



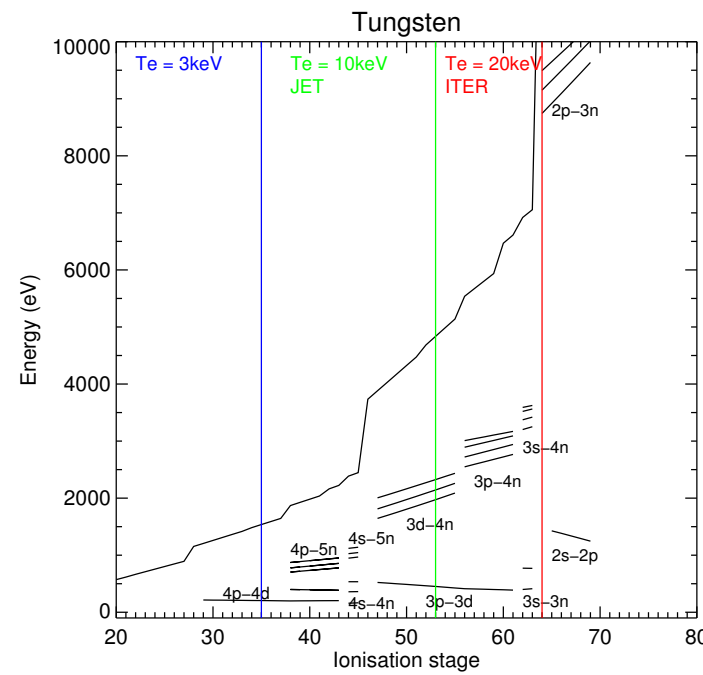
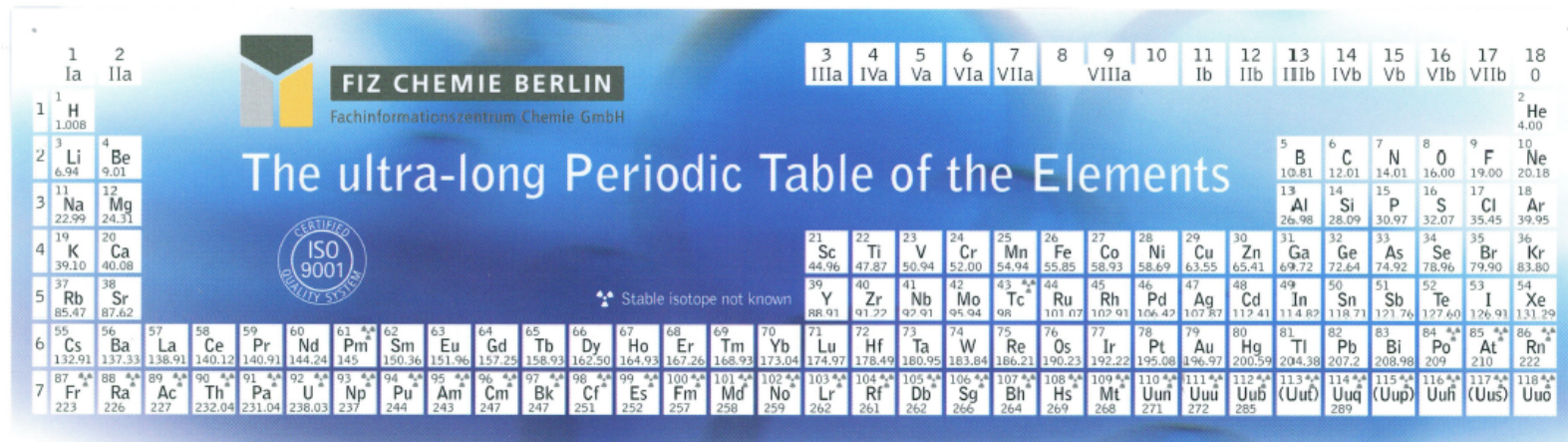
Heavy Species in ADAS

from the viewpoint of one lowly ion

Martin O'Mullane, Adam Foster, Hugh Summers and Allan Whiteford

Department of Physics
University of Strathclyde

Atomic data requirements



Three aspects of the heavy species question

The data we need

- ▶ Source functions — *adf11 acd, scd, ccd*
- ▶ Power coefficients — *adf11 plt, prb*
- ▶ Line emission — *adf15 pec*
- ▶ Spectral feature emission — *adf40 f-pec*

How to get it

- ▶ Scoping the problem
- ▶ Automated generation

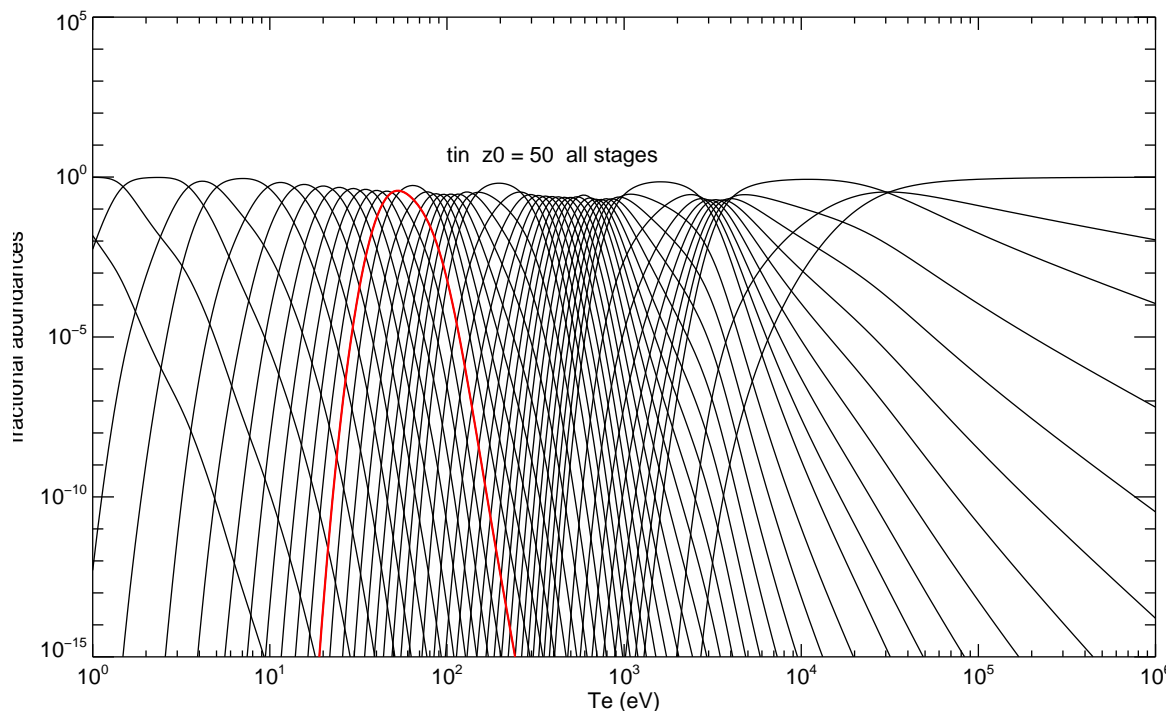
How to use it

- ▶ Potentially large datasets
- ▶ Partitioning and superstaging.

Let's choose tin (Z=50)

First questions: Where do its stages radiate? And what if there is no helpful ADAS *adf11* data?

```
preview_natural_partition, z0=50, plot_type=3, $  
    te_min=1, te_max=1e6, $  
    te_plot=1000,           $  
    frac=frac  
oplot, frac.te, frac.ion_all[13,*], color=5, thick=5
```



Now let's narrow our focus to Sn¹³⁺

- ▶ What is its ground state configuration?
- ▶ What configurations contribute to spectral emission?
- ▶ And to radiated power?
- ▶ How do we choose which ones to include?

