

Electronic Structure: from BlackBoard to Source Code

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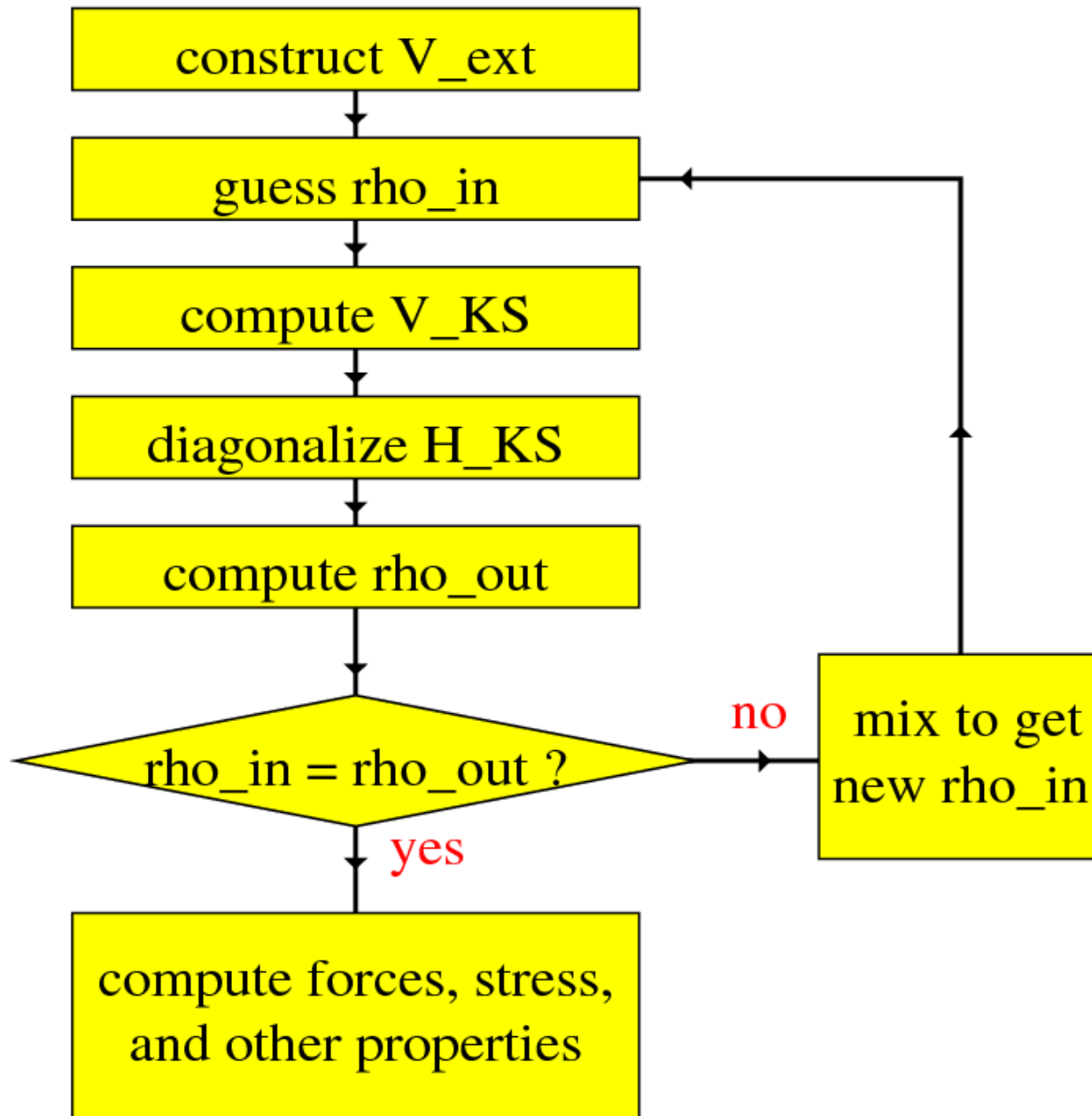
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PWSCF

and

new charge density



PWSCF

```
call read_input_file      (input.f90)

call run_pwscf

call setup                - - > SETUP
call init_run             - - > INIT_RUN
do
  call electrons          - - > ELECTRONS
  call forces
  call stress
  call move_ions
  call update_pot
  call hinit1
end do
```



