

MAGEMin_C, the julia interface: introduction

Example 1 - predefined compositions

This is an example of how to use it for a predefined bulk rock composition:

```
julia> using MAGEMin_C
julia> db = "ig" # database: ig, igneous (Holland et al., 2018); mp, metapelite (White et al., 2018)
julia> data = Initialize_MAGEMin(db, verbose=true);
julia> test = 0 #KLB1
julia> data = use_predefined_bulk_rock(data, test);
julia> P = 8.0;
julia> T = 800.0;
julia> out = point_wise_minimization(P,T, data);
```

which gives

```
Status      :      0
Mass residual : +5.34576e-06
Rank         :      0
Point        :      1
Temperature  : +800.00000 [C]
Pressure     : +8.00000 [kbar]

SOL = [G: -797.749] (25 iterations, 39.62 ms)
GAM = [-979.481432, -1774.104523, -795.261024, -673.747244, -375.070247, -917.557241, -829.990582, -1000.000000]

Phase :      spn      cpx      opx      ol
Mode  :  0.02799  0.14166  0.24228  0.58807
```

MAGEMin_C, the julia interface: installation



https://github.com/ComputationalThermodynamics/MAGEMin_C.jl



Install MAGEMin_C

```
julia> ] # opens the package manager
pkg> add MAGEMin_C # MAGEMin_C
```

Load MAGEMin_C

```
julia> using MAGEMin_C # load MAGEMin_C
```

```
seph42@DESKTOP-2V82075:~$ julia
Documentation: https://docs.julialang.org
Type "?" for help, "]" for Pkg help.
Version 1.10.0 (2023-12-25)
Official https://julialang.org/ release

(@v1.10) pkg> add MAGEMin_C
Resolving package versions...
Updating ~/.julia/environments/v1.10/Project.toml
[e5d170eb] + MAGEMin_C v1.5.5
No Changes to ~/.julia/environments/v1.10/Manifest.toml

julia> using MAGEMin_C
Using libMAGEMin.dylib from MAGEMin_jll
```

MAGEMin_C, the julia interface: online documentation



https://github.com/ComputationalThermodynamics/MAGEMin_C.jl

- Numerous examples outside what will be presented during the short course are given in the README.md
- Simply access the MAGEMin_C.jl github link and scroll down to see them!

README GPL-3.0 license Security

MAGEMin_C.jl

Julia interface to the MAGEMin C package, which performs thermodynamic equilibrium calculations. See the [MAGEMin](#) page for more details on the package & how to use it.

Using the julia interface

First install julia. We recommend downloading the official binary from the [julia](#) webpage.

Next, install the `MAGEMin_C` package with:

```
julia> ]
pkg> add MAGEMin_C
```

You can check if it works on your system by running the build-in test suite:

```
pkg> test MAGEMin_C
```

By pushing `backspace` you return from the package manager to the main julia terminal. This will download a compiled version of the library as well as some wrapper functions to your system.

Next, you can do calculations with:

Example 1 - predefined compositions

This is an example of how to use it for a predefined bulk rock composition:

```
julia> using MAGEMin_C
julia> db = "ig" # database: ig, igneous (Holland et al., 2018); mp, metapelite (White et al.
julia> data = Initialize_MAGEMin(db, verbose=true);
julia> test = 0
julia> data = use_predefined_bulk_rock(data, test);
julia> P = 8.0;
julia> T = 800.0;
julia> out = point_wise_minimization(P,T, data);
```

```
julia> data = Initialize_MAGEMin(db, verbose=true);
julia> test = 0
julia> data = use_predefined_bulk_rock(data, test);
julia> P = 8.0;
julia> T = 800.0;
julia> out = point_wise_minimization(P,T, data);
```

which gives

```
Status      : 0
Mass residual : +5.34570e-06
Rank        : 0
Point       : 1
Temperature  : +800.00000 [C]
Pressure     : +8.00000 [kbar]

SOL = [G: -797.749] (25 iterations, 39.62 ms)
GAM = [-979.481432, -1774.104523, -795.201024, -673.747244, -375.079247, -917.557241, -829.990582, -
```

