Scale Uncertainty in ALDEx2

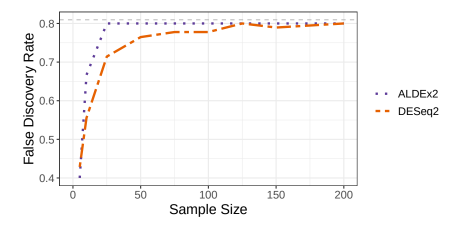
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May 13, 2024

Observed data (Y)	Sample 1	Sample 2	Sample 3	
Condition	Health	Health	Disease	Conclusion
Entity 1	5	10	100	Increase
Entity 2	10	25	3	Decrease
Entity 3	0	1	8	Increase
Entity 4	0	0	19	Increase
Sampling Depth	15	36	130	

System data (W)	Sample 1	Sample 2	Sample 3	
Condition	Health	Health	Disease	Conclusion
Entity 1	227	351	154	Decrease
Entity 2	684	891	3	Decrease
Entity 3	48	32	15	Decrease
Entity 4	43	39	27	Decrease
Scale (W [⊥])	1,002	1,313	200	

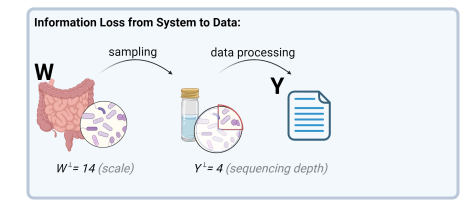
.. and lead to unacknowledged bias.



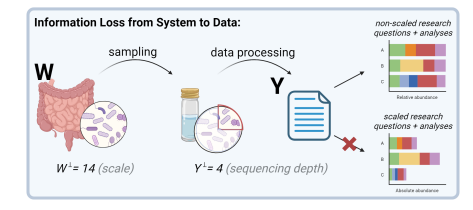
Section 1

Problem Set-Up

Observed Data as a Sample from the System



Observed Data as a Sample from the System



Notation

Problem Set-Up

• \mathbf{Y} : a measurement of the underlying system W.

$$\mathbf{W}_{dn} = \underbrace{\mathbf{W}_{dn}^{\parallel}}_{\text{composition}} \times \underbrace{\mathbf{W}_{n}^{\perp}}_{\text{scale}}$$

Problem Set-Up

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$$\mathbf{W}_{dn} = \underbrace{\mathbf{W}_{dn}^{\parallel}}_{\text{composition}} \times \underbrace{\mathbf{W}_{n}^{\perp}}_{\text{scale}}$$

- Compostion: $\mathbf{W}_{dn}^{\parallel} = \frac{\mathbf{W}_{dn}}{\sum_{d=1}^{D} \mathbf{W}_{dn}}$
- Scale: $W_n^{\perp} = \sum_{d=1}^{D} W_{dn}$

000000000000 Notation

Problem Set-Up

• \mathbf{Y} : a measurement of the underlying system W.

$$\mathbf{W}_{dn} = \underbrace{\mathbf{W}_{dn}^{\parallel}}_{\text{composition}} \times \underbrace{\mathbf{W}_{n}^{\perp}}_{\text{scale}}$$

• Compostion:
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- Scale: $W_n^{\perp} = \sum_{d=1}^D \mathbf{W}_{dn}$
- \bullet θ : what we want to estimate.

Example: Notation

System data (W)	Sample 1	Sample 2	Sample 3
Condition	Health	Health	Disease
Entity 1	0.27	0.27	0.77
Entity 2	0.68	0.68	0.02
Entity 3	0.05	0.02	0.08
Entity 4	0.04	0.03	0.13
Scale (W^{\perp})	1,002	1,313	200

Differential Abundance/Expression Analysis

 Question: How do entities (e.g., taxa or genes) change between conditions?

• In this case, θ is the log-fold change (LFC):

$$\theta_d = \mathsf{mean}_{\mathsf{case}}(\mathsf{log}\,\mathbf{W}_{dn}) - \mathsf{mean}_{\mathsf{control}}(\mathsf{log}\,\mathbf{W}_{dn})$$

The Original ALDEx2 Model

Scale Reliant Inference

Step 1: Model Sampling Uncertainty

$$\mathbf{Y}_{\cdot n} \sim \mathsf{Multinomial}(\mathbf{W}_{\cdot n}^{\parallel})$$

