$\label{thm:continuous} \mbox{Title: Soil redox potential as a predictor of endobenthos species composition in an intertidal zone }$ 

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# Soil redox potential as a predictor of endobenthos species composition in an intertidal zone

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### Abstract

Soil redox potential measurements can be done without much physical effort in a short amount of time, whereas measuring endobenthos species diversity involves more effort and time. This study explored if soil redox potential can be used as a sole predictor of the abundance of endobenthos. Both variables were measured at an intertidal zone at Schiermonnikoog at a transect from salt marsh to mudflat. Inundation was expected to be influential in species composition, and to correct for this, species abundances and redox values were obtained from two depths at each site. It was found that redox potential alone cannot be used as predictor of endobenthos composition.

## Introduction

Species composition is one of the first and most vital pieces of data a field-based ecological research needs to gather. Any way to reliably conclude the same information with less work will save researchers both time and resources. One piece of information that is easy to obtain is a soil its redox potential. Redox potential is not just a passive abiotic factor; it is a value of biological importance. Among others, redox potential can be a measure of oxygen in the soil, and it is known that plant roots can raise the oxygen level in the soil (and thus its redox potential) [Blossfeld et al. 2011]. This positive correlation, however, is not always correct, because at least one other study finds a negative correlation [Dong et al. 2014]. As far as I know, using redox potential as a predictor of bethos species being present or not, has not yet been investigated.

As redox potential is positively correlated with soil oxygen concentration, and all animals need oxygen for respiration, I expect some benthos animals to respond to soil redox potential. The most likely candidates are animals that live in the soil and breathe through their skin, as these animals might choke from a too low oxygen concentration and will probably move to a habitat with a higher oxygen concentration instead. Additionally, it might be that

microbenthos serving as prey aggregaates at a certain oxygen concentration, which might cause its predator to follow this distribution.

The hypothesis tested is if soil redox potential can be used as a sole predictor of endobenthos species abundance, when assuming species distribute themselves normally around a certain redox potential. This study makes a strong case that this hypothesis should be rejected.

### Material and methods

This study was carried out at the intertidal zone of Schiermonnikoog. In the Southwest of this island, a 2400 meter long transect was set up, from salt marsh to mudflat. The elevations of the transect range from 270 cm to -80 cm NAP. All measurements were done at September 9th and 10th. At those days, inundation times of the sample sites ranged from from 1-80% of a tidal period.

Soil samples of 20 cm deep were taken at different distances. The top 5 cm was separated. Both parts of the soil sample were scored for species.

Redox values were measured by a potentiometer using 4 platinum-tip electrodes and a solution of KCl as a reference. The electrodes were put in at three depths: 2, 5 and 10 cm, in this sequence. The potential read is the value that remained constant, when placing or changing the electrodes. The values read were transformed to use earth as a reference point, using the formula  $V=1.8847 \cdot V_{measured}-53.201$ .

In this study, the redox potential at 2 cm depth is coupled to the benthos species diversity at depth 0-5 cm, where the potential at 10 cm depth is coupled to the diversity of depth 5-20 cm.

Of all collected species, only species with at least 3 individuals at both depths were taken into account. The minimum value of 3 was chosen, because it is the minimum number to test for normality. The requirement for a species to be at two depths is to disrupt the effect of inundation, as there can be similar redox potentials for different inundation times.

For each species, a Shapiro-Wilk normality test was used to determine if abundance is distributed normally around a certain redox potential. This test is chosen, as it has the best power for a given significance [Razali & Wah 2011].

The script to analyze the data is written in R and can be downloaded at https://github.com/richelbilderbeek/EvoEcoResearchCourse2014, or viewed in the appendix.

### Results

The redox potentials measured can be seen in figure.

863 individuals of 18 different benthos species were collected at the site (see figure). Of these species, 8 species had at least 3 individuals at both depths. Out of these 8 species, only 4 could be used, as not all sites were measured for redox potential. From the 4 species left, only the 2 species occurring at multiple

redox potentials were analyzed. These two species were Hydrobia ulvae and Nereis diversicolor. Figure shows the abundance of both species at different redox potentials. A Shapiro-Wilk normality test shows that both species have a significant probability of not following a normal distribution ( $p_{Hydrobia} < 0.001$ ,  $p_{Nereis} < 0.05$ , see table for exact values).