Example 2 - custom composition

And here a case in which you specify your own bulk rock composition.

```
julia> using MAGEMin_C
julia> data = Initialize_MAGEMin("ig", verbose=false);
julia> P,T = 10.0, 1100.0
julia> Xoxides = ["SiO2", "Al2O3", "CaO", "MgO", "FeO", "Fe2O3", "K2O", "Na2O", "TiO2", "Cr2O3"]
julia> X = [48.43; 15.19; 11.57; 18.13; 6.65; 1.64; 0.59; 1.87; 0.68; 0.0; 3.8];
julia> sys_in = "wt"
julia> out = single_point_minimization(P, T, data, X=X, Xoxides=Xoxides, sys_in=sys_in)
```

which gives:

```
Pressure      : 10.0 [kbar]
Temperature    : 1100.0 [Celsius]
Stable phase | Fraction (mol fraction)
```

```
Stable phase | Fraction (mol fraction)
liq          | 0.75133
cpx          | 0.20987
opx          | 0.03877
Stable phase | Fraction (wt fraction)
liq          | 0.73091
cpx          | 0.22895
opx          | 0.04096
Gibbs free energy : -916.874646 (45 iterations; 86.53 ms)
Oxygen fugacity   : 2.0509883251350577e-8
```

After the calculation is finished, the structure `out` holds all the information about the stable assemblage, including seismic velocities, melt content, melt chemistry, densities etc. You can show a full overview of that with

```
julia> print_info(out)
```

If you are interested in the density or seismic velocity at the point, access it with

```
julia> out.rho
2755.2995530913095
julia> out.Vp
3.945646731595539
```

Once you are done with all calculations, release the memory with

```
julia> Finalize_MAGEMin(data)
```

Example 3 - Removing solution phase(s) from consideration

To suppress solution phases from the calculation, define a remove list `rm_list` using the `remove_phases()` function. In the latter, provide a vector of the solution phase(s) you want to remove and the database acronym as a second argument. Then pass the created `rm_list` to the `single_point_minimization()` function.

```
julia> using MAGEMin_C
julia> data = Initialize_MAGEMin("mp", verbose=1, solver=0);
julia> rm_list = remove_phases(["liq", "sp", "mp"]);
```

Moreover, the testing framework can provide more knowledge:
https://github.com/ComputationalThermodynamics/MAGEMin_C.jl/blob/main/test/tests.jl

MAGEMin_C, the julia interface: let's get started!

- MAGEMin_C allow you to write phase equilibrium **Julia** scripts fitted to your needs
- Numerous resources to learn Julia can be found online e.g., <https://julialang.org/learning/tutorials/>
- Overall, **Julia** programming style is quite like **Matlab** and being a Matlab user will help to write in Julia
- Here we are going to explore the basic functionality **MAGEMin_C** and we will go through one advanced example on how to compute iso-entropic paths (fixed entropy during decompression, applied to MORB genesis)
- Note that if you struggle, the scripts presented are available in the resources, in the GG2024 folder

MAGEMin_C, simple example

- The following example performs an equilibrium given the igneous database “ig” and the predefined test for KLB-1 peridotite

Note that: “ig”, igneous; “mp”, metapelite; “mb”, metabasite; “um”, ultramafic

```
using MAGEMin_C          # load MAGEMin

# Initialize database
data      = Initialize_MAGEMin("ig", verbose=true);
test      = 0             #KLB1
data      = use_predefined_bulk_rock(data, test);

# Call optimization routine for given P & T & bulk_rock
P          = 8.0
T          = 800.0
out        = single_point_minimization(P,T, data);
Finalize_MAGEMin(data)
```

THIS SNIPPET CAN BE COPIED

- Initialize MAGEMin with “ig” database
 - Selects test 0, i.e., KLB-1 peridotite
 - Set pressure and temperature
 - Performs the calculation
 - Deallocate memory
- “out” is a structure that stores all the information about the equilibrium. We will see how to create a vector of structure to stores multiple equilibrium point information

MAGEMin_C, first example

- Executing the previous set of commands gives a summary of the computed equilibrium

```
julia> out = single_point_minimization(P,T, data);
Status      :      0
Mass residual : +2.44463e-14
Rank        :      0
Point       :      1
Temperature  : +800.00000      [C]
Pressure     : +8.00000      [kbar]

SOL = [G: -797.749] (106 iterations, 87.90 ms)
GAM = [-979.481429,-1774.103896,-795.262450,-673.746205,-375.079335,-917.592609,-829.966523,-1023.697396,-256.999589,-1308.291347]

Phase :      spn      opx      cpx      ol
Mode  : 0.02799 0.24227 0.14166 0.58807
```

