

Introduction

Why radiative transfer?

- ▶ Sensitivity studies: detectability of molecules, etc.
- ▶ Forward model for atmospheric inverse problems

Why line-by-line (lbl)?

- ▶ Modeling and analysis of high resolution spectra
- ▶ “Training set” and benchmark for parameterized models

Why Python?

- ▶ Rapid prototyping
- ▶ Most (?) lbl models kind of “black-box”
- ▶ Difficult to see intermediate quantities

InfraRed Radiative Transfer

Schwarzschild equation: $I(\nu)$ radiance/intensity at wavenumber ν

$$I(\nu) = I_b(\nu) e^{-\tau_b(\nu)} + \int_0^{\tau_b(\nu)} B(\nu, T(\tau')) e^{-\tau'} d\tau'$$

Beer's law: transmission \mathcal{T} and optical depth τ

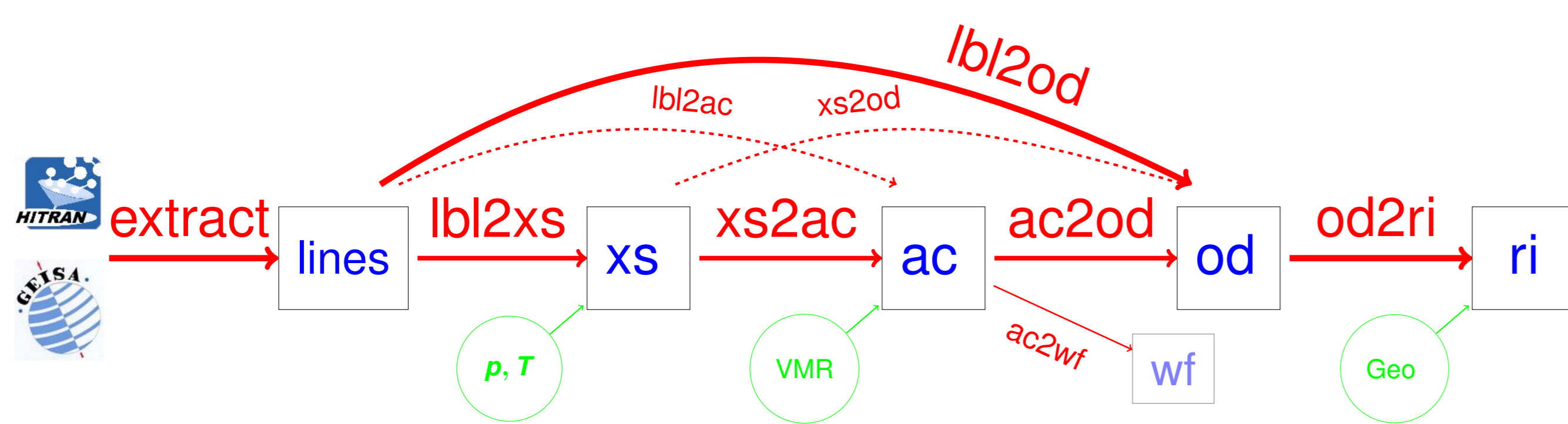
$$\mathcal{T}(\nu, \mathbf{s}) = e^{-\tau} = \exp\left(-\int_0^{\mathbf{s}} ds' \sum_m k_m(\nu, \mathbf{p}(s'), T(s')) n_m(s')\right)$$

Absorption coefficient α and cross section k : line-by-line

$$k(\nu, \mathbf{p}, T) = \sum_l S_l(T) g(\nu; \hat{\nu}_l, \gamma_l(\mathbf{p}, T))$$

Line shape function g : Voigt, Lorentz, VVH, VVW, ...

Line parameters: position $\hat{\nu}_l$, strength S_l , width(s) γ_l , ...