The *adf00* set archives ionisation potential and ground configurations:

tin	-50	0	1	2	3	4	5	6	7	8	9	10	11	12	13	14
		7.343d+00	1s2	2s2	2p6	3s2	3p6	3d10	4s2	4p6	4d10	4f0	5s2	5p2		
		1.463d+01	1s2	2s2	2p6	3s2	3p6	3d10	4s2	4p6	4d10	4f0	5s2	5p1		
								..								
								..								
12	2.744d+02	1s2	2s2	2p6	3s2	3p6	3d10	4s2	4p6	4d2						
13	2.995d+02	1s2	2s2	2p6	3s2	3p6	3d10	4s2	4p6	4d1						
14	3.959d+02	1s2	2s2	2p6	3s2	3p6	3d10	4s2	4p6							
								..								

What configurations should be considered?

With a ground state of $3d^{10}4s^24p^64d^1$ we can

- ▶ promote the valence 4d electron to any higher n/l shell
- ▶ allow 4s or 4p electrons to be excited
- ▶ or any other electron — from 2p perhaps?
- ▶ however where do we stop in Δn or Δl ?
- ▶ and how many configurations should we consider?

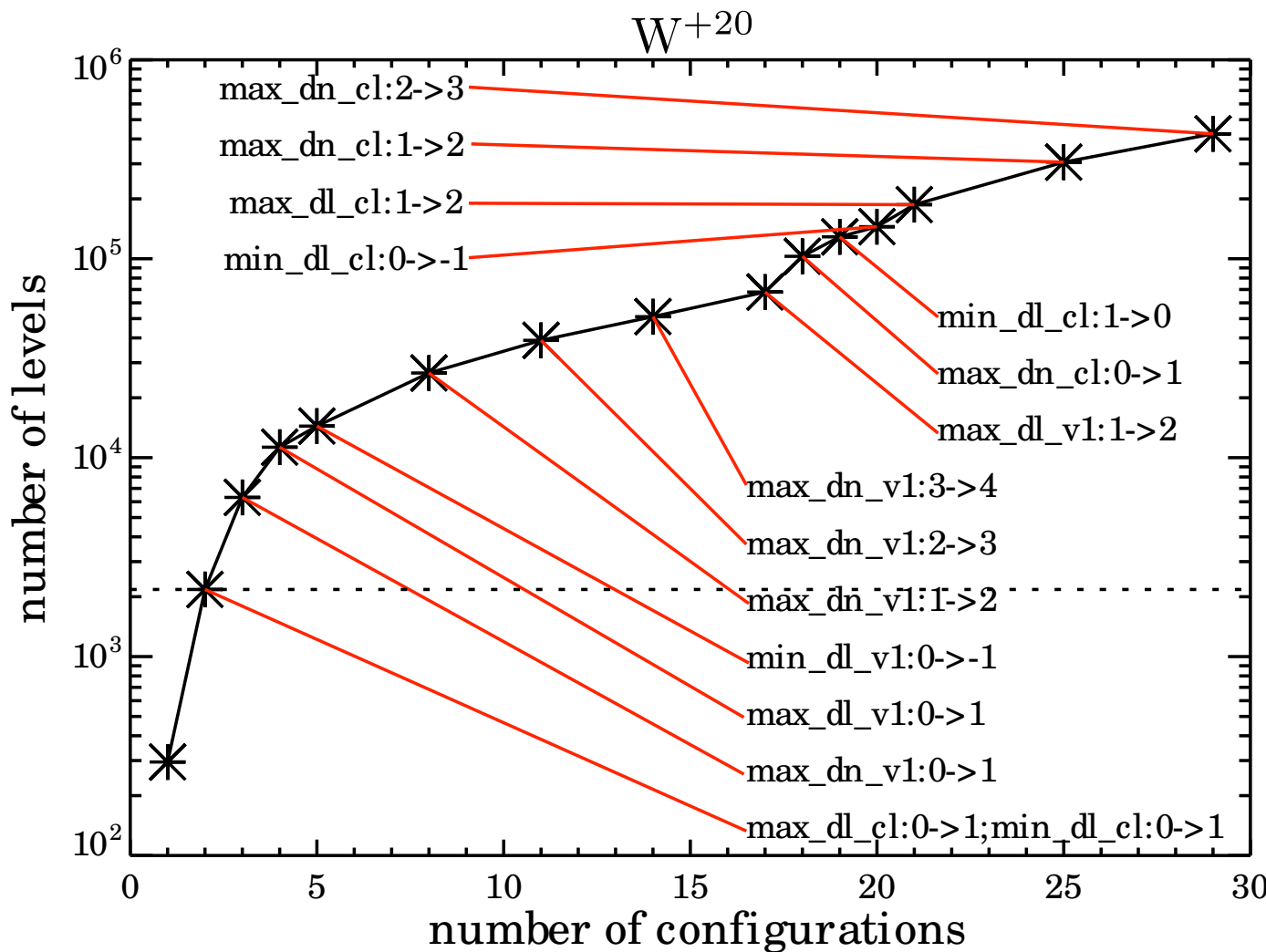
There are 180 distinct ground configurations (for elements up to Radon)

A rule based method is desirable (essential!)

ADAS rules for choosing where to promote electrons

<i>index[]</i>	:	index of ground configuration of each ion of element in <i>adf54</i> file
<i>config[]</i>	:	ground configuration for each ion of element
<i>n_el[]</i>	:	number of electrons for each ion of element
<i>no_v_shl[]</i>	:	number of open (valence) shells. Include outer-most shell even if closed.
<i>max_dn_v1[]</i>	:	maximum Δn promotion for first (outer-most) valence shell.
<i>min_dn_v1[]</i>	:	minimum Δn promotion for first (outer-most) valence shell. Negative value allows access to inner unoccupied or open shells
<i>max_dl_v1[]</i>	:	maximum delta Δl promotion for first (outer-most) valence shell.
<i>min_dl_v1[]</i>	:	minimum delta Δl promotion for first (outer-most) valence shell.
<i>max_dn_v2[]</i>	:	maximum Δn promotion for second (inner-most) valence shell.
<i>min_dn_v2[]</i>	:	maximum Δn promotion for second (inner-most) valence shell.
<i>max_dl_v2[]</i>	:	maximum delta Δl promotion for second (inner-most) valence shell.
<i>min_dl_v2[]</i>	:	minimum delta Δl promotion for second (inner-most) valence shell.
<i>prom_cl[]</i>	:	promote from inner shell closed shells (1=yes,0=no).
<i>max_n_cl[]</i>	:	maximum inner shell <i>n</i> from which promotions are permitted.
<i>min_n_cl[]</i>	:	minimum inner shell <i>n</i> from which promotions are permitted.
<i>max_l_cl[]</i>	:	maximum inner shell <i>l</i> from which promotions are permitted.
<i>min_l_cl[]</i>	:	minimum inner shell <i>l</i> from which promotions are permitted.
<i>max_dn_cl[]</i>	:	maximum Δn promotion from a permitted inner shell.
<i>min_dn_cl[]</i>	:	minimum Δn promotion from a permitted inner shell. Negative values of Δn allow access to inner unoccupied or open shells.
<i>max_dl_cl[]</i>	:	maximum Δl promotion from a permitted inner shell.
<i>min_dl_cl[]</i>	:	minimum Δl promotion from a permitted inner shell.
<i>fill_n_v1[]</i>	:	add all <i>nl</i> configurations of outer valence shell <i>n</i> (1=yes,0=no).
<i>fill_par[]</i>	:	if <i>n_fill</i> only add opposite parity to valence shell else add both parities (1=yes, 0=n0).
<i>for_tr_sel[]</i>	:	Cowan option for radiative transitions 1 - first parity, 2 or 3(default).
<i>last_4f[]</i>	:	shift an electron valence shell to unfilled 4f as extra ground.
<i>grd_cmplx[]</i>	:	include configurations of same complex as ground configuration for valence <i>n</i> -shell.

adf54 : rules for automatic data generation



Care needed!! resolved calculations (ic or LS) can overwhelm computers.