Note that “GAM” stands for gamma and the array is the chemical potential of the oxides of the system

- To access the information store in “out” within the terminal simply type “out.” then hit the “tab” key



```
julia> out.
G_system      Gamma      MAGEMin_ver  M_sys      PP_vec      P_kbar      SS_vec      T_C      V      Vp      Vp_S      Vs      Vs_S      X
aAl2O3      aFeO      aH2O      aMgO      aSiO2      aTiO2      alpha      bulk      bulkMod      bulkModulus_M      bulkModulus_S      bulk_F      bulk_F_wt      bulk_M
bulk_M_wt      bulk_S      bulk_S_wt      bulk_res_norm      bulk_wt      cp      dQFM      dataset      enthalpy      entropy      f02      frac_F      frac_F_wt      frac_M
frac_M_wt      frac_S      frac_S_wt      iter      mSS_vec      n_PP      n_SS      n_mSS      oxides      ph      ph_frac      ph_frac_vol      ph_frac_wt      ph_id
ph_type      rho      rho_F      rho_M      rho_S      s_cp      shearMod      shearModulus_S      status      time_ms
```

MAGEMin_C, output structure

- “out” structure is of type `MAGEMin_C.gmin_struct{Float64, Int64}`

- Some details about information stored in “out”

| | |
|--------------------------|---|
| <code>out.dataset</code> | → Dataset used for the calculation |
| <code>out.oxides</code> | → List of oxides |
| <code>out.Gamma</code> | → Chemical potential of oxides |
| <code>out.bulk</code> | → Bulk rock composition in mol |
| <code>out.bulk_M</code> | → Bulk composition of melt if any |
| <code>out.bulk_S</code> | → Bulk composition of solid part if any |
| <code>out.ph</code> | → Stable phases names |
| <code>out.ph_frac</code> | → Stable phases fraction in mol |
| <code>out.frac_M</code> | → Melt fraction in mol |
| <code>out.frac_S</code> | → Solid fraction in mol |
| <code>...</code> | |

| | |
|-----------------------------|---------|
| <code>out.bulk_wt</code> | → in wt |
| <code>out.bulk_M_wt</code> | → in wt |
| <code>out.bulk_S_wt</code> | → in wt |
| <code>out.ph_frac_wt</code> | → in wt |
| <code>out.frac_M_wt</code> | → in wt |
| <code>out.frac_S_wt</code> | → in wt |

MAGEMin_C, output structure

- Phases information can be access as:

| | | |
|-------------------------|---|--------------------------------------|
| <code>out.n_SS</code> | → | Number of stable solution phases |
| <code>out.n_PP</code> | → | Number of stable pure phases |
| <code>out.SS_vec</code> | → | Vector{MAGEMin_C.LibMAGEMin.SS_data} |
| <code>out.PP_vec</code> | → | Vector{MAGEMin_C.LibMAGEMin.SS_data} |

- You can access the details of solution phase 1 as:

`out.SS_vec[1].`

then hit the “tab” key



```
julia> out.SS_vec[1].
Comp      Comp_wt      G      V      Vp
Vs         alpha      bulkMod  compVariables  compVariablesNames
cp         deltaG      emChemPot  emComp      emComp_wt
emFrac      emFrac_wt      emNames  enthalpy      entropy
f           rho      shearMod  siteFractions  siteFractionsNames
julia> out.SS_vec[1].
```

- Then you can access any content of the solution model as

| | | |
|------------------------------------|---|---|
| <code>out.SS_vec[1].Comp</code> | → | Composition of the solution in mol fraction |
| <code>out.SS_vec[1].emNames</code> | → | End-member names |
| <code>out.SS_vec[1].emFrac</code> | → | End-member fraction |
| <code>...</code> | | |