Py4CATS — Implementation

- ▶ (Numeric and Scientific) Python version of Fortran 2008
- ▶ “Generic Atmospheric Radiation Lbl Ir Code” GARLIC [4]
- ▶ Series of scripts for IR & μ Wave radiative transfer, e.g.,
 - ▶ `extract` lines of relevant molecules in the spectral range of interest
 - ▶ `l2xs` line-by-line cross sections for given pressure(s) & temperature(s)
 - ▶ `xs2ac` multiply cross sections with densities and sum over all molecules
 - ▶ `ac2od` integrate absorption coefficients along line-of-sight to optical depth
 - ▶ `od2ri` integrate Schwarzschild eq. along line-of-sight to radiance/intensity
 - ▶ `ac2wf` compute weighting functions $\partial\mathcal{T}/\partial\mathbf{z} \propto \alpha\mathcal{T}$
 - ▶ and some shortcuts, e.g., `l2ac` or `l2od`
- ▶ **New:** functions accessible within (I)Python shell
- ▶ Sub-classed numpy arrays `xsArray`, `acArray`, `odArray`, ... for cross sections, absorption coefficients, optical depths, ... to store “spectra” along with attributes (e.g. `xs.p` and `xs.t`)
- ▶ Numerics:
 - ▶ Complex error function: Humlíček [1] – Weideman [5] combination [3]
 - ▶ Multigrid line-by-line (fine grid near line center only) [2]
 - ▶ Schwarzschild integral: B linear or exponential in τ
- ▶ Limitations:
 - ▶ Plots for quicklook only, not “publication-ready”
 - ▶ Plane-parallel atmosphere, no scattering, continua, ...
 - ▶ No “package” yet, no distutils etc. (coming soon)

References:

- [1] J. Humlíček. Optimized computation of the Voigt and complex probability function. *JQSRT*, 27:437–444, 1982.
- [2] F. Schreier. Optimized evaluation of a large sum of functions using a three-grid approach. *Comp. Phys. Comm.*, 174:783–802, 2006.
- [3] F. Schreier. Optimized implementations of rational approximations for the Voigt and complex error function. *JQSRT*, 112:1010, 2011.
- [4] F. Schreier, S. Gimeno García, P. Hedelt, M. Hess, J. Mendrok, M. Vasquez, and J. Xu. GARLIC – a general purpose atmospheric radiative transfer line-by-line infrared-microwave code: Implementation and evaluation. *JQSRT*, 137:29–50, 2014.
- [5] J.A.C. Weideman. Computation of the complex error function. *SIAM J. Num. Anal.*, 31:1497–1518, 1994.

IPython Demo

(output largely deleted)

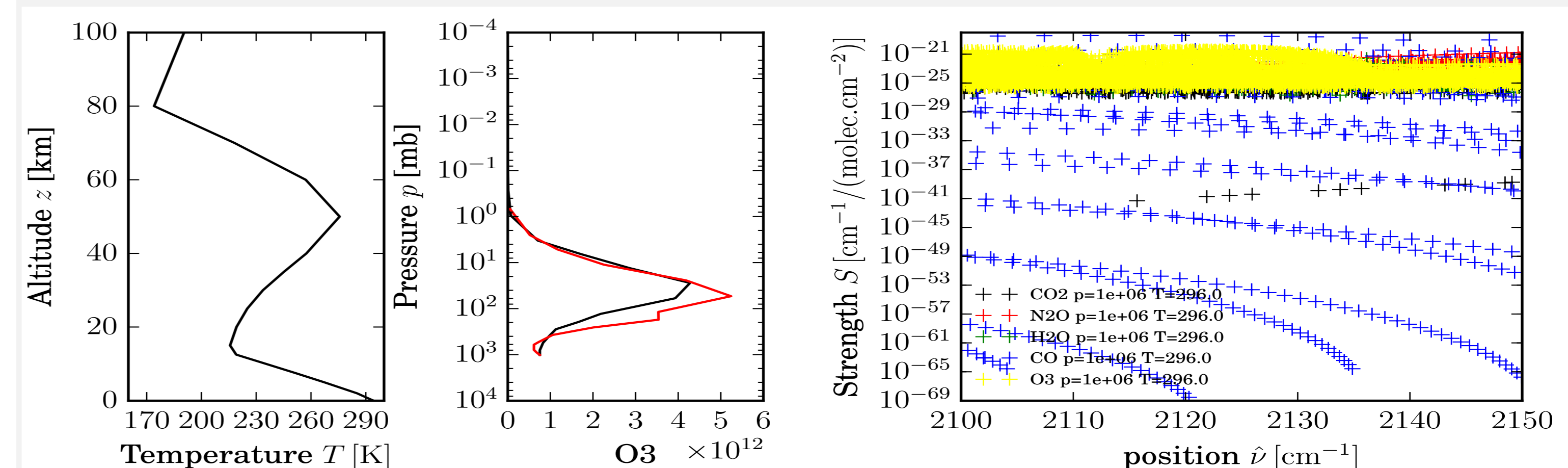
```
Python 2.7.3 (default, Apr 14 2012, 08:58:41) [GCC]
IPython 2.0.0 -- An enhanced Interactive Python.
```

```
In[1]: # get two mid latitude atmospheres
...: mls = atmos1D('/data/atmos/20/mls.xy')
...: mlw = atmos1D('/data/atmos/20/mlw.xy', zToA=50)
```