ELECTRONS

```
call electron_scf
  do iter = 1, niter
    call c_bands      --> C_BANDS
    call sum_band    --> SUM_BAND
    call mix_rho
    call v_of_rho
  end do iter
```

SUM_BAND

```
call weights
do ik = 1, nks
  call get_buffer      (evc)
  call init_us_2      (vkb)
  do ibnd = 1, nbnd
    eband = eband + wg(ibnd, ik) * et(ibnd, ik)
    evc(ig) → psi(nl(igk(ig))) → psi(ir)
    call get_rho      --> GET_RHO
    call sum_bec      --> SUM_BEC
  end do ibnd
end do ik
call addusdens
```

GET_RHO

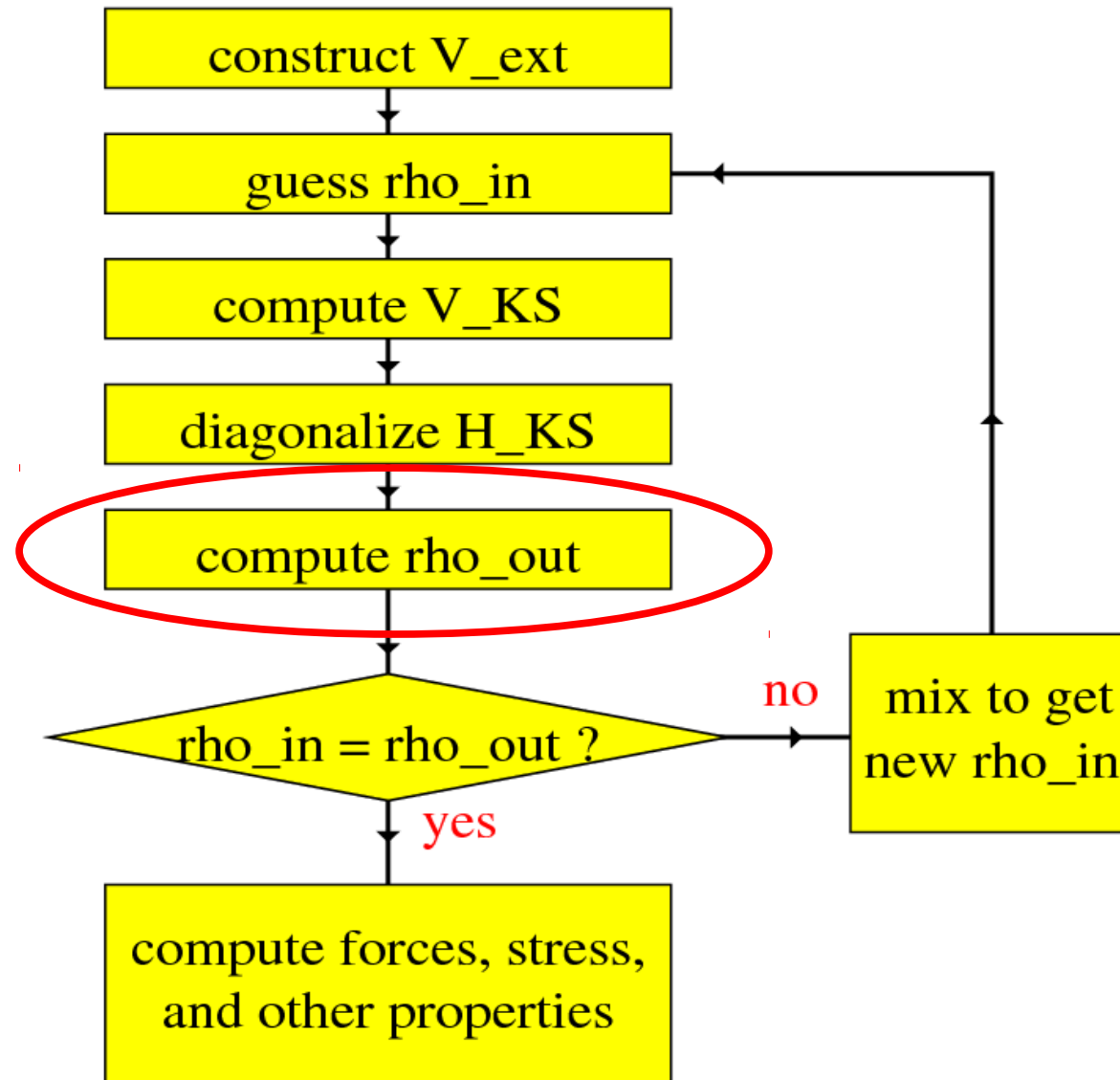
```
rho(ir) = rho(ir) + wg * |psi(ir)|**2
```