MAGEMin_C, custom bulk-rock composition

- Equilibrium calculation with custom bulk-rock composition is straightforward

```
data      = Initialize_MAGEMin("ig", verbose=false);  
  
P,T       = 10.0, 1100.0  
Xoxides   = ["SiO2"; "Al2O3"; "CaO"; "MgO"; "FeO"; "Fe2O3"; "K2O"; "Na2O"; "TiO2"; "Cr2O3"; "H2O"];  
X         = [48.43; 15.19; 11.57; 10.13; 6.65; 1.64; 0.59; 1.87; 0.68; 0.0; 3.0];  
sys_in    = "wt"  
out       = single_point_minimization(P, T, data, X=X, Xoxides=Xoxides, sys_in=sys_in)  
Finalize_MAGEMin(data)
```

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- Xoxides → string array of oxides names
- X → bulk-rock composition, provide either FeO,O or FeO,Fe2O3
- sys_in → system unit: mol or wt

```
julia> out = single_point_minimization(P, T, data, X=X, Xoxides=Xoxides, sys_in=sys_in)  
Pressure      : 10.0      [kbar]  
Temperature   : 1100.0    [Celsius]  
Stable phase | Fraction (mol fraction)  
  opx         0.04245  
  liq         0.72699  
  cpx         0.23056  
Stable phase | Fraction (wt fraction)  
  opx         0.04473  
  liq         0.70443  
  cpx         0.25084  
Stable phase | Fraction (vol fraction)  
  opx         0.03743  
  liq         0.75082  
  cpx         0.21174  
Gibbs free energy : -916.874646 (45 iterations; 90.29 ms)  
Oxygen fugacity   : -7.688086637813585  
Delta QFM         : 1.2159866924221605
```

MAGEMin_C, parallel calculation

- When Julia is launched in parallel, one can perform equilibrium calculation of several points in parallel:

```
data = Initialize_MAGEMin("ig", verbose=false);
P = [10.0, 12.0]
T = [1100.0, 1000.0]
Xoxides = ["SiO2"; "Al2O3"; "CaO"; "MgO"; "FeO"; "Fe2O3"; "K2O"; "Na2O"; "TiO2"; "Cr2O3"; "H2O"];
X1 = [48.43; 15.19; 11.57; 10.13; 6.65; 1.64; 0.59; 1.87; 0.68; 0.0; 3.0];
X2 = [49.43; 14.19; 11.57; 10.13; 6.65; 1.64; 0.59; 1.87; 0.68; 0.0; 0.0];
X = [X1,X2]
sys_in = "wt"
out = multi_point_minimization(P, T, data, X=X, Xoxides=Xoxides, sys_in=sys_in)
Finalize_MAGEMin(data)
```

```
julia> out = multi_point_minimization(P, T, data, X=X, Xoxides=Xoxides, sys_in=sys_in)
2-element Vector{MAGEMin_C.gmin_struct{Float64, Int64}}:
  Pressure : 10.0 [kbar]
  Temperature : 1100.0 [Celsius]
    Stable phase | Fraction (mol fraction)
      opx 0.04245
      liq 0.72699
      cpx 0.23056
    Stable phase | Fraction (wt fraction)
      opx 0.04473
      liq 0.70443
      cpx 0.25084
    Stable phase | Fraction (vol fraction)
      opx 0.03743
      liq 0.75082
      cpx 0.21174
  Gibbs free energy : -916.874646 (45 iterations; 91.5 ms)
  Oxygen fugacity : -7.688086637813585
  Delta QFM : 1.2159866924221605
```

```
Pressure : 12.0 [kbar]
Temperature : 1000.0 [Celsius]
  Stable phase | Fraction (mol fraction)
    g 0.00767
    fsp 0.31479
    cpx 0.42243
    fsp 0.01294
    opx 0.23891
    ru 0.00326
  Stable phase | Fraction (wt fraction)
    g 0.00789
    fsp 0.30308
    cpx 0.43189
    fsp 0.01261
    opx 0.24054
    ru 0.00399
  Stable phase | Fraction (vol fraction)
    g 0.0065
    fsp 0.34987
    cpx 0.40516
    fsp 0.01522
    opx 0.22028
    ru 0.00298
  Gibbs free energy : -943.792662 (16 iterations; 36.2 ms)
  Oxygen fugacity : -8.176560770673227
  Delta QFM : 1.912657699586645
```

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MAGEMin_C, parallel calculation: reading output