Work through Sn¹³⁺

- ▶ Within ADAS the generation of heavy species data is almost exclusively a non-GUI activity.
- ▶ The outputs are standard *adf11*, *adf15* and *adf40* datasets which can be used and examined with the GUI interactive system.

At the IDL command line:

```
; Let's choose Sn13+
```

```
z_nuc = 50  
z_ion = 13  
tag = xxesym(z_nuc, /lower) + string(z_ion, format='(i2.2)')
```

```
; Use promotion rules from W work
```

```
a54file = '/u/adas/adas/adf54/promotion_rules_w_adf54.dat'
```

```

adas8xx_promotion_rules, z0_nuc = z_nuc, z_ion = z_ion, ionpot = ip,  $
                                prom_rules=rules, a54file = file
help, rules, /st

** Structure <9b54e9c>, 25 tags, length=60, data length=60, refs=1:
CONFIG      STRING ' 1s2  2s2  2p6  3s2  3p6  3d10 4s2  4p6  4d1'
INDEX       INT    129
NO_V_SHL   INT    1
MAX_DN_V1  INT    3
MIN_DN_V1  INT    0
MAX_DL_V1  INT    2
MIN_DL_V1  INT   -2
MAX_DN_V2  INT    0
MIN_DN_V2  INT    0
MAX_DL_V2  INT    0
MIN_DL_V2  INT    0
PROM_CL    INT    1
MAX_N_CL   INT    4
MIN_N_CL   INT    4

```

MAX_L_CL	INT	1
MIN_L_CL	INT	0
MAX_DN_CL	INT	1
MIN_DN_CL	INT	0
MAX_DL_CL	INT	2
MIN_DL_CL	INT	0
FILL_N_V1	INT	1
FILL_PAR	INT	0
FOR_TR_SEL	INT	3
LAST_4F	INT	0
GRD_CMPLX	INT	0

```

adas8xx_promotions, z0_nuc = z_nuc, z_ion = z_ion, ionpot = ip,  $
                           prom_rules          = rules,           $
                           promotion_results = results

help, results, /st

** Structure <9b530dc>, 11 tags, length=2496, data length=2496, refs=1:
   GRD_CFG           STRING    '4d1  '
   GRD_OCC           INT       Array[36]
   EX_CFG            STRING    Array[25]
   GRD_PAR           INT       0
   EX_PAR            INT       Array[25]
   GRD_ZC_COW        LONG      -14
   EX_ZC_COW         LONG      Array[25]
   OC_STORE          INT       Array[36, 26]
   NO_CONFIGS        LONG      Array[7]
   NO_TERMS          LONG      Array[7]
   NO_LEVELS         LONG      Array[7]

```

```
print, results.grd_occ
```

2	2	6	2	6	10	2	6	1	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0

```
print, results.oc_store[:,1]
```

2	2	6	2	6	10	2	6	0	1
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0

```
print, results.oc_store[:,2]
```

2	2	6	2	6	10	2	6	0	0
1	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0
0	0	0	0	0	0	0	0	0	0

```

; Write CA driver files for restricted plasma parameters

files = { adf34_file      : 'adf34_ca_' + tag + '.dat', $  

          adf42_ca_file : 'adf42_ca_' + tag + '.dat', $  

          adf04_ca_file : 'adf04_ca_' + tag + '.dat', $  

          adf40_ca_file : 'adf40_ca_' + tag + '.dat', $  

          adf15_ca_file : 'adf15_ca_' + tag + '.dat', $  

          adf11_ca_file : 'adf11_ca_' + tag + '.dat' }

plasma = {theta           : [ 1.0e3, 2.0e3, 5.0e3, 1.0e4, 1.5e4, $  

                               2.0e4, 5.0e4, 1.0e5], $  

          idx_theta       : indgen(8),  

          rho            : [ 1.0e8, 1.0e10, 1.0e12, 1.0e14],  

          idx_rho         : indgen(4),  

          npix           : [ 128,    256],  

          wvlmin         : [100.0,   1.0],  

          wvlmax         : [150.0, 500.0],  

          idx_wvl        : indgen(2),  

          theta_noscale : 0,  

          rho_scale      : 0

```

```
adas8xx_create_drivers, z0_nuc=z_nuc, z_ion=z_ion, ionpot=ip, $  
promotion_results=results, $  
plasma=plasma, files=files
```

The driver file for ADAS801 (Cowan code):

2	-5	2	10	1.0	5.d-09	5.d-11-2	0130	1.0	0.65	0.0	0.5
50	-14	Sn	ground	z1=13	0	4d1					
50	-14	Sn	cfg	01	0	5s1					
50	-14	Sn	cfg	02	0	5d1					
50	-14	Sn	cfg	03	0	5g1					
50	-14	Sn	cfg	04	0	6s1					
50	-14	Sn	cfg	05	0	6d1					
50	-14	Sn	cfg	06	0	6g1					
50	-14	Sn	cfg	07	0	7s1					
50	-14	Sn	cfg	08	0	7d1					
50	-14	Sn	cfg	09	0	7g1					
50	-32	Sn	cfg	10	0	3d10	4s1	4p6	4d2		
50	-32	Sn	cfg	11	0	3d10	4s1	4p6	4d1	5s1	

50	-32	Sn	cfg	12	0	3d10	4s1	4p6	4d1	5d1
50	-32	Sn	cfg	13	0	3d10	4s2	4p5	4d1	4f1
50	-32	Sn	cfg	14	0	3d10	4s2	4p5	4d1	5p1
50	-32	Sn	cfg	15	0	3d10	4s2	4p5	4d1	5f1
50	-14	Sn	cfg	16	1	4f1				
50	-14	Sn	cfg	17	1	5p1				
50	-14	Sn	cfg	18	1	5f1				
50	-14	Sn	cfg	19	1	6p1				
50	-14	Sn	cfg	20	1	6f1				
50	-14	Sn	cfg	21	1	7p1				
50	-14	Sn	cfg	22	1	7f1				
50	-32	Sn	cfg	23	1	3d10	4s1	4p6	4d1	5p1
50	-32	Sn	cfg	24	1	3d10	4s2	4p5	4d2	
50	-32	Sn	cfg	25	1	3d10	4s2	4p5	4d1	5d1
-1										

Back to the IDL command line:

```
; Run the CA structure code
```

```
adas8xx_create_ca_adf04, z_ion, $  
      z_nuc, $  
      results.oc_store, $  
      ionpot          = ip, $  
      plasma          = plasma, $  
      adf04_t3_file   = files.adf04_ca_file
```