SUM_BEC

```
becsum(m, m') = Sum_ik wg <psi|beta_m><beta_m'|psi>
```



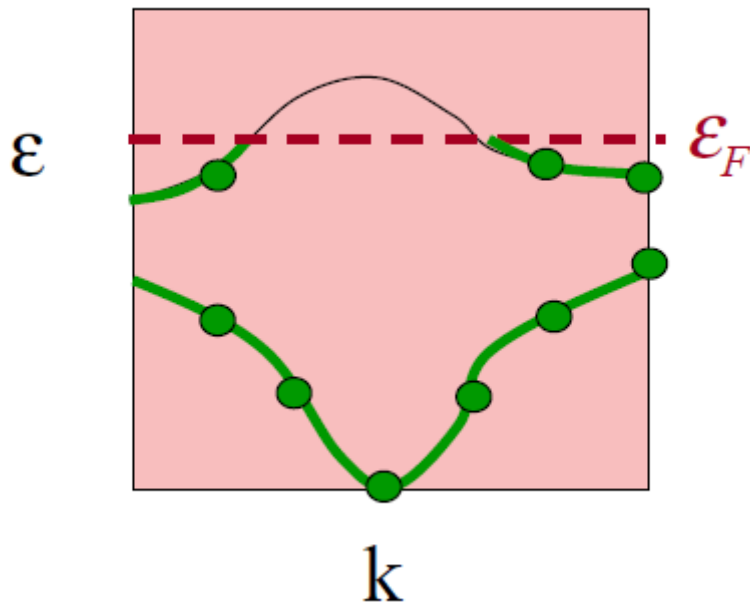
Step 5 : new charge density



Brillouin Zone Sums

Many quantities (e.g., n , E_{tot}) involve sums over k .

- In principle, need infinite number of k 's.
- In practice, sum over a finite number: BZ "Sampling".
- Number needed depends on band structure.
- Typically need more k 's for metals.
- Need to test convergence wrt k -point sampling.



$$\langle P \rangle = \frac{1}{N_k} \sum_{k \in \text{BZ}} P(k) w_k$$

Types of K-points used

Special Points: [Chadi & Cohen]

Points designed to give quick convergence for particular crystal structures.

Monkhorst-Pack grids:

Equally spaced mesh in reciprocal space.

