

Selecting lines for HiLo F1s

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Start with SeeD GWAS GBS lines

I want all lines we select to be here to be able to take advantage of potential genotype info

Load the SeeDs data and match up with the CIMMYT data to extract CIMMYT availability and IDs

Load CIMMYT data for availability:

CIMMYT data and name conversion table, match up, and add bank numbers to CIMMYT_data table

Add CIMMYT info to SeeD:

Load GRIN data for Lat/Long/Elevation of some lines

Load GRIN data

Identify lines with multiple CIMMYTMA IDs. Exclude these lines. Numbers of lines with 0-2 CIMMYTMA IDs:

```
##      0      1      2
## 36021 9878    28
```

Identify lines that share the same CIMMYTMA ID. Exclude these lines.

```
## [1] 121
```

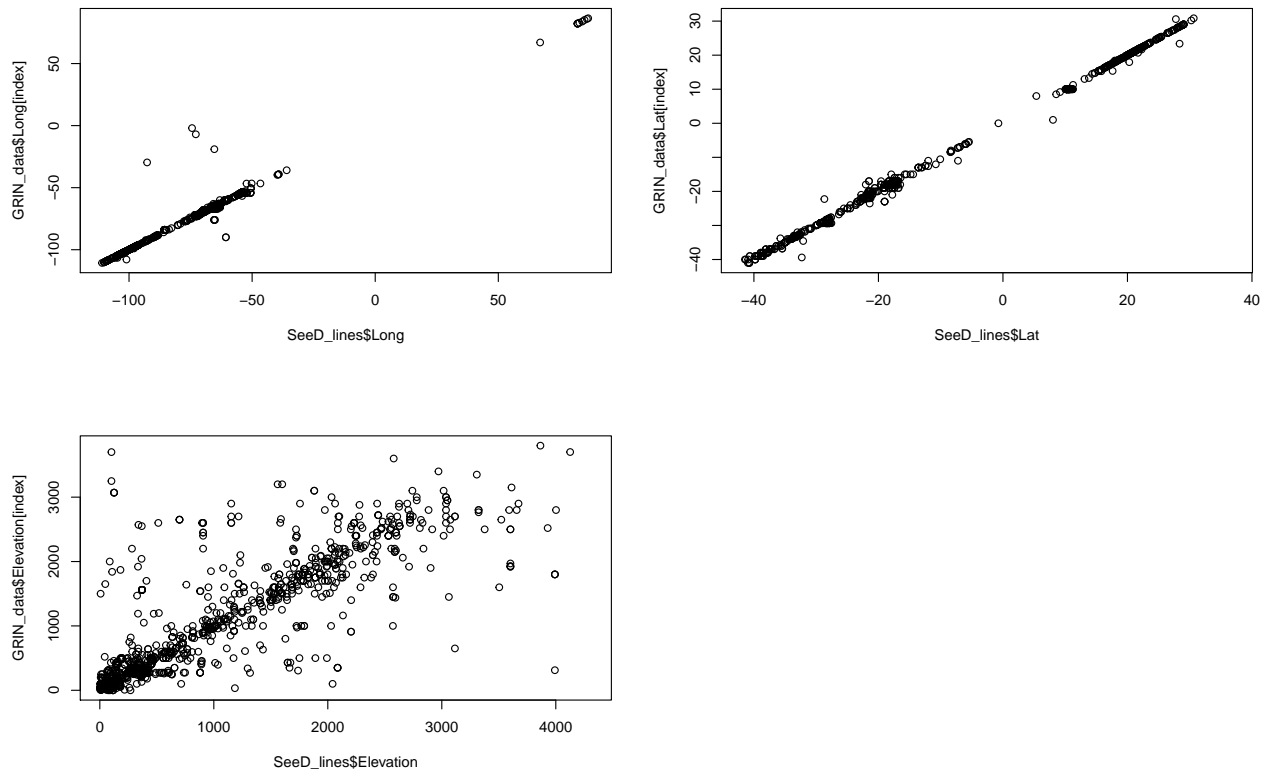
Match up GRIN and SeeD and add missing Elevation.

When info differs for the same line, use SeeD for Lat/Long, and GRIN for elevation

```
## [1] 0
```

```
## [1] 0
```

```
## [1] 44
```



Drop lines without coordinates

Load Current F1s

Make all available lines database

```
## [1] 16120      7
```

Which lines have GBS?