adf04 file for Sn¹³⁺

Sn+13	50	14	2415630.6								
1 19		(0)0(4.5)								0.0
2 606527558529		(0)0(134.5)								604454.8
3 1B		(0)0(0.5)								656865.3
4 1A		(0)0(6.5)								664371.3
5 1C		(0)0(2.5)								810958.4
6 1D		(0)0(4.5)								1048671.2
7 606517568529		(0)0(44.5)								1052972.9
8 60652755851951A		(0)0(419.0)								1259268.0
9 1E		(0)0(6.5)								1290521.2
10 1G		(0)0(0.5)								1291113.7
11 1H		(0)0(2.5)								1366810.5
12 1F		(0)0(8.5)								1400752.2
13 60652755851951C		(0)0(179.5)								1412966.0
14 1I		(0)0(4.5)								1485688.9
15 1J		(0)0(6.5)								1609362.8
16 1M		(0)0(0.5)								1611744.1
17 60652755851951D		(0)0(299.5)								1649394.9
18 1N		(0)0(2.5)								1654541.6
19 1K		(0)0(8.5)								1668939.1
20 60651756851951B		(0)0(19.5)								1715621.3
21 1O		(0)0(4.5)								1722798.2
22 1P		(0)0(6.5)								1795180.9
23 1Q		(0)0(8.5)								1831229.6
24 60651756851951C		(0)0(59.5)								1867168.5
25 60652755851951E		(0)0(419.0)								1891442.4
26 60651756851951D		(0)0(98.5)								2103915.7
-1											
14.0	3	1.96+05	3.92+05	9.80+05	1.96+06	2.94+06	3.92+06	9.80+06	1.96+07		
4	1	1.92+11	7.92+00	8.17+00	8.96+00	1.02+01	1.12+01	1.20+01	1.54+01	1.87+01	
3	1	7.69+06	1.99-01	2.02-01	2.09-01	2.17-01	2.23-01	2.27-01	2.39-01	2.47-01	
5	1	1.07+11	3.89-01	4.07-01	4.70-01	5.77-01	6.73-01	7.59-01	1.13+00	1.50+00	
6	1	1.17+07	7.21-01	7.29-01	7.54-01	7.91-01	8.20-01	8.42-01	9.17-01	9.69-01	
9	1	2.22+10	1.34-01	1.35-01	1.35-01	1.37-01	1.39-01	1.41-01	1.59-01	1.83-01	
12	1	1.21+08	5.57-01	5.62-01	5.81-01	6.11-01	6.36-01	6.57-01	7.33-01	7.89-01	
10	1	2.97+06	2.19-02	2.19-02	2.22-02	2.25-02	2.27-02	2.29-02	2.35-02	2.39-02	

Eissner notation — quick recap

Each occupation/orbital-list pair is separated from the next by 5 (or 6)

1s

1

2s 2p

2 3

3s 3p 3d

4 5 6

4s 4p 4d 4f

7 8 9 A

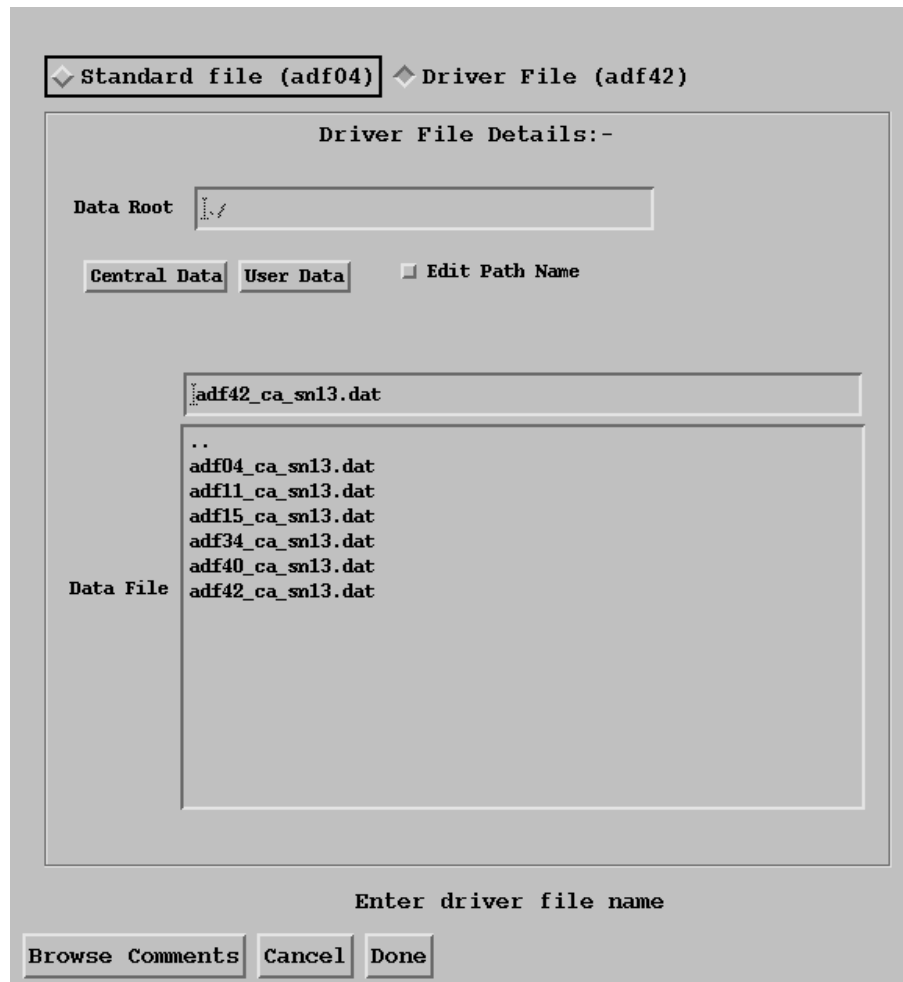
5s 5p 5d 5f 5g

B C D E F

etc.

Generating spectral and power data — ADAS810

Process the *adf42* file made by `adas8xx_create_drivers` with ADAS810 to generate *adf11/plt*, *adf15* and *adf40* datasets.



Title for Run [] Nuclear Charge: 50 Ion Charge: 13

Input from adf42 file : ./adf42_ca_sn13.dat [Browse Comments](#)

adf04 file : adf04_ca_sn13.dat [Browse Comments](#)
No expansion data

Temperatures				
INDEX	Electron	Ion	Neutral Hydrogen	Input Value
1	1.689E+01	0.000E+00	0.000E+00	1.689E+01
2	3.378E+01	0.000E+00	0.000E+00	3.378E+01
3	8.444E+01	0.000E+00	0.000E+00	8.445E+01
4	1.689E+02	0.000E+00	0.000E+00	1.689E+02
5	2.533E+02	0.000E+00	0.000E+00	2.533E+02

Temperature Units: eV

[Edit Table](#)

Default: Standard Set

Densities			
INDEX	Electron Densities	Ion Densities	
1	1.000E+08	0.000E+00	
2	1.000E+10	0.000E+00	
3	1.000E+12	0.000E+00	
4	1.000E+14	0.000E+00	
5			

Density Units: cm⁻³

[Edit Table](#)

Default: Standard Set

Metastable State			
<input type="checkbox"/> 19			
<input type="checkbox"/> 606527558529			
<input type="checkbox"/> 1B			
<input type="checkbox"/> 1A			

For a single metastable normalise PLT and PEC ?

NO

YES

[Edit Table](#)

Lower limit of A-value: 0.00000

Include Reactions:

- Ionisation Rates
- Charge Exchange
- Recombination
- Inner Shell Ionisation
- Include Projection Data
- Proton Impact Collisions

Zeff : 3.00000

Spectral Intervals

INDEX	# pixels	min wave	max wave
1	128	100.00	150.00
2	256	1.00	500.00
3			
4			
5			

Edit the processing options data and press Done to proceed

[Cancel](#) [Done](#)

Text Output Replace Default File Name

File Name : paper-810.txt

PEC (adf15) file Replace Default File Name

File Name : adf15_ca_sn13.dat

Feature PEC (adf40) file Replace Default File Name

File Name : adf40_ca_sn13.dat

Total power (adf11/plt) unfiltered Replace Default File Name

File Name : adf11_ca_sn13.dat

Total power (adf11/plt) filtered Replace Default File Name

File Name : NULL

Choose output options

 Cancel Done

Where lies the emission?