# Discussion

This study makes a strong case that soil redox potential cannot be used to predict species abundances, for both *Hydrobia ulvae* and *Nereis diversicolor*.

As *Hydrobia ulvae* is an epibenthic grazer [Newell 1965], it seems rather obvious that is not influenced by the oxygen level of the soil underneath it. Less obvious is that individuals were found in benthos 5 cm below the surface. This finding appears not to be an experimental error, as *Hydrobia ulvae* is found in deeper soil samples at multiple distances. It might be that the *Hydrobia ulvae* found in the deeper soil were not individuals, but only the shells left.

Nereis diversicolor creates burrows in the mud and is a predator and scavanger [Witte & Wilde 1979]. As it does not live in the soil itself, nor does it feed on something in the soil itself, it is not surprising that also this species is unaffected by soil redox potential.

Coupling a redox potential at a single depth to species abundances at a range of depths may have been too much of a simplification. Redox potential changes when probing at different depths in the soil, but whether this change is monotonous was unknown. After drawing the conclusions, the change of redox potential was analyzed for its orderedness. Because in the experiments also the redox potential at 5 cm deep was measured, it could be tested if redox potential changes monotonously for depths 2, 5 and 10 cm. It was found that this was the case in 10 out of 13 locations (see table). Thus, in 3 out of 13 cases, the redox potential at intermediate depth was the heighest or lowest value measured at that location. There can be two explanations for this unexpected pattern: (1) the soil redox potential is a complex abiotic variable that does not follow a monotonic change, or (2) the noise in the redox measurement is higher than the change in 'true' redox potential between depths.

## References

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Table 1: All 18 species and the number of individuals found per species per depth.

Species name	Depth: 2 cm	Depth: 10 cm
Arenicola marina	12	1
Bathy pore ia	2	0
$Carcinus\ maenas$	3	14
$Cerastoderma\ edule$	5	22
$Crassostrea\ gigas$	4	6
$Eteone\ long a$	0	5
$Gammarus\ locusta$	1	3
$Hemigropsus\ takanoi$	4	11
$Heteromastus\ filliform is$	1	13
$Hydrobia\ ulvae$	131	369
$Lanice\ conchilega$	2	31
$Littorina\ littorea$	11	51
$Macoma\ balthica$	0	31
$Mytilus\ edulis$	7	78
$Nere is\ diversicolor$	14	10
$Nere is\ virens$	3	0
$Scoloplas \ armiger$	1	16
$Scrobicularia\ plana$	1	0

Table 2: Shapiro-Wilk normality test of the species abundances on redox potential. n: number of individuals. p: chance the species abundances do not follow a normal distribution for a redox potential.

$_{ m Name}$	$\mathbf{n}$	p	$\operatorname{significance}$
Hydrobia ulvae	294	< 2.2e-16	***
$Nere is\ diversicolor$	9	0.04965	*

Table 3: the way the redox potentials are ordered when measuring a redox potential at 2, 5 and 10 cm in this order. LTH: low-to-high (the lowest redox potential was measured at 2cm deep, the heighest at 10 cm), HTL: high-to-low, UNO: unordered.

Distance	${\rm Orderedness}$	
220	LTH	
250	UNO	
300	UNO	
350	$_{ m LTH}$	
400	$_{ m LTH}$	
500	$_{ m LTH}$	
1000	UNO	
1150	$_{ m LTH}$	
1400	$\mathrm{HTL}$	
1600	$\mathrm{HTL}$	
1800	$\mathrm{HTL}$	
1970	$\mathrm{HTL}$	
2250	$\mathrm{HTL}$	
2300	$\mathrm{HTL}$	
2350	$\mathrm{HTL}$	
2400	$\mathrm{HTL}$	

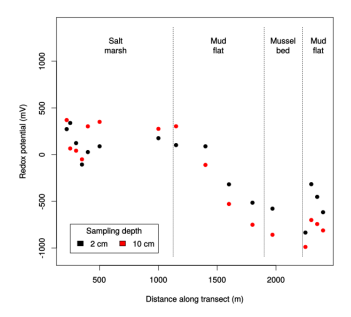


Figure 1: Redox potentials along the transect.