 $\mathbf{W}_{\cdot n}^{\parallel} \sim \mathsf{Dirichlet}(lpha)$

Step 2: Centered Log-Ratio Transformation

$$\log \mathbf{W}_{\cdot n} = \left[\log \mathbf{W}_{1n}^{\parallel} - \operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel}), ..., \log \mathbf{W}_{Dn}^{\parallel} - \operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel})\right]$$

Step 3: Calculate LFCs and Test if Different from Zero.

$$\theta_d = \mathsf{mean}_{\mathsf{case}}(\mathsf{log}\,\mathbf{W}_{dn}) - \mathsf{mean}_{\mathsf{control}}(\mathsf{log}\,\mathbf{W}_{dn})$$

Coding Changes to ALDEx2

Implied Assumptions about Scale

Problem Set-Up

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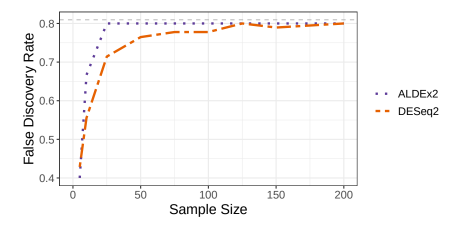
Implied Assumptions about Scale, cont.

Using the relationship $\mathbf{W}_{dn} = \mathbf{W}_{dn}^{\parallel} W_n^{\perp}$ and some math, the CLR normalization implies:

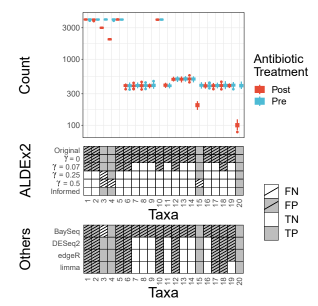
$$\log W_n^{\perp} = -\operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel}).$$

What does this mean?

Unacknowledged bias!



Adding Uncertainty in Scale can Help.



Scale Reliant Inference

- The CoDA perspective: Research questions that depend on W^{\perp} (scale) are not possible.
- The Normalization perspective: Research questions that depend on W^{\perp} (scale) can be answered after normalization.
- Who is right?

Scale Reliant Inference: The Basics

- The CoDA perspective: Research questions that depend on W^{\perp} (scale) are not possible.
- The Normalization perspective: Research questions that depend on W^{\perp} (scale) can be answered after normalization.
- Who is right?
- The CoDA perspective: Yes, but this is limiting in practice.
- The Normalization perspective: Not correct, but attempting to answer relevant questions.

Problem Set-Up

Scale Reliant Inference: The Basics

- What happens if θ depends on W^{\perp} ?
- Consider LFCs: how are taxa changing between two conditions?

$$egin{aligned} heta_d &= \mathsf{mean}_{\mathsf{case}}(\log \mathbf{W}_{dn}) - \mathsf{mean}_{\mathsf{control}}(\log \mathbf{W}_{dn}) \ &= ... \ &= \underbrace{\mathsf{mean}_{\mathsf{case}}(\log \mathbf{W}_{dn}^{\parallel}) - \mathsf{mean}_{\mathsf{control}}(\log \mathbf{W}_{dn}^{\parallel})}_{ heta^{\parallel}} \ &- \underbrace{\mathsf{mean}_{\mathsf{case}}(\log W_n^{\perp}) - \mathsf{mean}_{\mathsf{control}}(\log W_n^{\perp})}_{ heta^{\perp}} \end{aligned}$$

Scale Reliant Inference: Theory Intro

Recall for LFCs:

Problem Set-Up

$$egin{aligned} heta_d &= \mathsf{mean}_\mathsf{case}(\mathsf{log}\,\mathbf{W}_{dn}) - \mathsf{mean}_\mathsf{control}(\mathsf{log}\,\mathbf{W}_{dn}) \ &= heta^{\parallel} + heta^{\perp} \end{aligned}$$

• What can we say about θ from θ^{\parallel} alone?

Scale Reliant Inference: Theory Intro

Recall for LFCs:

Problem Set-Up

$$egin{aligned} heta_d &= \mathsf{mean}_\mathsf{case}(\log \mathbf{W}_{dn}) - \mathsf{mean}_\mathsf{control}(\log \mathbf{W}_{dn}) \ &= heta^\parallel + heta^\perp \end{aligned}$$

- What can we say about θ from θ^{\parallel} alone?
- Statistical perspective: θ is not identifiable without θ^{\perp} .
- Practical issues: unbiased estimators, calibrated confidence sets, and type-I error control **NOT** possible!
- See Nixon et al. (2023) for details.