Back to the IDL command line!

```
read_adf40,file='adf40_ca_sn13.dat', fulldata=all
```

```
help, all, /st
```

ESYM	STRING	'Sn'
IZ0	LONG	50
IS	LONG	13
IS1	LONG	14
NBLOCK	LONG	2
NPIX	LONG	Array[2]
WAVE_MIN	DOUBLE	Array[2]
WAVE_MAX	DOUBLE	Array[2]
NTE	LONG	Array[2]
TE	DOUBLE	Array[8, 2]
NDENS	LONG	Array[2]
DENS	DOUBLE	Array[4, 2]
FPEC	DOUBLE	Array[256, 8, 4, 2]
TYPE	STRING	Array[2]

```

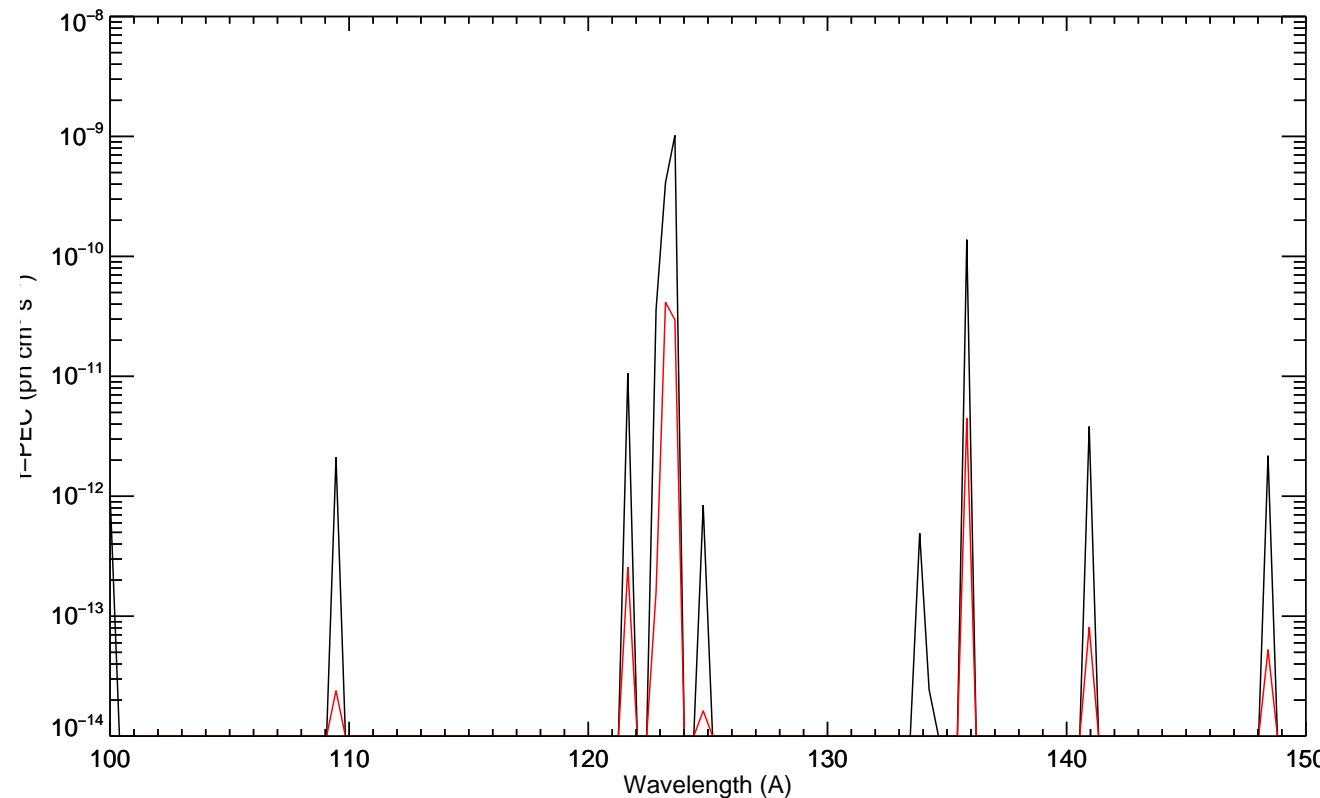
wave=adas_vector(low=all.wave_min[0], high=all.wave_max[0], $  

                  num=all.npix[0], /linear)

plot_io, wave, all.fpec[* , 7, 2, 0] > 1e-14, $  

          xtitle='Wavelength (A)', ytitle = 'f-PEC (ph cm!u3!n s!u-1!n)'
oplot, wave, all.fpec[* , 1, 2, 0] > 1e-14, color=5

```



How to identify contributing configurations

It depends of the width of the spectral region of interest

C	lv	configuration	(2S+1)L(w-1/2)	energy (cm^-1)
C	---	-----	-----	-----
C	1	19	(0)0(-4.5)	0.0
C	2	606527558529	(0)0(134.5)	604454.8
C	3	1B	(0)0(-0.5)	656865.3
C	4	1A	(0)0(-6.5)	664371.3
C	5	1C	(0)0(-2.5)	810958.4
C	6	1D	(0)0(-4.5)	1048671.2
C	7	606517568529	(0)0(-44.5)	1052972.9
C			-	-
C	20	60651756851951B	(0)0(19.5)	1715621.3
C	21	10	(0)0(-4.5)	1722798.2
C	22	1P	(0)0(-6.5)	1795180.9
C			-	-
C	12	94.9692	7(0)0(-44.5)-	1(0)0(-4.5)
C	13	99.5453	19(0)0(-8.5)-	4(0)0(-6.5)
C	14	100.233	18(0)0(-2.5)-	3(0)0(0.5)
C	15	109.668	21(0)0(-4.5)-	5(0)0(-2.5)
C	16	121.756	14(0)0(-4.5)-	4(0)0(-6.5)
C	17	122.821	24(0)0(-59.5)-	7(0)0(-44.5)
C	18	123.311	5(0)0(-2.5)-	1(0)0(-4.5)
C	19	123.684	13(0)0(179.5)-	2(0)0(134.5)
C	20	124.877	16(0)0(0.5)-	5(0)0(-2.5)
C				excit 1 1 13 17 48 12
C				excit 1 1 13 33 14 13
C				excit 1 1 13 44 43 14
C				excit 1 1 13 54 39 15
C				excit 1 1 13 31 28 16
C				excit 1 1 13 117 18 17
C				excit 1 1 13 3 6 18
C				excit 1 1 13 124 5 19
C				excit 1 1 13 52 44 20

- ▶ Overplot/look at PEC *adf15* data.
- ▶ Refine promotion rules or *adf34* driver to home-in
- ▶ Note that structure codes are not spectroscopically accurate.

Where next?

Identify emission region of interest — treat these in intermediate coupling.

For Sn¹³⁺ :