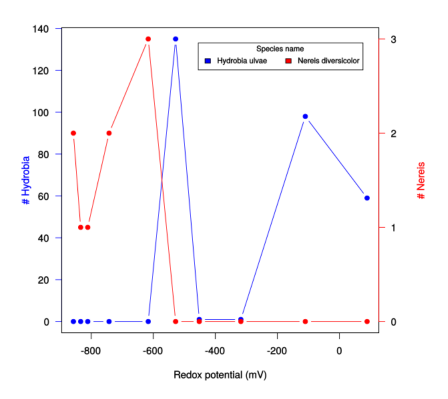


Figure 2: Number of individuals at the different redox potentials

```
# Manuscript, script to generate figures and tables for the Evolution & Ecology
Research course 2014
# Copyright (C) 2014 Richel Bilderbeek
  \# This program is free software: you can redistribute it and/or modify \# it under the terms of the GNU General Public License as published by \# the Free Software Foundation, either version 3 of the License, or \# (at your option) any later version.
  #
# This program is distributed in the hope that it will be useful,
# but WITHOUT ANY WARRANTY; without even the implied warranty of
# MERCHANTABLITY or FITNESS FOR A PARTICULAR PURPOSE. See the
# GNU General Public License for more details.
# You should have received a copy of the GNU General Public License
# along with this program If not, see <a href="http://www.gnu.org/licenses/">http://www.gnu.org/licenses/</a>>.
# File --- b.
  #
# File can be downloaded from https://github.com/richelbilderbeek/
EvoEcoResearchCourse2014
  ^{\pi} Research question: Can I predict where species are from a redox potential?
  # Experiment:

# * Couple shallow (0-5 \text{ cm}) benthos abundances with redox measurement of 2 cm

# * Couple deep (5-20 \text{ cm}) benthos abundances with redox measurement of 10 cm
  # Answer:
 rm(list = ls())
setwd("~/GitHubs/EvoEcoResearchCourse2014")
library(reshape2)
library(testit)
 # Create benthos data as such:
# dist_m depth_cm species_name
# 2350 2 Bathyporeia_spec.
# 2400 10 Nereis_diversicolor
# Removes species at unknown depths
CreateDataBenthos <- function()
        data_benthos <- read.table("benthos_species_diversity.csv",header=TRUE,sep="#Remove useless columns data_benthos <- data_benthos [c("dist_m","soil_layer","species_name")] #Only keep rows with 'D' or 'S' (for deep and shallow/top part of the core) data_benthos <- data_benthos | layer | = "t" & data_benthos | layer | = "t" | #Drop the unneeded levels | #Orop the unneeded levels | #Of data_benthos | layer <- droplevels (data_benthos | layer | = "t" | #Of data_benthod, change the column 'soil_layer' to 'depth_cm': # if soil_layer == "S" -> depth_cm = 2 | # if soil_layer == "D" -> depth_cm = 10 | names (data_benthos | layer == "D" -> depth_cm | layer 
          \mathtt{data\_benthos} \ <- \ \mathtt{read.table} ( \ \mathtt{"benthos\_species\_diversity.csv"}, \mathtt{header} \equiv \mathtt{TRUE}, \mathtt{sep} \equiv \mathtt{","})
  # Count the species occurring at two depths
  # species_name 2 10
# Arenicola_marina 12 1
# Bathyporeia_spec. 2 0
GetSpeciesCountAtDepths <- function()
        species_to_depth <- dcast(
  CreateDataBenthos(),
  species_name ~ depth_cm,
  value.var = "species_name",
  fill = 0,</pre>
                 fun.aggregate=length # fun.aggregate may also be mean
          species_to_depth
# Obtain the species names occuring at least thrice in both of the two depths in the
           species\_to\_depth \ <- \ GetSpeciesCountAtDepths()
        species_to_depth <- GetSpeciesCountAtDepths()
selected_species_to_depth <- subset(species_to_depth,
    species_to_depth$"2" > 2 & species_to_depth$"10" > 2)
selected_species_to_depth <- droplevels(selected_species_to_depth)
# Collect all species' names
species_list <- levels(selected_species_to_depth$species_name)