- For multi point minimization the outputs are store as `Vector{MAGEMin_C.gmin_struct{Float64, Int64}}`
- To access individual point information simply do

```
julia> out[1].
G_system      Gamma      MAGEMin_ver  M_sys      PP_vec      P_kbar      SS_vec      T_C      V      Vp      Vp_S      Vs      Vs_S      X
aAl2O3        aFeO        aH2O        aMgO        aSiO2        aTiO2        alpha      bulk      bulkMod      bulkModulus_M  bulkModulus_S  bulk_F  bulk_F_wt  bulk_M
bulk_M_wt     bulk_S      bulk_S_wt   bulk_res_norm  bulk_wt     cp          dQFM      dataset  enthalpy  entropy      fO2      frac_F  frac_F_wt  frac_M
frac_M_wt     frac_S      frac_S_wt   iter        mSS_vec     n_PP        n_SS      n_mSS    oxides    ph           ph_frac  ph_frac_vol  ph_frac_wt  ph_id
ph_type      rho         rho_F      rho_M      rho_S      s_cp        shearMod   shearModulus_S  status    time_ms
```

```
julia> out[2].
G_system      Gamma      MAGEMin_ver  M_sys      PP_vec      P_kbar      SS_vec      T_C      V      Vp      Vp_S      Vs      Vs_S      X
aAl2O3        aFeO        aH2O        aMgO        aSiO2        aTiO2        alpha      bulk      bulkMod      bulkModulus_M  bulkModulus_S  bulk_F  bulk_F_wt  bulk_M
bulk_M_wt     bulk_S      bulk_S_wt   bulk_res_norm  bulk_wt     cp          dQFM      dataset  enthalpy  entropy      fO2      frac_F  frac_F_wt  frac_M
frac_M_wt     frac_S      frac_S_wt   iter        mSS_vec     n_PP        n_SS      n_mSS    oxides    ph           ph_frac  ph_frac_vol  ph_frac_wt  ph_id
ph_type      rho         rho_F      rho_M      rho_S      s_cp        shearMod   shearModulus_S  status    time_ms
```

- The number of points can be retrieved using

`length(out)`



```
julia> length(out)
2
```

- Information of each minimization can otherwise be accessed as described in slides 7-8

MAGEMin_C, some useful commands

- Bulk-rock composition can be converted to MAGEMin system unit (mol) with

```
bulk_in_ox = ["SiO2"; "Al2O3"; "CaO"; "MgO"; "FeO"; "Fe2O3"; "K2O"; "Na2O"; "TiO2"; "MnO"; "H2O"];  
bulk_in    = [69.64; 13.76; 1.77; 1.73; 4.32; 0.4; 2.61; 2.41; 0.80; 0.07; 0.0];  
bulk_rock,ox = convertBulk4MAGEMin(bulk_in,bulk_in_ox,"wt","mp");
```

Note that here, we convert for the “mp” (metapelite) database from wt to mol and transforming FeO,Fe2O3 to FeO,O

```
julia> bulk_rock  
11-element Vector{Float64}:  
 76.57038397179574  
  8.914984523583415  
  2.0849576977131403  
  2.835783318610597  
  4.30275071755529  
  1.8302970975627948  
  2.568605789798099  
  0.6615823604771729  
  0.16546809116073818  
  0.06518643174302832  
  0.0
```

```
julia> ox  
11-element Vector{String}:  
 "SiO2"  
 "Al2O3"  
 "CaO"  
 "MgO"  
 "FeO"  
 "K2O"  
 "Na2O"  
 "TiO2"  
 "O"  
 "MnO"  
 "H2O"
```

- Declaring an “array of output structure” to store results of several minimizations can be done with

```
n    = 10  
out = Vector{MAGEMin_C.gmin_struct{Float64, Int64}}(undef, n)
```

Note that n can be any size

MAGEMin_C, fractional crystallization: definition

- In the following example we are going to fractionate a basaltic composition using the igneous database. We are also going to use “Plots” package to visualize the results
- First declare the database, initialize MAGEMin and provide bulk-rock composition, oxide list and system unit

```
using MAGEMin_C
using Plots

dtb      = "ig"
data     = Initialize_MAGEMin(dtb);

Xoxides  = ["SiO2", "Al2O3", "CaO", "MgO", "FeO", "K2O", "Na2O", "TiO2", "O", "Cr2O3", "H2O"]
X        = [50.0, 8.7, 11.7, 12.14, 7.7, 0.2, 2.5, 1.0, 0.5, 0.01, 10.0]
sys_in   = "mol";
```