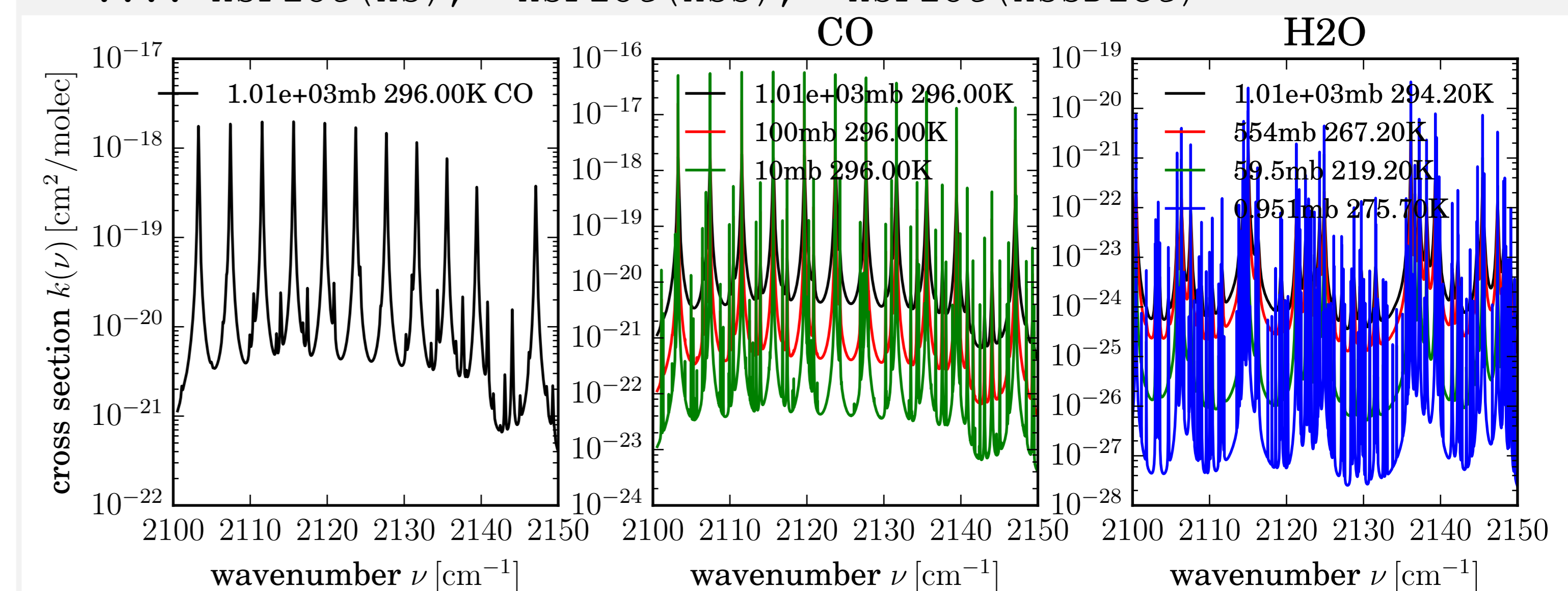
```
Atmos1d: got p, T, air and 7 gases at 20 levels
Atmos1d: got p, T, air and 7 gases at 16 levels
```

```
In [2]: # IASI microwindow for CO retrieval: HITRAN-GeiSa-exTRACT
...: dictLineLists = higstract('/data/geisa/87/lines',
...:                           (2100,2150), molecule='main')
9771 lines of 5 molecule(s), returning a dictionary
```

```
In [3]: atmPlot(mls); atmPlot([mls,mlw], 'O3', 'mb')
...: atlas(dictLineLists) # plot line data (default strength)
```



