

# Cellana

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Today

## Libraries

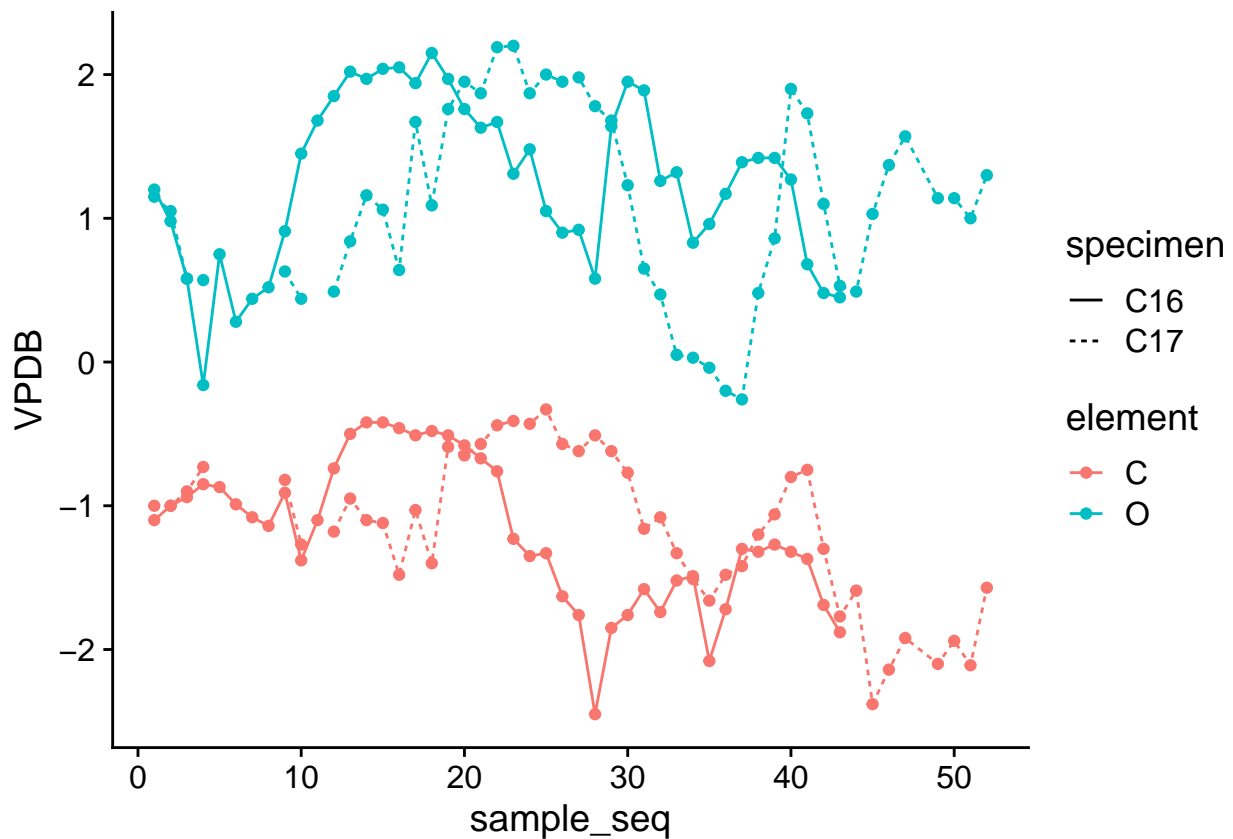
## Raw Data

## Plots

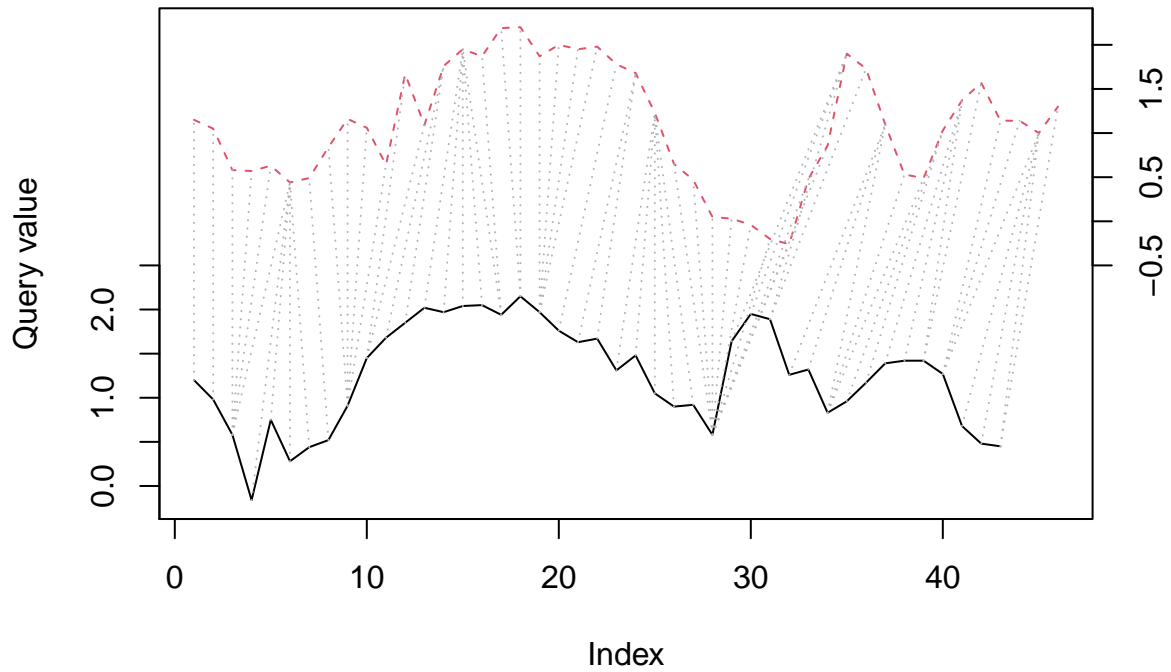
### Basic data

```
## Warning: Removed 16 row(s) containing missing values (geom_path).
```

```
## Warning: Removed 26 rows containing missing values (geom_point).
```