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| | | | |
|-------------------------------|---|---------|---|
| Pressure | → | P | = 5.0 |
| Number of fractionation steps | → | n_steps | = 64 |
| Starting temperature | → | Ts | = 1200.0 |
| Ending temperature | → | Te | = 600.0 |
| Define temperature range | → | T | = Array(range(Ts, stop=Te, length=n_steps)) |
| Declare output vector | → | out | = Vector{MAGEMin_C.gmin_struct{Float64, Int64}}(undef, n_steps) |

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MAGEMin_C, fractional crystallization: calculation

- In its simplest form, fractional crystallization uses the composition of the stable melt at a given temperature, as the starting bulk-rock composition at decreased temperature

```
for i in 1:n_steps
    out[i] = deepcopy(single_point_minimization(P, T[i], data, X=X, Xoxides=Xoxides, sys_in=sys_in))
    if "liq" in out[i].ph                                     # If liquid phase is
        X = deepcopy(out[i].bulk_M)
    end
end
```

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- **In the above code:**
 1. We loop through all defined steps (64 as defined in previous slide)
 2. Perform the equilibrium calculation given the temperature array $T[i]$ and the bulk composition X . Note that the result are stored in “out[i]”
 3. If “liq” is stable, we then update the bulk-rock composition X to be equal to the composition of the melt “out[i].bulk_M”
otherwise, the bulk is unchanged

MAGEMin_C, fractional crystallization: data extraction

- Now that the calculation is performed, we need to be able to transform the saved data in a useful way to be plotted. The code snippet below, gives you some hints on how it can be done

```
frac_M      = [out[i].frac_M for i in 1:n_steps]
frac_M_tot  = accumulate(*, frac_M)

SiO2_id     = findfirst(out[1].oxides .== "SiO2")
dry_id      = findall(out[1].oxides .!= "H2O")

SiO2_M_dry  = [ (out[i].bulk_M[SiO2_id] / sum(out[i].bulk_M[dry_id])*100.0) for i in 1:n_steps]

rho_M       = [ (out[i].rho_M) for i in 1:n_steps]
rho_M[rho_M .== 0.0] .= NaN;
```

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- In the code above, we extract and clean several information**
 - The fraction of stable melt in mol
 - The accumulated stable melt fraction with respect to the starting mol quantity
 - We retrieve the array index of SiO₂ and the indices of all oxides excluding water, this for anhydrous normalization
 - We compute the SiO₂ content of the melt on an anhydrous basis
 - We retrieve the melt density and set to NaN the values equal to 0.0 (when melt is not stable)