selected_species <- selected_species_to_depth$species_name
selected_species <- droplevels(selected_species)
selected_species</pre>
```

```
}
 GetDataBenthosSelected <- function()
   data benthos <- CreateDataBenthos()
   # Prepare the redox data
# dist_m depth_cm replicate redox_calib
# 220 2 1 304.89200
# 220 2 2 2 2 20.8326
 # 220
                                                   302.63036
 "CreateDataRedox <- function()
   data_redox <- read.table("Redox.csv",header=TRUE,sep="\t")
data_redox <- subset(data_redox,depth_cm != 5)
data_redox
   data redox
{\tt CreateDataRedoxAll} \ < - \ \ {\tt function} \ (\,)
    data redox <- read.table("Redox.csv", header=TRUE, sep="\t")
   #Remove the replicate column data_redox <- data_redox [c("dist_m", "depth_cm", "redox_calib")]
# Take the average redox values, so that every distance and depth has a single redox value
    data_redox <- aggregate(data_redox, list(data_redox$depth_cm,data_redox$dist_m),
   \begin{array}{lll} & & \\ & \text{data\_redox} < - & \text{data\_redox} \left[ \text{ c} \left( \text{"dist\_m","depth\_cm","redox\_calib"} \right) & \\ & \text{data\_redox} \end{array} \right] \\ & & \text{data\_redox} \end{array}
\label{eq:GetRedoxPerDistance} G\,et\,R\,ed\,o\,x\,P\,er\,D\,i\,st\,a\,n\,c\,e\ <-\ f\,u\,n\,c\,t\,i\,o\,n\,\left(\,\right)
   data_redox <- read.table("Redox.csv",header=TRUE,sep="\t")
#Remove the replicate column
data_redox <- data_redox[c("dist_m","depth_cm","redox_calib")]
# Take the average redox values, so that every distance and depth has a single
    redox value
data_redox <- aggregate(data_redox,list(data_redox$depth_cm,data_redox$dist_m),</pre>
            mean)
   mean)
data_redox <- data_redox[c("dist_m","depth_cm","redox_calib") ]
data_redox
# Get the 32 redox values used in this study
# [1] -987.68238 -857.44961 -834.07933 -811.22734 -750.16306 -742.48291 -699.70022
-616.30224 -577.85436 -528.05116
# [31] 350.83156 370.14974
GetRedoxValues <- function()
   redox_values <- subset(CreateDataRedox(), select="redox_calib")
redox_values <- unique(redox_values)
redox_values <- sort(redox_values$redox_calib)
redox_values</pre>
\# Obtain the distances of all redox sites \# [1] 220 250 300 350 400 500 1000 1150 1400 1600 1800 1970 2250 2300 2350
         2400
 Get Distances <- function ()
   \begin{array}{ll} distances <- & subset (\,C\,reateD\,ataRed\,ox\,(\,)\,\,,\,selec\,t="dist\_m\,"\,) \\ distances <- & unique(\,distances) \\ distances <- & sort\,(\,distances\$dist\_m\,) \\ \end{array}
    distances
# Get all species per redox potential

# redox_calib species_name

# -78.26751 Hydrobia_ulvae

# -78.26751 Hydrobia_ulvae

GetDataCombined <- function()
   data_combined <- merge(CreateDataRedox(),GetDataBenthosSelected(),by=c("dist_m"," depth_cm"),all=FALSE) data_combined$species_name <- droplevels(data_combined$species_name)
```

```
data_combined <- data_combined[ c("redox_calib","species_name") ] data_combined
# Get the redox potentials of Hydrobia
# [1] -78.26751 -78.26751 -78.26751 -7
-78.26751 -78.26751 -78.26751
                                                                      -78.26751 -78.26751 -78.26751 -78.26751
 GetRedoxesHydrobia <- function()
   redoxes_hydrobia <- subset(GetDataCombined(), species_name == "Hydrobia_ulvae") redoxes_hydrobia <- subset(redoxes_hydrobia, select=c("redox_calib")) redoxes_hydrobia
