.5%
CLV:DO
           7.6%
CL.V: PH
          32.9%
CLV: CON
           2.7%
CLV:CODMn
           2.1%
CL.V:TP
           8.8%
CL.V:TN
           0.5%
CLV:NH3.N 18.0%
CLV:NO3.N 0.5%
CLV:NO2.N
           6.2%
CLV:Chla
           0.7%
CLV:SS
           0.2%
CI.V:WD
           0.3%
```

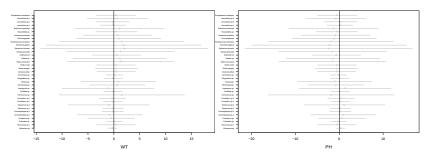
What has changed?

### What has changed?

The effect of Chlorophyll and NH3 content has **disappeared**!



# Coefficient plots (model 2)



Still extremely high uncertainty



Both models can be argued to be a reasonable, but suggest different drivers of species diversity.

- Still, some indication that PH and water temperature are important factors to look into (stood out in both models)
- Large uncertainties like this often a feature of negative binomial models

If the goal was prediction, we could have done automatic model selection or similar to (probably) get a simpler model with clearer estimates



Highlights the need to not trust model outputs blindly

The tools showed here can be useful in figuring out what is going on.

Next time: Try to collect more data in a way that could fit a Poisson model?

- Some unclearness about the actual sampling effort and pooling of samples in this study
- Unbalanced study design can have had an influence(more sampling in alkaline lakes than



### In other words:

► We can say both **that** more data is needed, and something about **why** more data is needed:)



GLLVMs A-Å

### References

### References

- Dou, Qianming, Xue Du, Yanfeng Cong, Le Wang, Chen Zhao,
   Dan Song, Hui Liu, and Tangbin Huo. 2022. "Influence of
   Environmental Variables on Macroinvertebrate Community
   Structure in Lianhuan Lake." *Ecology and Evolution* 12 (2): e8553. https://doi.org/10.1002/ece3.8553.
- Zuur, Alain F., Elena N. Ieno, and Chris S. Elphick. 2010. "A Protocol for Data Exploration to Avoid Common Statistical Problems." *Methods in Ecology and Evolution* 1 (1): 3–14. https://doi.org/10.1111/j.2041-210X.2009.00001.x.