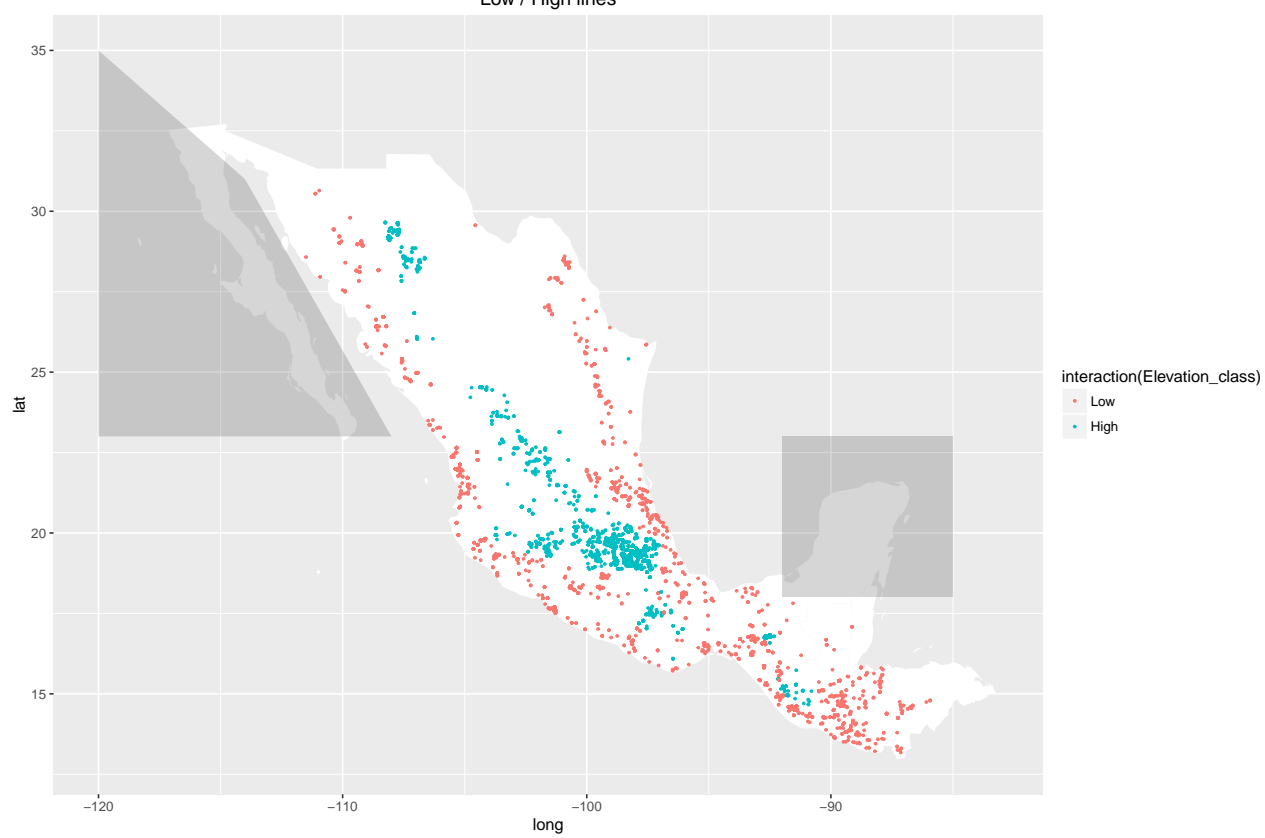