Problem Set-Up

Goal: Estimate $\theta = f(\mathbf{W}^{\parallel}, W^{\perp})$.

- **①** Draw samples of \mathbf{W}^{\parallel} from a measurement model (can depend on \mathbf{Y}).
- ② Draw samples of W^{\perp} from a scale model (can depend on \mathbf{W}^{\parallel}).
- **3** Estimate samples of $\theta = f(\mathbf{W}^{\parallel}, W^{\perp})$.

Section 3

The Updated ALDEx2 Software

Coding Changes to ALDEx2

ALDEx2 as an SSRV

Step 1: Model Sampling Uncertainty

Scale Reliant Inference

$$\mathbf{Y}_{\cdot n} \sim \mathsf{Multinomial}(\mathbf{W}_{\cdot n}^{\parallel})$$

 $\mathbf{W}_{\cdot n}^{\parallel} \sim \mathsf{Dirichlet}(\alpha)$

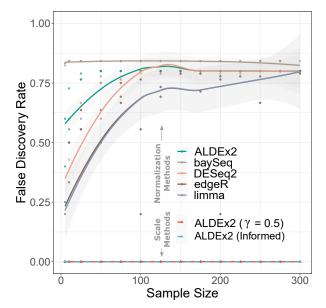
Step 2: Draw Samples from a Scale Model

$$\begin{split} \log W_n^\perp &= -\mathsf{mean}(\log \mathbf{W}_{\cdot n}^\parallel) + \epsilon, \ \epsilon \sim \mathit{N}(0, \gamma^2) \\ \log \mathbf{W}_{\cdot n} &= \log \mathbf{W}_{\cdot n}^\parallel + \log W_n^\perp \end{split}$$

Step 3: Calculate LFCs and Test if Different from Zero.

$$\theta_d = \text{mean}_{\text{case}}(\log \mathbf{W}_{dn}) - \text{mean}_{\text{control}}(\log \mathbf{W}_{dn})$$

Benefits of Moving Past Normalizations to Scale



Coding Changes to ALDEx2

Intro to Scale Models

Scale Reliant Inference

Normalizations are replaced by a scale model:

$$\log W_n^{\perp} = - \mathrm{mean}(\log \mathbf{W}_{\cdot n}^{\parallel}) + \epsilon$$
 $\epsilon \sim N(0, \gamma^2)$

What about other options?

There are no restrictions on what scale models can be, although there are some helpful options:

- Based on normalizations. (Stochastic normalizations)
- ② Based on biological knowledge.
- Based on outside measurements.

Scale Models based on Biological Knowledge

What do past studies or biological mechanisms tell about the scale of the system?

Problem Set-Up

What do past studies or biological mechanisms tell about the scale of the system?

- You are confident that taking an antibiotic will kill at least some microbes in the gut.
- A past study showed that a certain disease (e.g., Crohn's disease) leads to lower microbial load in the gut.
- You believe the total microbial load in the mouth changes after brushing your teeth.

This type of information can be used in scale model building.

Scale Models based on Outside Measurements

How can outside measurements be used to quantify scale?

Scale Models based on Outside Measurements

How can outside measurements be used to quantify scale?

- These measurements can be used *if* they relate to your scale of interest.
- 2 Examples include flow cytometry, qPCR, etc.
- Scale models can incorporate measurement uncertainty.

Section 4

Coding Changes to ALDEx2

Including scale

The new ALDEx2 model removes normalizations in lieu of scale models.

Including scale

The new ALDEx2 model removes normalizations in lieu of scale models.

Major updates:

- A new argument gamma which makes it easy to incorporate scale uncertainty.
- ② A new function aldex.senAnalysis to see how analysis results change as a function of scale uncertainty.

The gamma argument

- Added as argument to the aldex and aldex.clr function.
- gamma can either be a single numeric or a matrix.
 - Single numeric: controls the noise on the default scale model.
 - **2** Matrix: A $N \times S$ matrix of samples of W^{\perp} .
- gamma = NULL returns the default behavior of ALDEx2.

The default scale model is based on errors in the CLR normalization.

$$\log \hat{W}_n^{\perp(s)} = -\mathrm{mean}\left(\log \hat{W}_n^{\parallel(s)}\right) + \Lambda^{\perp} x_n$$

$$\Lambda^{\perp} \sim N(0, \gamma^2).$$

Advantages of the Default Scale Model

- 1 It is built off the status quo for ALDEx2.
- ② Any value of $\gamma>0$ will reduce false positives compared to the CLR normalization.
- It has a concrete interpretation to contextualize scale assumptions.

