


A

 **SCIPION** v2.0 (2019-)

View Model building

- Model building
  - Imports
    - import volumes
    - import atomic structure
    - import sequence
  - Preprocess map
    - xmipp3 - create 3d mask
    - xmipp3 - local MonoRes
    - xmipp3 - localdeblur sharpening
    - xmipp3 - extract unit cell
  - Initial model
    - chimera - model from template
  - Rigid fitting
    - powerfit\_sciption - powerfit
    - chimera - chimera rigid fit
  - Flexible fitting
    - phenix - real space refine
    - ccp4 - coot refinement
    - ccp4 - refmac
  - Validation
    - phenix - emringer**
    - phenix - molprobability
    - phenix - validation\_cryoem
  - Tools-Calculators
    - phenix - superpose pdbs
    - atomstructutils - operator
    - atomstructutils - convert\_sym
    - chimera - chimera operate
    - chimera - chimera restore session
    - chimera - contacts
  - Others
    - xmipp3 - 3d bionotes
  - Exports
    - export to EMDB

B

Protocol Run: PhenixProtRunEMRinger

**Phenix** Protocol: phenix - emringer [Cite](#) [Help](#)

**Run**

Run name phenix - emringer [✎](#) Comment  [✎](#)

Run mode ☒ Continue ☐ Restart [?](#) Host localhost [?](#)

Use queue? ☐ Yes ☒ No [?](#)

Wait for  [?](#)

**Input**

Input Volume  [Q](#) [✎](#) [👁](#) [?](#)

Input atomic structure  [Q](#) [✎](#) [👁](#) [?](#)

[✕](#) Close [💾](#) Save [⚙️](#) Execute