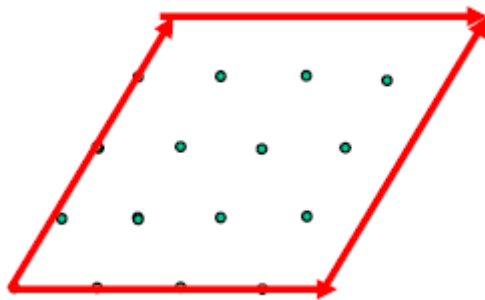
May be centred on origin ['non-shifted'] or not ['shifted']

`K_POINTS {tpiba|crystal|automatic|gamma}`

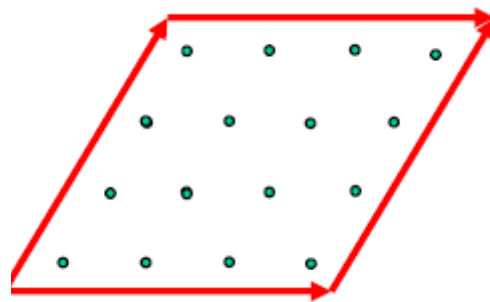
If 'automatic' use M-P grids

`nk1, nk2, nk3, ik1, ik2, ik3`

4 4 4 0 0 0

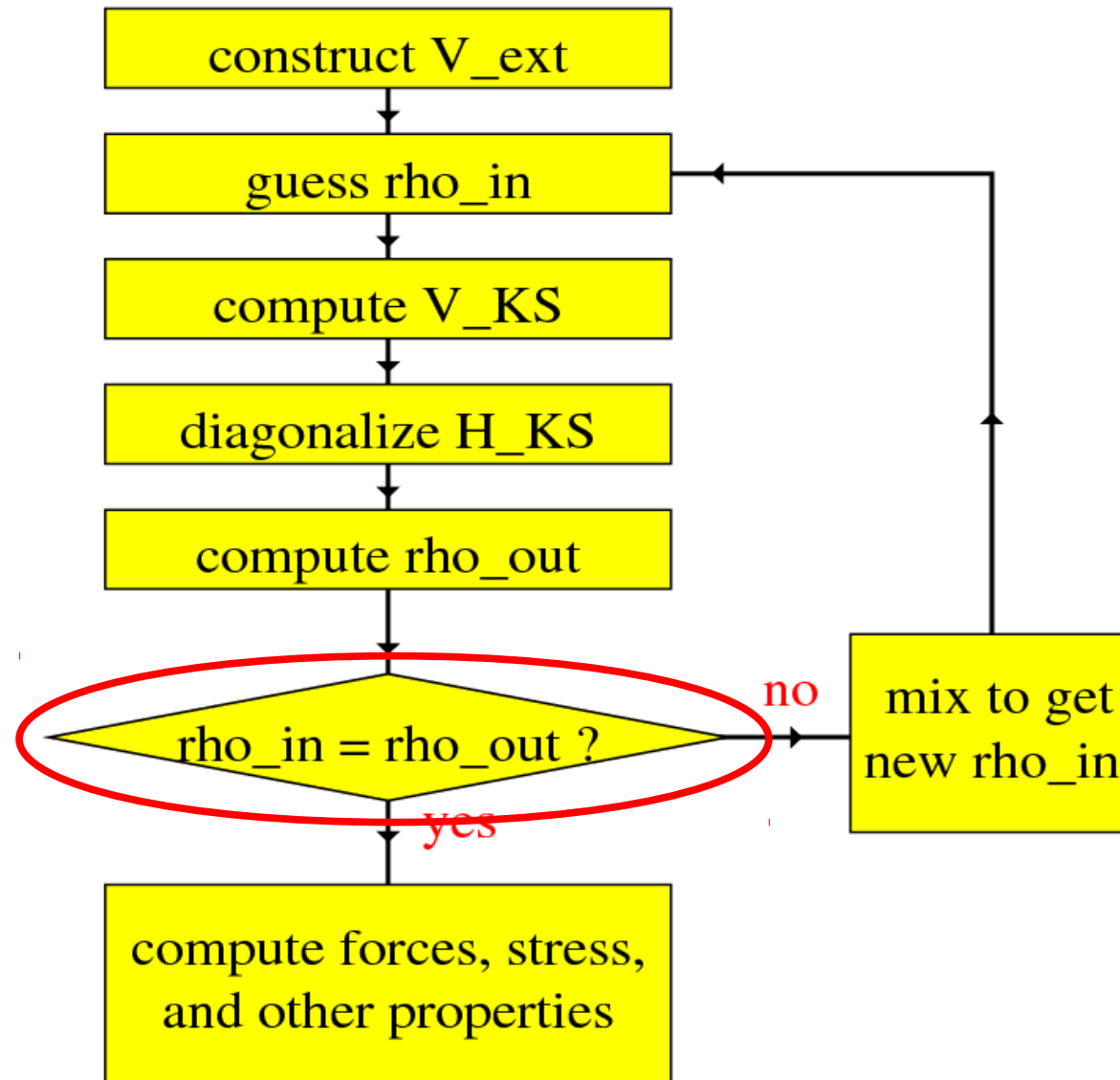