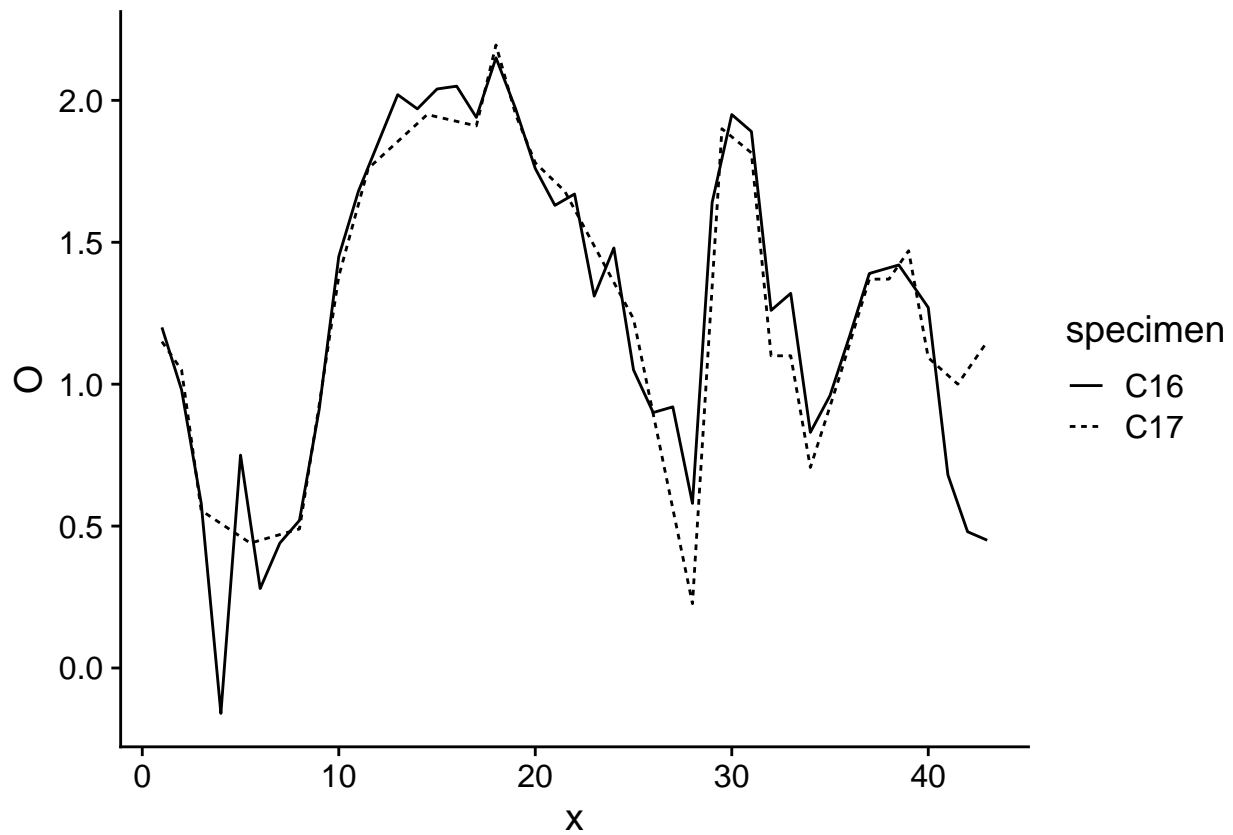
## Dynamic Time Warping



Add new x-values

```
## 'summarise()' has grouped output by 'x'. You can override using the '.groups' argument.
```

```
## 'summarise()' has grouped output by 'specimen'. You can override using the '.groups' argument.
```



Adding d13C values to check alignment

## 'summarise()' has grouped output by 'x'. You can override using the '.groups' argument.

## 'summarise()' has grouped output by 'specimen'. You can override using the '.groups' argument.