 GetRedoxesNereis <- function()
   redoxes_nereis <- subset (GetDataCombined () ,species_name == "Nereis_diversicolor") redoxes_nereis <- subset (redoxes_nereis ,select=c("redox_calib"))
    redoxes nereis
{\tt TallySpeciesPerRedox} \;\; < - \;\; {\tt function} \; \left( \; \right)
   redox_to_species <- dcast(
GetDataCombined(),
redox_calib ~ species_name,
value.var = "species_name"
f::1 - 0
                                  species_name,
        fill = 0.
        fun.aggregate=length # fun.aggregate may also be mean
   )
redox_calib <- redox_to_species$redox_calib
# Note that not all species are present anymore that often. This is due to that
not all distances are redoxed
redox_to_species
 {\tt Tally Selected Species Per Redox} \ < - \ function \ ( \, )
   redox_to_species <- TallySpeciesPerRedox()
redox_to_species <- redox_to_species[, colSums(redox_to_species) > 6]
redox_to_species <- cbind(TallySpeciesPerRedox(),redox_to_species)
redox_to_species <- redox_to_species[, c("redox_calib","Hydrobia_ulvae","
Nereis_diversicolor")]
redox_to_species
}
\# Can the redox potentials at a certain distane be assumed to be linear at a certain depth?
 CalcOrderednessPerDistance <- function()
    orderedness\_per\_distance <- \ data.frame(dist\_m = numeric(), order = factor()) \\ for \ (i \ in \ GetDistances())
    {
       \begin{array}{lll} data <- \ subset (CreateDataRedoxAll(), dist\_m == i) \\ low <- \ subset (data, depth\_cm == 2) \$redox\_calib \\ mid <- \ subset (data, depth\_cm == 5) \$redox\_calib \\ high <- \ subset (data, depth\_cm == 10) \$redox\_calib \\ if (low < mid && mid < high) \\ \end{array}
       if (low < mid && mid < nigh)
{
   orderedness_per_distance <- rbind(orderedness_per_distance,data.frame(dist_m =
        i, order = "LTH"))
} else if(low > mid && mid > high)
{

       .
deness_per_distance <- rbind(orderedness_per_distance,data.frame(dist_m = i, order = "UNO"))
       }
    orderedness_per_distance
# Generate figure for species abundances for the range of redox potentials # in two vertically aligned plots
CreateFigureSpeciesAbundancesSeperate <- function()
    svg\left(\,file\,n\,a\,m\,e\,=\,\,"\,F\,i\,g\,u\,r\,e\,\_\,s\,p\,e\,c\,i\,e\,s\,\_\,a\,b\,u\,n\,d\,a\,n\,c\,e\,s\,\_\,s\,e\,p\,e\,r\,a\,t\,e\,\,.\,\,sv\,g\,\,"\,\right)
    par(mfrow=c(2,1))
    plot (
        Hydrobia_ulvae ~ redox_calib ,
data = TallySpeciesPerRedox() ,
t = "b".
                        ulvae ~ redox
       t = "b",
pch = 19,
col = "black"
       #main = "Hydrobia ulvae abundance",
xlab = "Redox potential (mV)",
ylab = "# Hydrobia ulvae"
    plot (
        Nereis_diversicolor ~ redox_calib,
```

```
\begin{array}{lll} \mathtt{data} &=& \mathtt{TallySpeciesPerRedox} \; (\;) \;\;, \\ \mathsf{t} &=& \mathtt{"} \; \mathsf{b} \; \mathtt{"} \;, \end{array}
        t = "b",
pch = 19,
col = "black"
       #main = "Nereis diversicolor abundance",
xlab = "Redox potential (mV)",
ylab = "# Nereis Diversicolor",
ylim = c(0,4)
    par (mfrow=c(1,1))
dev.off()
# Generate figure for species abundances for the range of redox potentials # in the same plot
CreateFigureSpeciesAbundances <- function()
   svg(filename="Figure_species_abundances.svg")
par(mar = c(5, 4, 4, 4) + 0.3)  # Leave space for z axis
par(new = FALSE)