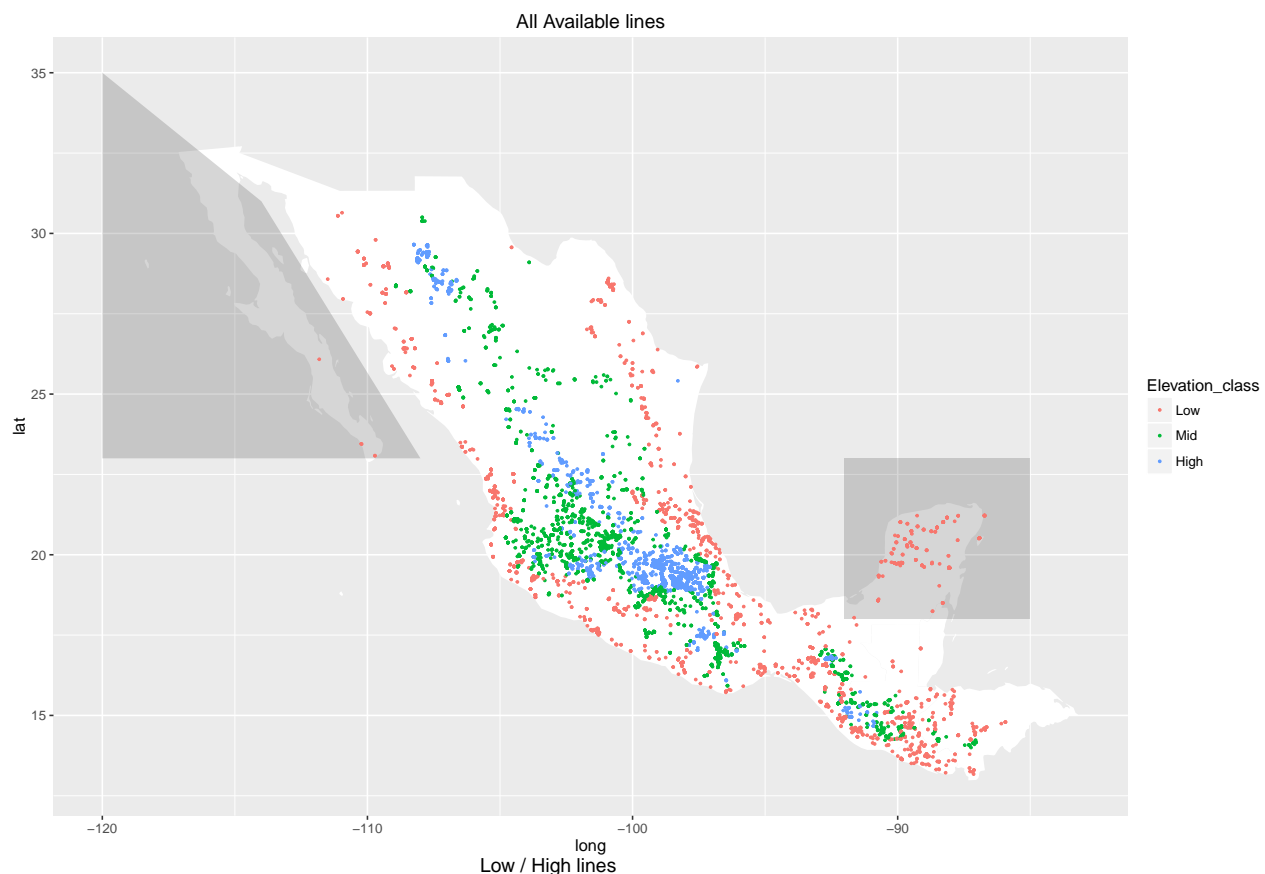
Divide lines by elevation:

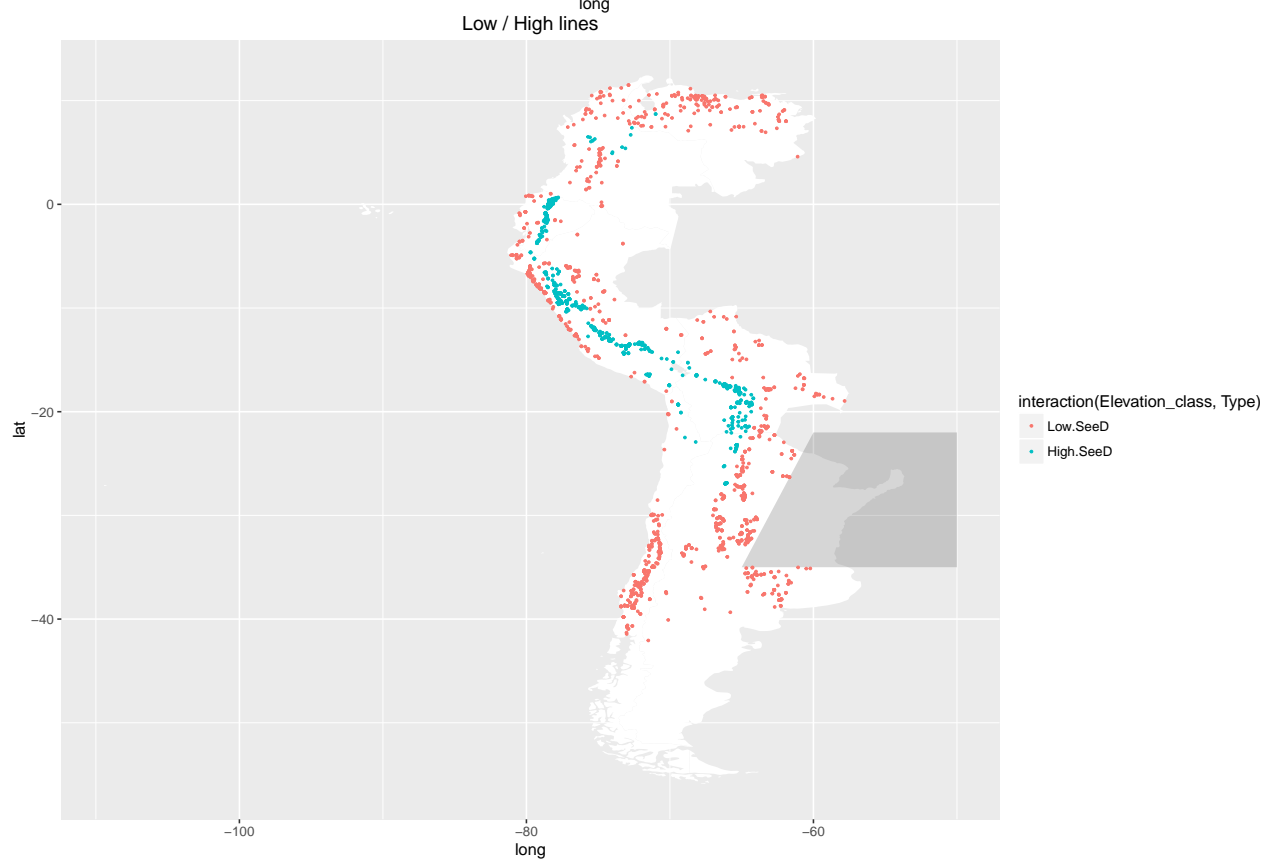
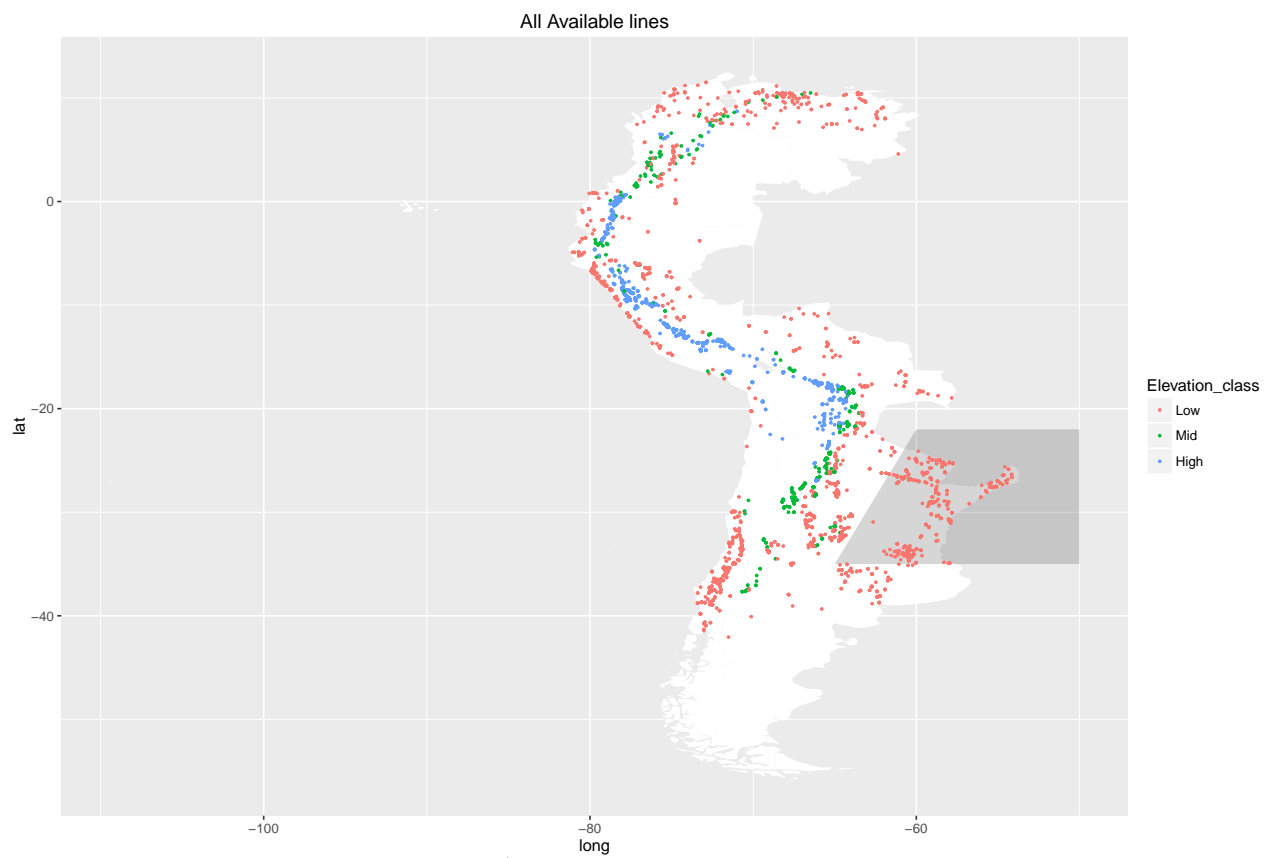
Subset lines by regions (Central America vs South America). Plot