MAGeMin_C, fractional crystallization: visualization

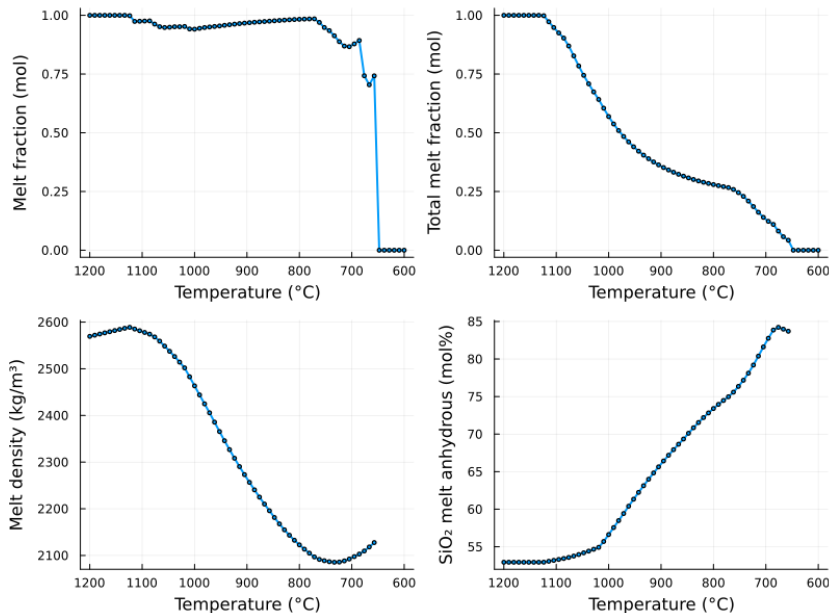
- Now that some data have been converted, we can visualize it. Here is provided one example of visualization using Plots, you are of course welcome to use any other package

```
p1 = plot(T,frac_M, xflip=true, xlabel="Temperature (°C)", marker = :circle, markersize = 2, lw=2, ylabel="Melt fraction (mol)", legend=false)
p2 = plot(T,frac_M_tot, xflip=true, xlabel="Temperature (°C)", marker = :circle, markersize = 2, lw=2, ylabel="Total melt fraction (mol)", legend=false)
p3 = plot(T,rho_M, xflip=true, xlabel="Temperature (°C)", marker = :circle, markersize = 2, lw=2, ylabel="Melt density (kg/m³)", legend=false)
p4 = plot(T,SiO2_M_dry, xflip=true, xlabel="Temperature (°C)", marker = :circle, markersize = 2, lw=2, ylabel="SiO₂ melt anhydrous (mol%)", legend=false)

fig = plot(p1, p2, p3, p4, layout=(2, 2), size=(800, 600))
savefig(fig,"frac_crystallization.png")
```

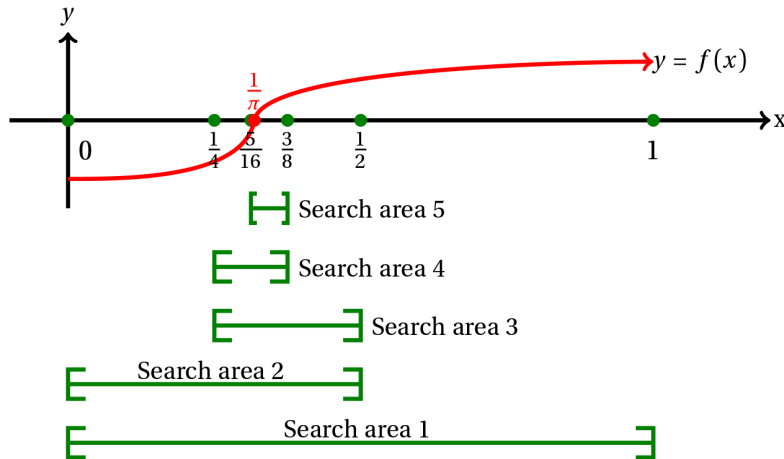
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- Which yields



MAGEMin_C, advanced example: isentropic path

- Isentropic paths (Adiabatic decompression) can be generated directly in MAGEMinApp, but here we are going to see how this can be achieved using MAGEMin_C
- The objective is to provide a starting PT condition, compute the entropy of the system, then gradually decrease pressure while keeping entropy constant and computing the corresponding temperature
- Because the temperature at decreased pressure that keeps entropy constant is unknown, we need a strategy to constrain it. Here we are going to use the bisection method.
- The idea is to reformulate the problem as a root finding problem and use the bisection method as described below:



- $f(x) = S_{ref} - S_x$
where x is temperature
 S_{ref} is entropy of reference
 S_x is the entropy at tentative temperature x