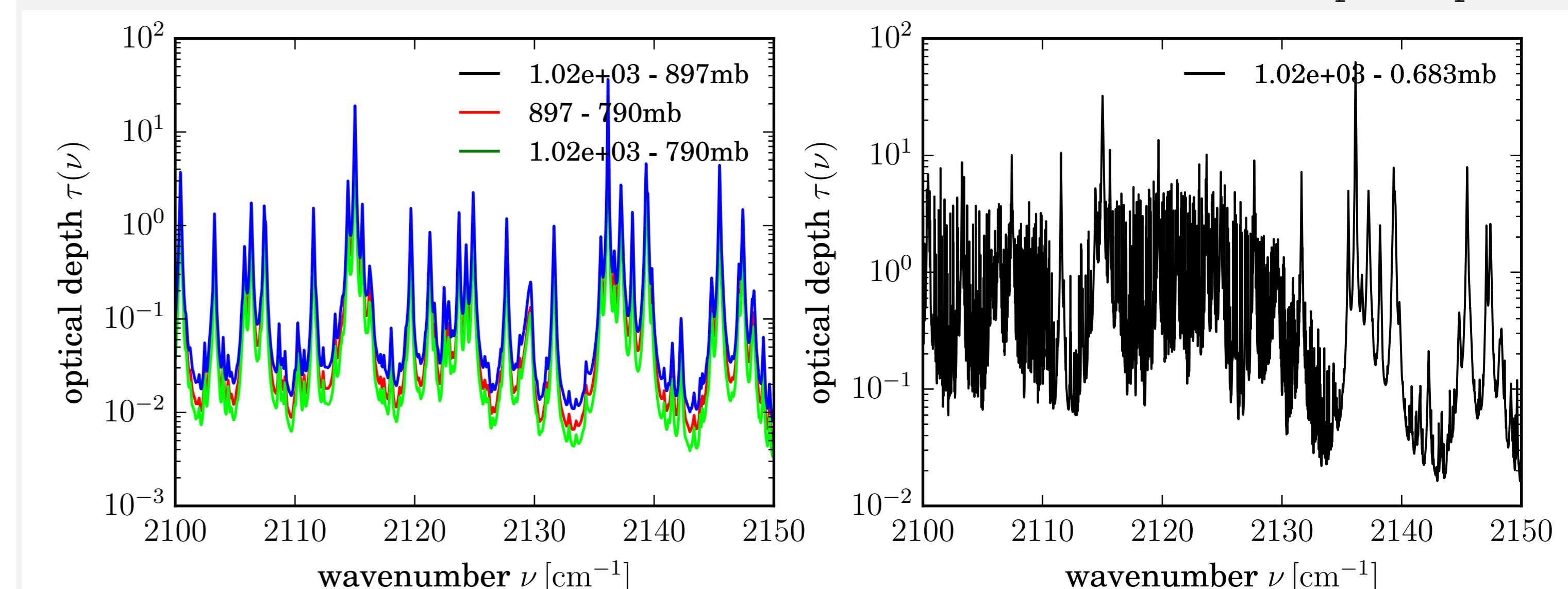
```
In [4]: # CO cross section at database pressure and temperature
...: xs = lbl2xs(dictLineLists['CO'])
...: # a list of cross sections for three pressures
...: xss = lbl2xs(dictLineLists['CO'], [1013,100,10,'mb'])
...: # a dictionary of x-section lists (for all p, T, gases)
...: xssDict = lbl2xs(dictLineLists,mls['p'],mls['T'])
...: # ... and some plots (not all are shown here)
...: xsPlot(xs); xsPlot(xss); xsPlot(xssDict)
```



```
In [5]: # proceed step-by-step
...: acList = xs2ac(mls, xssDict) # absorption coefficients
...: dodList = ac2dod(acList) # delta optical depths
```

```
In [6]: # alternatively bypass intermediate quantities, e.g.
...: dodList = lbl2dod(mls,dictLineLists) # delta opt.depths
```

```
In [7]: # sum/combine optical depths and plot
...: odPlot([dodList[0], dodList[1]]) # the bottom layers,
...: odPlot(dodList[0]+dodList[1]) # ... their sum,
...: odPlot(dod2tod(dodList)) # and total opt.depth
```



```
In [8]: # radiation intensity seen by uplooking observer at BoA
...: vGrid, radUp = dod2ri(dodList)
...: # and downlooking observer at ToA (incl. surface @ 294K)
...: vGrid, radNadir = dod2ri(dodList, 180, 294.2)
```