Selecting candidate lines

Selection based on:

1. Available in CIMMYT
2. Either < 1000m or > 2000m
3. In SeeD database (although elevation info may have come from GRIN)





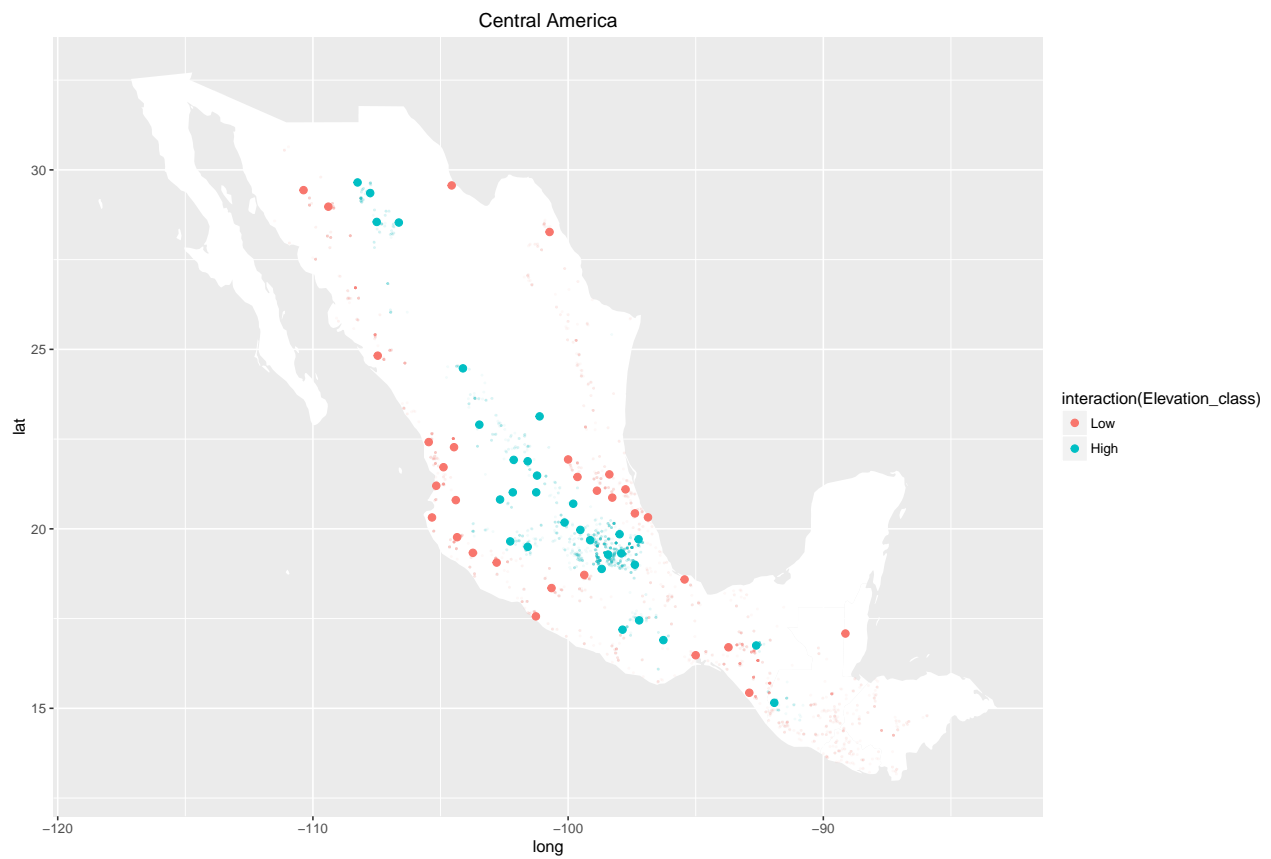
Selection algorithm 1

In this algorithm, I randomly select a number of high elevation lines, with the restriction that they cannot be too close together. Then for each high line, I pick a low line in the same latitude band, also not too close to any other line