- ▶ 26 configurations, 226 terms, 554 levels

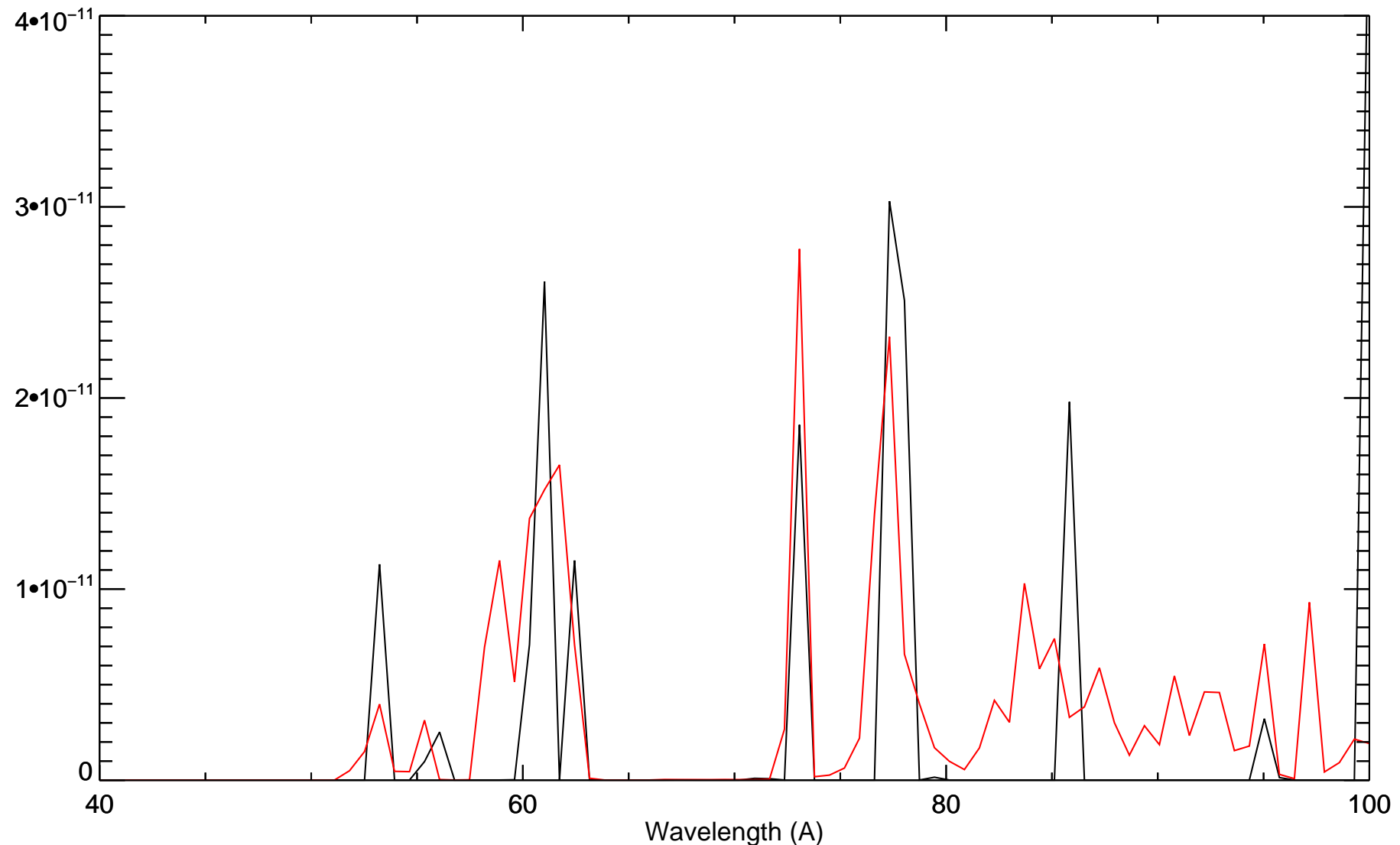
```
read_adf40,file='fpec40#sn_ca#sn13.dat', fulldata=all_ca
read_adf40,file='fpec40#sn_ic#sn13.dat', fulldata=all_ic

wave=adas_vector(low=all_ca.wave_min[1], high=all_ca.wave_max[1], $
                  num=all_ca.npix[1], /linear)

plot, wave, all_ca.fpec[*, 7, 2, 1] > 1e-14,           $
  xtitle = 'Wavelength (A)',                         $
  ytitle = 'f-PEC (ph cm!u3!n s!u-1!n)',          $
  xrange = [40, 100], yrange = [1e-12, 4e-11]

oplot, wave, all_ic.fpec[*, 7, 2, 1] > 1e-14, color=5
```

Configuration average (CA) vs intermediate coupling (ic)



In Reality

- ▶ Do not consider each stage by hand — by element is preferable.
- ▶ Many scripts available within ADAS to aid this task.
- ▶ These impose a directory structure.
- ▶ Baseline data identified by the year '40' tag.
- ▶ Full instructions in forthcoming ADAS technical report.

Ionisation

Very similar to excitation — driven by *adf56* collection of rules

<i>index[]</i>	: index of ground configuration of each ion of element in <i>adf56</i> file
<i>config[]</i>	: ground configuration for each ion of element
<i>n_el[]</i>	: number of electrons for each ion of element
<i>no_v_shl[]</i>	: number of shells to treat as valence shells. Max. 2 relevant to relating ion and parent.
<i>v1_shl[]</i>	: first valence shell position in <i>adf56</i> configuration specifications.
<i>v2_shl[]</i>	: second valence shell position in <i>adf56</i> configuration specifications. zero if none defined.
<i>drct_eval_v[]</i>	: evaluate direct ionisation from the valence shell(s).
<i>drct_eval_cl[]</i>	: evaluate direct ionisation from other non-valence (closed) shells.
<i>min_shl_cl[]</i>	: lowest closed shell to include (position in <i>adf56</i> configuration specifications).
<i>exca_eval_v2[]</i>	: evaluate excitation/autoionisation from second valence shell if identified.
<i>max_dn_v2[]</i>	: maximum change in v2 n-shell to be included.
<i>min_dn_v2[]</i>	: minimum change in v2 n-shell to be include.
<i>max_dl_v2[]</i>	: maximum change in v2 l-shell to be included.
<i>min_dl_v2[]</i>	: minimum change in v2 l-shell to be include.
<i>exca_eval_cl[]</i>	: evaluate excitation/autoionisation from other non-valence (closed) shells.
<i>max_dn_cl[]</i>	: maximum change in closed n-shell to be included.
<i>min_dn_cl[]</i>	: minimum change in closed n-shell to be included.
<i>max_dl_cl[]</i>	: maximum change in closed l-shell to be included.
<i>min_dl_cl[]</i>	: minimum change in closed l-shell to be included.
<i>exst_eval[]</i>	: evaluate ionisation from excited states.
<i>exst_adf00_prt[]</i>	: assume parent for building excited states is as present in the <i>adf00</i> data set for the ion.
<i>exst_prt_hole_shl[]</i>	: specify position of shell in ground configuration to form parent if not from <i>adf00</i> above.
<i>max_n_exst[]</i>	: maximum n-shell for excited states to be included.
<i>max_l_exst[]</i>	: maximum l-shell for excited states to be included.
<i>drct_eval_exst_v[]</i>	: evaluate direct ionisation from excited state valence shells.
<i>drct_eval_exst_cl[]</i>	: evaluate direct ionisation from excited state non-valence (closed) shells.
<i>exca_eval_exst_v[]</i>	: evaluate excitation/autoionisation for excited states from valence shells (v1 and v2 above).
<i>exca_eval_exst_cl[]</i>	: evaluate excitation/autoionisation for excited states from non-valence (closed) shells.

adf32 is the driver file for CADW ionisation code from the Auburn group.