Interpreting the Default Scale Model

$$\log \hat{W}_n^{\perp(s)} = -\mathrm{mean}\left(\log \hat{W}_n^{\parallel(s)}\right) + \Lambda^{\perp} x_n$$

$$\Lambda^{\perp} \sim N(0, \gamma^2).$$

Empirical Rule: 95% of the samples of Λ^{\perp} fall within a factor of $\pm 2\gamma$ from zero.

Interpreting the Default Scale Model, cont.

$$\log \hat{W}_n^{\perp(s)} = -\text{mean}\left(\log \hat{W}_n^{\parallel(s)}\right) + \Lambda^{\perp} x_n$$
$$\Lambda^{\perp} \sim N(0, \gamma^2).$$

For case/control experiments:

- If $x_n = 1$: 95% of samples of $\log \hat{W}_n^{\perp(s)}$ fall within a factor of $\pm 2\gamma$ of the negative geometric mean.
- ② If $x_n = 0$: $\log \hat{W}_n^{\perp(s)}$ is equal to the negative geometric mean.

Interpreting the Default Scale Model, cont.

Recall that with the CLR normalization:

$$\log W_n^{\perp} = -\operatorname{mean}(\log \mathbf{W}_{\cdot n}^{\parallel}) = -\operatorname{GM}(\mathbf{W}_{\cdot n}^{\parallel}).$$

Thus, when using the CLR normalization:

$$\theta^{\perp} = \mathsf{mean}_{\mathsf{case}}(-\mathsf{GM}(\mathbf{W}_{\cdot n}^{\parallel})) - \mathsf{mean}_{\mathsf{control}}(-\mathsf{GM}(\mathbf{W}_{\cdot n}^{\parallel}))$$

This is same mean that the default scale model is centered on.

Interpreting the Default Scale Model, cont.

Taken together, the default scale model implies that:

- The value of θ^{\perp} is within $\pm 2\gamma$ of the value of θ^{\perp} implied by the CLR normalization.
- ② With 95% certainty, the true difference in scales falls within the the range $2^{\theta^{\perp}\pm2\gamma}$.

Option 2: More Complex Scale Models

Alternatively, can pass a matrix of scale samples to gamma so long as:

- The dimension is $N \times S$.
- 2 They are samples of W^{\perp} not $\log W^{\perp}$.

Reasons to do this:

- Biological beliefs: Scale is guided by the biological system or the researcher's prior beliefs.
- Outside Measurements: These can be used in building a scale model if they are informative on the scale of interest (e.g., qPCR, flow cytometry).

Sensitivity Analyses

- \bullet Recall that the default scale model has a parameter γ controlling the amount of noise added.
- Instead of picking γ , why not test over a range instead?
- Enter sensitivity analyses.

Sensitivity Analyses

Step 1: Model Sampling Uncertainty

Scale Reliant Inference

$$\mathbf{Y}_{\cdot n} \sim \mathsf{Multinomial}(\mathbf{W}_{\cdot n}^{\parallel})$$

 $\mathbf{W}_{\cdot n}^{\parallel} \sim \mathsf{Dirichlet}(lpha)$

Step 2: Draw Samples from a Scale Model For a given γ :

$$\log W_n^{\perp,\gamma} = -\text{mean}\left(\log \hat{W}_n^{\parallel(s)}\right) + \Lambda^{\perp} x_n$$

$$\Lambda^{\perp} \sim N(0, \gamma^2)$$

$$\log \mathbf{W}^{\gamma}_{\cdot n} = \log \mathbf{W}_n^{\parallel} + \log W_n^{\perp,\gamma}$$

Step 3: Calculate LFCs and Test if Different from Zero.

Step 4: Repeat for all desired values of γ .

Section 5

Data Examples

Simulation Study

Real Example: SELEX

Real Example: Vandputte



References

References

Scale Reliant Inference/Updates to ALDEx2:

- Nixon, et. al. (2023) "Scale Reliant Inference." ArXiv Preprint 2201.03616.
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- Fernandes et. al. (2014). "Unifying the analysis of high-throughput sequencing datasets: characterizing RNA-seq, 16S rRNA gene sequencing and selective growth experiments by compositional data analysis." *Microbiome*.

References

Problem Set-Up

Data Sources:

- McMurrough et. al. (2014)."Control of catalytic efficiency by a coevolving network of catalytic and noncatalytic residues." PNAS.
- Vandputte et. al. (2017). "Quantitative microbiome profiling links gut community variation to microbial load." *Nature*.