4 4 4 1 1 1



shift

Step 6 : test for convergence



How to decide if converged ?

Check for self-consistency. Could compare:

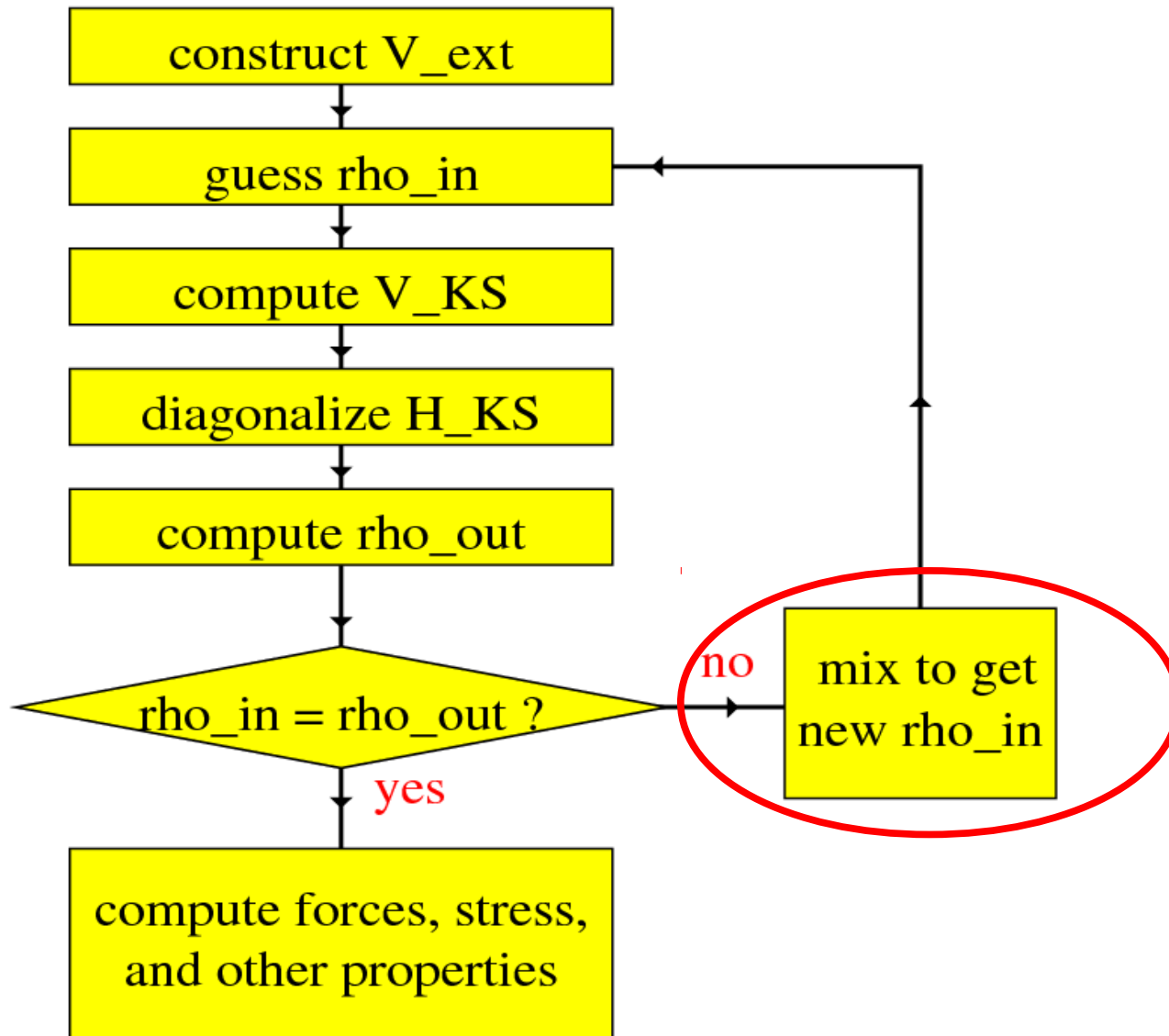
- New and old wavefunctions / charge densities.
- New and old total energies.
- Compare with energy estimated using Harris-Foulkes