plot(Hydrobia_ulvae ~ redox_calib, data = TallySpeciesPerRedox(), pch=19, axes=
    FALSE, xlab="Redox potential (mV)", ylab="",
    type="b",col="blue"
    #main="Species abundandances\nfor different redox potentials"
}
    ) axis (1, col="black", las=1) #'las=1' align labels horizontally axis (2, col="blue", las=1) # las=1' align labels horizontally mtext("# Hydrobia", side=2, line=2.5, col="blue")
     par(new = TRUE) # Prevents R from clearing the area
        pch = 19
     axis (
         side = 4
         at=seq (0,3,1), #Otherwise, 0.5 would be shown as a tick mark col="red",
        las=1 #Align labels horizontally
    mtext("# Nereis", side=4, line=3,col="red")
legend("topright",
       egend ("topright", inset = 0.05, title = "Species name", c("Hydrobia ulvae", "Nereis diversicolor"), horiz=TRUE, fill=c("blue", "red"), cex = 0.75
    dev.off()
\# Redox potentials along the transect for the two depths CreateFigureRedoxPerDistance <- function()
   ) segments (1125,y_min,1125,y_max,col="black",lty=3) segments (1900,y_min,1900,y_max,col="black",lty=3) segments (2225,y_min,1900,y_max,col="black",lty=3) segments (2225,y_min,2225,y_max,col="black",lty=3) text ((1125 + 150) / 2,y_text,"Salt\nmarsh") text ((1900 + 1125) / 2,y_text,"Muk\nflat") text ((2225 + 1900) / 2,y_text,"Muk\nflat") text ((225 + 1900) / 2,y_text,"Mussel\nbed") text (-50 + ((2600 + 2225) / 2),y_text,"Musl\nhat{muk\nhat{n}lat"}) legend ("bottomleft", inset=0.05,title = "Sampling depth", c("2 cm", "10 cm"),horiz=TRUE, fill=c(1,2,3), cex = 1.0
   rm(dist_to_redox)
rm(y_min)
rm(y_max)
   rm(y_text)
dev.off()
assert ("CreateDataBenthos: 863 individuals were scored at known depths", length (
```

```
CreateDataBenthos() % dist_m) == 863)
assert("GetSpeciesCountAtbepths: All 863 individuals must be seperated correctly at their depths", length(CreateDataBenthos() % dist_m) == sum(GetSpeciesCountAtDepths () 8 "2") + sum(GetSpeciesCountAtDepths () 8 "0")
assert("GetSpeciesCountAtDepths: 20 species were scored at all depths", length() assert("GetSpeciesCountAtDepths: 20 species were scored at all depths", length() assert("GetSelectedSpecies: 8 species were found at both depths, at each depth occurring at lenst thrice", length(GetSelectedSpecies()) == 8) assert("GetDataBenthosSelecte: 740 individuals of the 8 selected species were scored at known depths", length(GetDataBenthosSelected() % species_name) == 740) assert("GetDataBenthosSelected: 740 individuals of the 8 selected species were scored at known depths", length(GetDataBenthosSelected() % species_name) == 740) assert("GetRedoxValues: 32 redox potentials are investigated", length(GetRedoxValues) assert("GetRedoxOstules: 32 redox potentials are investigated", length(GetRedoxValues) () == 32) assert("GetRedoxostombined: needs to be 1176 Hydrobia", length(GetDataCombined), assert("GetDataCombined: needs to be 1176 Hydrobia", length(GetDataCombined), assert("GetRedoxostyldrobia: must be 20 selected species_name) == 9) assert("GetRedoxostyldrobia: must be 204 values", length(GetRedoxostyldrobia: must be 205 clumns (redoxes and each species its frequency", length(TallySpeciesPerRedox() == 9) assert("TallySelectedSpeciesPerRedox() == 3) assert("TallySelectedSpeciesPerRedox() == 3) assert("TallySelectedSpeciesPerRedox() == 8) assert("TallySelectedSpeciesPe
```