MAGEMin_C, isentropic path: definition

ProgressMeter to display a progress bar

Igneous database

Initialize MAGEMin while deactivating verbose

Select predefined KLB-1 test

Load predefined test

Mantle potential temperature

Adiabatic gradient

Starting depth

Mantle density

Starting temperature

Starting pressure

Ending pressure

Number of steps

Maximum number of bisection iterations

Tolerance in K

Pressure range for the isentropic path

Output structure Vector

Temporary output structure

Compute reference point at Ts and Ps

Reference entropy

```
using MAGEMin_C
using Plots
using ProgressMeter

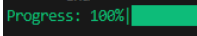
dtb      = "ig"
data     = Initialize_MAGEMin(dtb,verbose=-1)
test     = 0           # KLB-1
data     = use_predefined_bulk_rock(data, test)

MPT      = 1350.0
adiabat  = 0.55
Depth    = 100.0
rho_Mantle = 3300.0

Ts        = MPT + adiabat*Depth
Ps        = Depth*1e3*9.81*rho_Mantle/1e5/1e3
Pe        = 0.001
n_steps  = 32
n_max    = 32
tolerance = 0.1

P         = Array(range(Ps, stop=Pe, length=n_steps))
out       = Vector{MAGEMin_C.gmin_struct{Float64, Int64}}(undef, n_steps)
out_tmp   = MAGEMin_C.gmin_struct{Float64, Int64};
out[1]    = deepcopy( single_point_minimization(Ps,Ts, data));
Sref      = out[1].entropy
```

MAGEMin_C, isentropic path: calculation

- This code snippet uses the bisection method to find the temperature fitting the entropy of reference within given tolerance
- The command “@showprogress” place before the pressure loop allow to display the progress bar in the Julia terminal 
- **a** and **b** define the starting temperature bounds
n is the number of bisection
c is the bisected temperature
result is the misfit ($f(x) = S_{ref} - S_x$)
sign_c is the sign of the misfit a $T = c$
- If the bounds distance is $< \text{tolerance}$, then solution is accepted
else, the bounds are updated

```
@showprogress for j = 2:n_steps
    a      = out[j-1].T_C - 50.0
    b      = out[j-1].T_C
    n      = 1
    conv   = 0
    n      = 0
    sign_a = -1

    while n < n_max && conv == 0
        c      = (a+b)/2.0
        out_tmp = deepcopy( single_point_minimization(P[j],c, data));
        result = out_tmp.entropy - Sref
        sign_c  = sign(result)

        if abs(b-a) < tolerance
            conv = 1
        else
            if sign_c == sign_a
                a = c
                sign_a = sign_c
            else
                b = c
            end
        end
        n += 1
    end
    out[j] = deepcopy(out_tmp)
end

Finalize_MAGEMin(data)
```

MAGEMin_C, isentropic path: data extraction

- Similarly to the factional crystallization example, data output need to be reformatted.

```
S          = [out[i].entropy for i in 1:n_steps]
frac_M     = [out[i].frac_M for i in 1:n_steps]
frac_M[frac_M .== 0.0] .= NaN
T          = [out[i].T_C for i in 1:n_steps]
SiO2_id    = findfirst(out[1].oxides .== "SiO2")
dry_id     = findall(out[1].oxides .!= "H2O")
SiO2_M_dry = [ (out[i].bulk_M[SiO2_id] / sum(out[i].bulk_M[dry_id])*100.0) for i in 1:n_steps];
rho_M      = [ (out[i].rho_M) for i in 1:n_steps];
rho_M[rho_M .== 0.0] .= NaN;
```

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- In the code above, we extract and clean several information**
 - The entropy of all points for checking
 - The fraction of stable melt in mol and we set to NaN the values equal to 0.0 (when melt is not stable)
 - We retrieve the array index of SiO₂ and the indices of all oxides excluding water, this for anhydrous normalization
 - We compute the SiO₂ content of the melt on an anhydrous basis
 - We retrieve the melt density and set to NaN the values equal to 0.0 (when melt is not stable)

MAGeMin_C, isentropic path: visualization

- Visualization is achieved in a similar way as for the fractional crystallization example

```
p1 = plot(T,P, xlabel="Temperature (°C)", marker = :circle, markersize = 2, lw=2, ylabel="Pressure (kbar)", legend=false)
p2 = plot(frac_M,P, xlabel="Melt fraction (mol)", marker = :circle, markersize = 2, lw=2, ylabel="Pressure (kbar)", legend=false)
p3 = plot(rho_M,P, xlabel="Melt density (kg/m³)", marker = :circle, markersize = 2, lw=2, ylabel="Pressure (kbar)", legend=false)
p4 = plot(SiO2_M_dry,P, xlabel="SiO₂ melt anhydrous (mol%)", marker = :circle, markersize = 2, lw=2, ylabel="Pressure (kbar)", legend=false)

fig = plot(p1, p2, p3, p4, layout=(2, 2), size=(800, 600))
savefig(fig,"isentropic_path.png")
```

THIS SNIPPET CAN BE COPIED

- Which yields