Input parameter `conv_thr`
typically ok to use `1.d-8`

Input parameter `electron_maxstep`
maximum number of scf steps performed



Step 7 : mixing



Mixing

Once iteration n of the self-consistent cycle has completed ... how to get next guess for ρ ?

direct iteration in which ρ_{out} is fed directly in ρ_{in}

$$\rho_{in}(n) \rightarrow \rho_{out}(n) \rightarrow \rho_{in}(n+1)$$

usually **doesn't converge**.

One needs to mix, take some combination of input and output densities (may include information from several previous iterations).

Goal is to achieve self consistency ($\rho_{out}=\rho_{in}$) in as few iterations as possible.



Mixing

Simplest prescription: linear mixing

$$\rho_{in}(n+1) = \beta * \rho_{out}(n) + (1-\beta) \rho_{in}(n).$$

Usually slow but should converge for small enough values of β

There exist more sophisticated prescriptions ([Broyden mixing](#), [modified Broyden](#) mixing of various kinds...) based on Quasi Newton Raphson methods.

Input parameter `mixing_mode`

plain | TF | local-TF

Input parameter `mixing_beta`

-Typical values between 0.1 & 0.7
(depend on type of system)



Broyden Mixing

$$\rho_{in}^i = \bar{\rho} + \delta\rho_{in}^i \longrightarrow \rho_{out}^i = \bar{\rho} + \delta\rho_{out}^i$$

In the linear regime if M iterations have been accumulated

$$\rho_{in} = \rho_{in}^M + \sum_{i=1}^{M-1} \alpha_i (\rho_{in}^i - \rho_{in}^{i+1}) \longrightarrow \rho_{out} = \rho_{out}^M + \sum_{i=1}^{M-1} \alpha_i (\rho_{out}^i - \rho_{out}^{i+1})$$

BM determines ρ_{in}^{best} and ρ_{out}^{best} in the already explored manifold by minimizing the norm of $\Delta\rho_{I/O}$ w.r.t. the α_i coefficients and then applies SM to them.

$$\rho_{in}^{new} = \rho_{in}^{best} + \beta \Delta\rho_{I/O}^{best} = (1 - \beta)\rho_{in}^{best} + \beta\rho_{out}^{best}$$



Simple Mixing Revisited

Ideally one would like

$$\rho_{in}^{new} = \rho_{in} - \delta\rho_{in} = \bar{\rho}$$

but we only have access to

$$\Delta\rho_{I/O} = -\chi_0\chi^{-1}\delta\rho_{in}$$

If some simple approximation A to $\chi\chi_0^{-1}$ is available one can then use it to improve the new trial density

$$\delta\rho_{in} \approx A\Delta\rho_{I/O}$$

$$\rho_{in}^{new} = \rho_{in} + \beta A\Delta\rho_{I/O} \approx \rho_{in} - \beta\delta\rho_{in}$$

Thomas-Fermi screening can provide a useful approximate inverse; for very inhomogeneous systems a local TF scheme may be required.



The end : convergence achieved