Select Mex lines

```
##          SeeD Current_F1
##          56           4
```

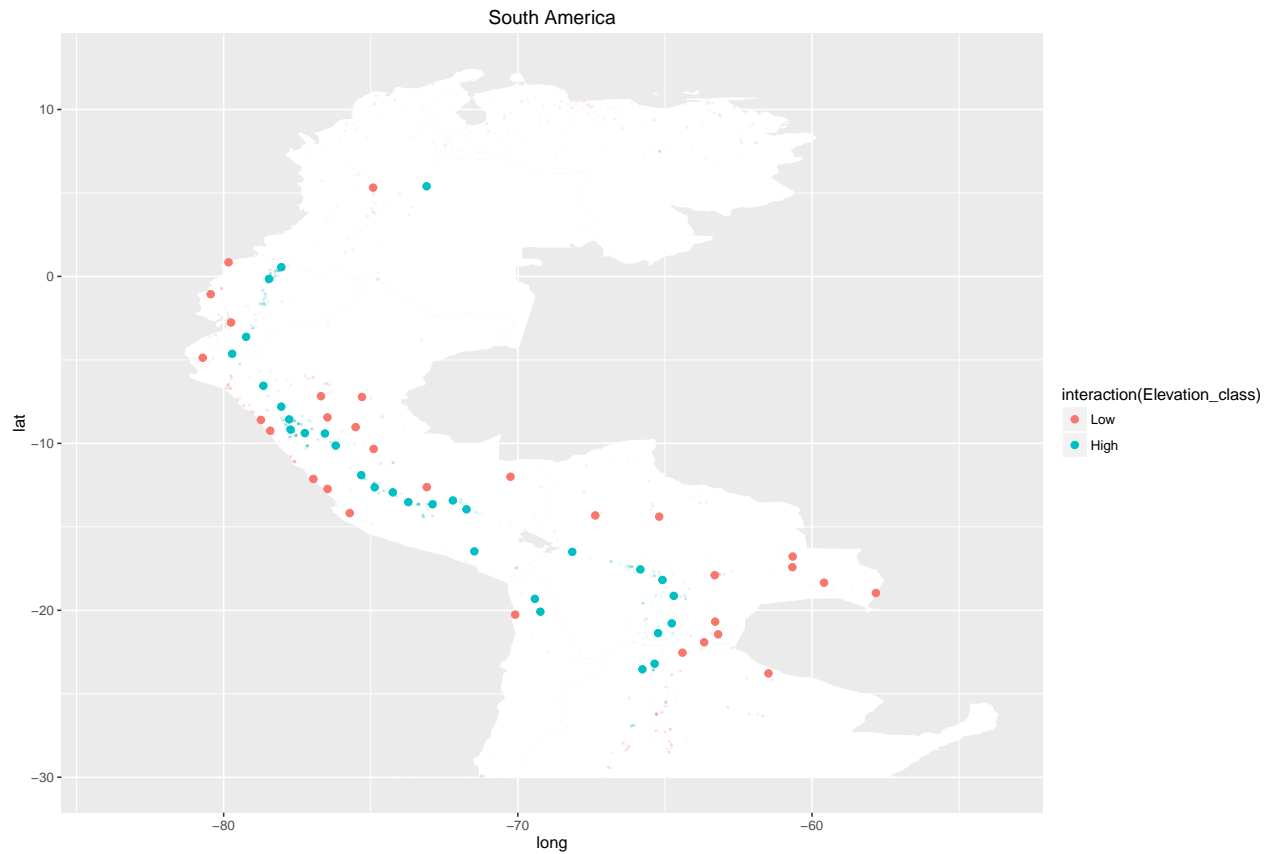
```
## Low  Mid High
##  30   0  30
```



Select SA lines

```
##          SeeD Current_F1
##          60           0
```

```
## Low  Mid High
##  30   0  30
```



Save lines in a DF

```
##          ID          Pair          Long          Lat
## CIMMYTMA-000449: 1 Mex_1 : 2 Min. : -110.37 Min. : -23.780
## CIMMYTMA-000547: 1 Mex_10 : 2 1st Qu.: -100.67 1st Qu.: -12.783
## CIMMYTMA-000554: 1 Mex_11 : 2 Median : -84.92 Median : 10.276
## CIMMYTMA-000645: 1 Mex_12 : 2 Mean : -86.24 Mean : 4.357
## CIMMYTMA-000686: 1 Mex_13 : 2 3rd Qu.: -73.57 3rd Qu.: 20.498
## CIMMYTMA-000861: 1 Mex_14 : 2 Max. : -57.83 Max. : 29.650
## (Other) :114 (Other):108
## Elevation Race GBS Elevation_class
## Min. : 2.0 :45 Mode :logical Low :60
## 1st Qu.: 303.8 TUXPEN :10 FALSE:107 Mid : 0
## Median :1505.5 CELAYA : 6 TRUE :13 High:60
## Mean :1443.4 CUBAAM : 6 NA's :0
## 3rd Qu.:2404.5 CONICO : 5
## Max. :3929.0 CONNOR : 5
## (Other):43
## id general_identifier bank_number
## Min. : 189 Min. : 5474 CIMMYTMA-BANK-000449: 1
## 1st Qu.:13690 1st Qu.:248942 CIMMYTMA-BANK-000547: 1
## Median :18102 Median :253356 CIMMYTMA-BANK-000554: 1
## Mean :17205 Mean :224272 CIMMYTMA-BANK-000645: 1
## 3rd Qu.:22148 3rd Qu.:257401 CIMMYTMA-BANK-000686: 1
## Max. :30844 Max. :267450 (Other) :111
## NA's :4 NA's :4 NA's : 4
```

```

##      collnumb      colldate
## AGUC67 : 1          :15
## ANC.526 : 1    1954-01-01: 8
## APUC.169: 1    1968-01-01: 5
## AREQ.164: 1    1943-01-01: 3
## ARGE521 : 1    1947-01-01: 3
## (Other) :111    (Other)   :82
## NA's    : 4      NA's     : 4
## Sample.ID.of.DNA.from.single.plants.used.in.GWAS      CIMMYT_ID
##              :103                                     CIMMYTMA 10248: 1
## SEEDGWAS1031: 1                                     CIMMYTMA 11854: 1
## SEEDGWAS156 : 1                                     CIMMYTMA 11857: 1
## SEEDGWAS2628: 1                                     CIMMYTMA 12085: 1
## SEEDGWAS2678: 1                                     CIMMYTMA 12231: 1
## (Other)      : 9                                     CIMMYTMA 12290: 1
## NA's         : 4                                     (Other)      :114
## Seed_Elevation  Elevation_flag
## Min.           : 2.0   Mode :logical
## 1st Qu.: 319.5   FALSE:116
## Median : 973.0   NA's :4
## Mean      :1428.4
## 3rd Qu.:2421.0
## Max.      :3929.0
## NA's      :4

```