Once again to the IDL command line!

```
; Add offline-ADAS IDL library to the path  
  
!path = expand_path('/u/adas/offline_adas/adas8#2/idl') + ':' + !path  
  
; Promotion rules - compiled by Adam Foster (arf)  
  
a56file = '/u/adas/adas/adf56/large_arf09.dat'  
  
; Sn13+ !!  
  
adas8xx_ionis_promotion_rules, z_nuc      = 50, $  
                                z_ion      = 13, $  
                                a56file   = a56file, $  
                               adf32     = 'adf32_ca_sn13.dat', $  
                                comments  = [ 'C-----', '$  
                                'C I made this!', '$  
                                'C-----', '$]  
                                ]
```

```

elem  = Sn
stage = 13
ip_z  = 3193147.3
ip_z1 = 2415629.2
seq   = rb
-----
Type = Direct /number=3/
#
200-51 1 2 01. 1. 5.0E-08 1.0E-11-2 0130 0 1.00 0.65 71. 0.5 0.70
  50 14 sn+13 ground 4d1
  50 15 sn+14 from 4d 3d10 4s2 4p6
  -1
#
200-51 1 2 01. 1. 5.0E-08 1.0E-11-2 0130 0 1.00 0.65 73. 0.5 0.70
  50 14 sn+13 ground 4d1
  50 15 sn+14 from 4s 3d10 4s1 4p6 4d1
  -1
#
200-51 1 2 01. 1. 5.0E-08 1.0E-11-2 0130 0 1.00 0.65 72. 0.5 0.70
  50 14 sn+13 ground 4d1
  50 15 sn+14 from 4p 3d10 4s2 4p5 4d1
  -1
-----
Type = InDirect /number=2/
#
20 -51 0 2 10 1.0 5.e-08 1.e-11-2 130 1.0 0.65 66. 0.5 0.7
  50 14 sn+13 ground 4d1
  50 14 sn+13 via 4d 3d10 4s1 4p6 4d2
  50 14 sn+13 via 4f 3d10 4s1 4p6 4d1 4f1
  50 14 sn+13 via 5s 3d10 4s1 4p6 4d1 5s1
  -
  -
  50 14 sn+13 via 7h 3d10 4s1 4p6 4d1 7h1
  50 14 sn+13 via 7i 3d10 4s1 4p6 4d1 7i1
  -1
#
20 -51 0 2 10 1.0 5.e-08 1.e-11-2 130 1.0 0.65 66. 0.5 0.7
  50 14 sn+13 ground 4d1
  50 14 sn+13 via 4d 3d10 4s2 4p5 4d2
  50 14 sn+13 via 4f 3d10 4s2 4p5 4d1 4f1
  -
  -
  50 14 sn+13 via 7h 3d10 4s2 4p5 4d1 7h1
  50 14 sn+13 via 7i 3d10 4s2 4p5 4d1 7i1
  -1
-----
C-----
C I made this!
C-----

```

Next to the unix command line

```
/u/adas/offline_adas/adas8#2/adas8#2.pl adf32_ca_sn13.dat adf23_ca_sn13
```

Return to IDL to inspect the results

```
read_adf23, file='adf23_ca_sn13.dat', fulldata=all, szd_total=szd  
help, szd,/st
```

```
** Structure <a3e784c>, 7 tags, length=6576, data length=6576, refs=1:  
TE           DOUBLE   Array[12]  
Q_ION        DOUBLE   Array[1, 3, 12]  
IS_Q_ION     LONG     Array[1, 3, 12]  
Q_EXC        DOUBLE   Array[1, 41, 12]  
IS_Q_EXC     LONG     Array[1, 41, 12]  
QTOT         DOUBLE   Array[1, 1, 12]  
IS_QTOT      LONG     Array[1, 1, 12]
```

```

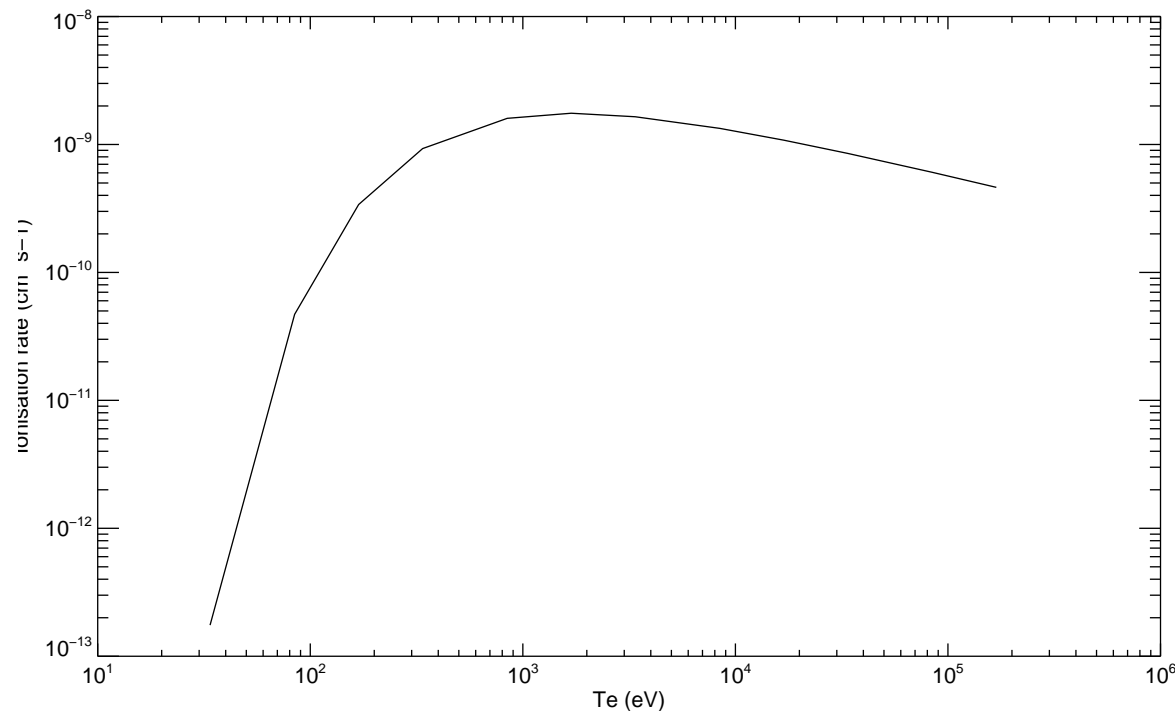
te = reform(szd.te) / 11605.0
szd = reform(szd.qtot*10.0^szd.is_qtot) > 1.0e-36

plot_oo, te, szd, $  

    xtitle='Te (eV)', $  

    ytitle = 'Ionisation rate (cm!u3!n s-1!n)'

```



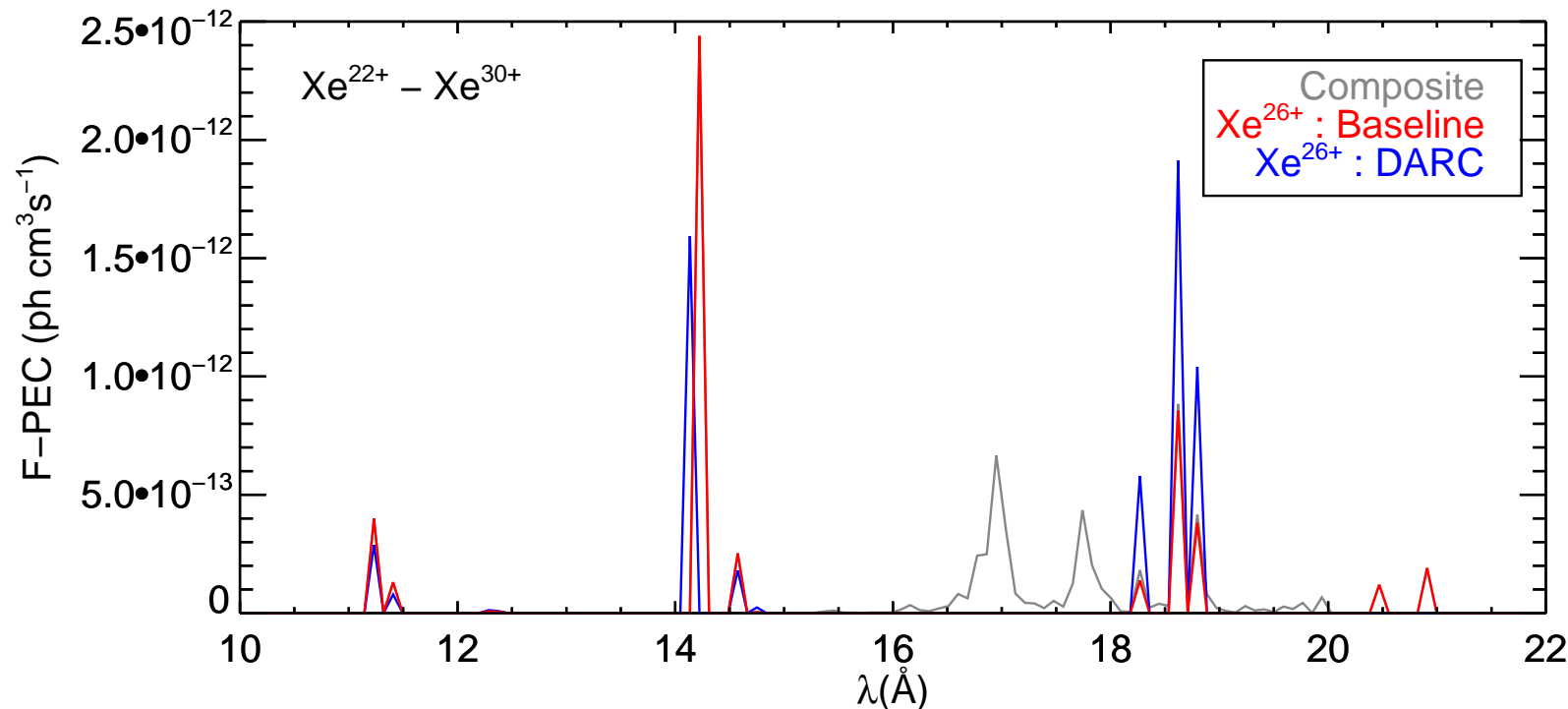
Recombination

- ▶ *adf55* rules are imminent.
- ▶ However, use ADAS407/ADAS408 for now.

Selectively uplift the quality of baseline

- ▶ With increasing atomic number relativistic effects assume a greater importance.
- ▶ Compare the baseline Born data to DARC to assess its validity.

Consider Ni-like Xe^{+26} with a $3d^{10}$ ground configuration:



Handling heavy species data

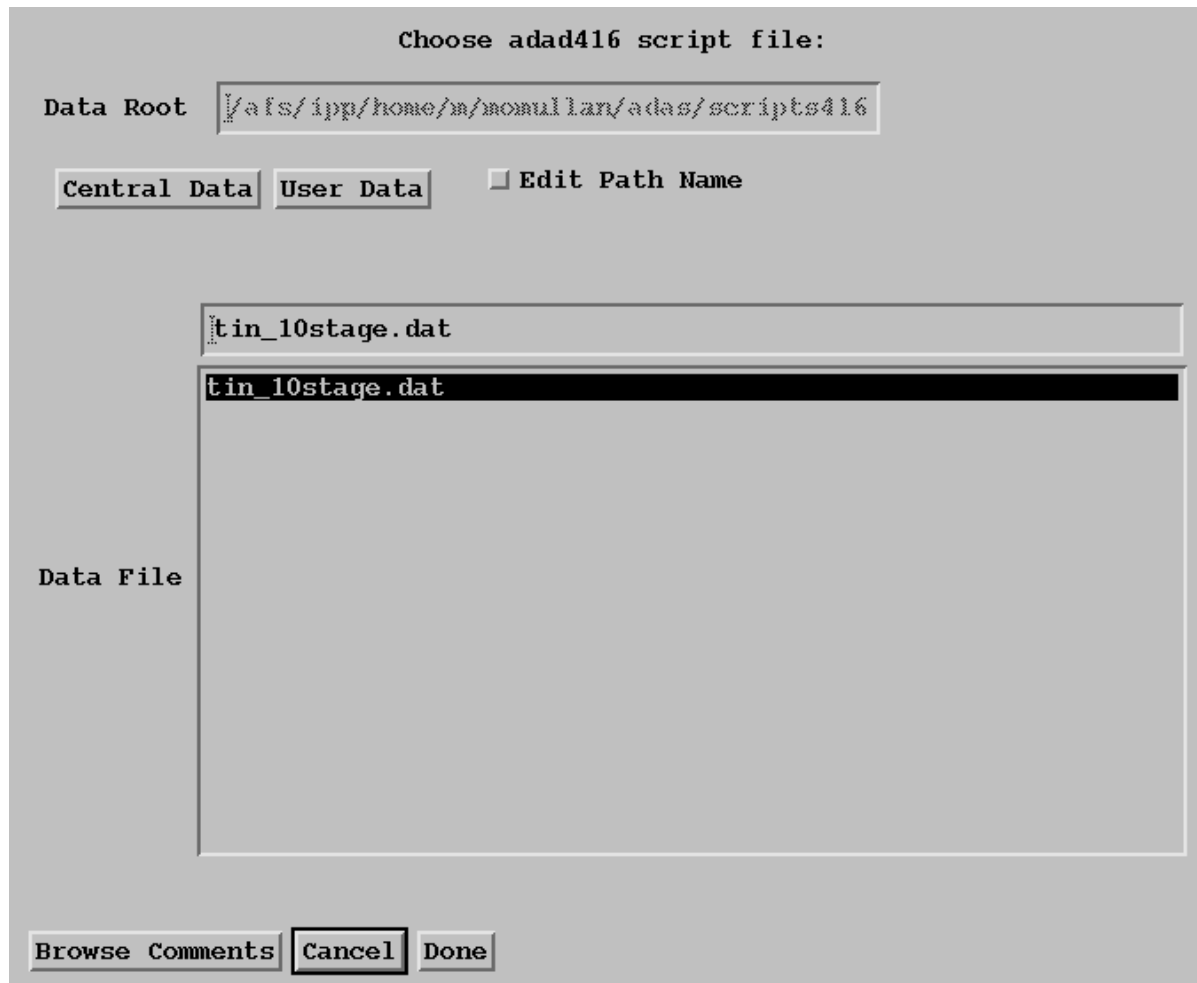
It may not always be necessary to consider all ionisation stages of an element.

Again, for tin, consider the partiton (extract from *scripts416* driver file:

```
//#02/p00/ 00/  
    p01/ 01 02 03 04 05 06 07/  
    p02/ 08/  
    p03/ 09/  
    p04/ 10/  
    p05/ 11/  
    p06/ 12/  
    p07/ 13/  
    p08/ 14/  
    p09/ 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 31 32  
        33 34 35 36 37 38 39 40 41 42 43 44 45 46 47 48 49/  
    p10/ 50/
```

Generate partitioned *adf11* data

Process with ADAS416 : See /u/momullan/adas/scripts416/tin_10stage.dat



Note: *adf11* datasets in /u/momullan/ADAS-EU_course/.

Compare ionisation equilibrium balance

At the IDL command line

```
te = adas_vector(low=1, high=1000, num=40)
dens = fltarr(40) + 1e12

; Explicity name adf11 files

files = {acd : 'acd66_sn#10stage.dat', $
          scd : 'scd66_sn#10stage.dat'  }

run_adas405, uid='adas', elem='sn', year=89, te=te, dens=dens, $
               files=files, frac=frac_par

files = {acd : 'acd89_sn.dat', $
          scd : 'scd89_sn.dat'  }

run_adas405, uid='adas', elem='sn', year=89, te=te, dens=dens, $
               files=files, frac=frac
```

```
xmin = min(te, max=xmax)
ymin = 0.001
ymax = 1.5

plot_oo, [xmin, xmax], [ymin, ymax], /nodata, ystyle=1, $
    xtitle = 'Te (eV)', ytitle = 'Fractional abundance'

for j = 0, n_elements(frac.stage)-1 do begin
    oplot, te, frac.ion[*,j]
endfor

for j = 0, n_elements(frac_par.stage)-1 do begin
    oplot, te, frac_par.ion[*,j], color=5, thick=5
endfor
```

We assume that we have no great interest outside our chosen ions!

