## Interpretation of Infrared (IR) Spectra: a Basic Approach Dr. Laurie S. Starkey, Cal Poly Pomona

"Read" the spectrum from left to right and pick out any of the following obvious bands. Next, look to the weaker bands for additional evidence of the FG's you suspect are present.

| Obvious bands (cm <sup>-1</sup> ) | Functional Group                        | Notes/Additional evidence         |
|-----------------------------------|---|-----------------------------------|
| 3500–3200 (m)                     | O–H N–H                                 | broad signal (water?!)            |
| 3200–3000 (m-s)                   | C–H, sp <sup>2</sup> (aromatic, alkene) | substitution patterns <1000       |
| 3000-2800 (s)                     | C–H, sp <sup>3</sup> (alkane)           | C–H bend at 1460, 1380            |
| 2250–2000 (m-w)                   | C≡N C≡C                                 | often weak                        |
| 1800–1600 (s)                     | C=0                                     | see table, lowered by conjugation |
|                                   |   |                                   |

(s) = strong, (m) = medium, (w) = weak, (v) = variable

## Other peaks to look for:

| ≡с−н   | sp C–H stretch is sharp peak around 3300 (s)  |                                    |                |   |  |
|--|---|------------------------------------|----------------|---|--|
| о<br>—с—н  | aldehyde C-   | -H stretch appears as two          | o sharp peak   | s 2900–2800 and 2800–2700 (w)                                     |  |
| C=C  | stretches oc  | cur 1680–1630 with varia           | able intensity | (v) and <u>will</u> <u>disappear</u> if <u>symmetrical</u>        |  |
| с-о  | stretches occur 1150–1050 and are typically strong (s)  |                                    |                |   |  |
| о<br>—С-он   | Carboxylic acids should be obvious because the H-bonded dimer has a HUGE $O_H$ stretch (3400–2400) that overlaps with the C-H stretch bands. Carboxylic acids will also have a C=O stretch (1725–1700) and a C-O stretch (1320–1210). |                                    |                |   |  |
| O<br>II<br>—C-OR   | Because esters have <u>two</u> C–O bonds, they typically have two C–O stretches.<br>Together with the C=O stretch, it's known as the "rule of 3" (~1700, 1200, 1100).   |                                    |                |   |  |
| alkene C–<br>benzene (   | H bend<br>C–H bend  | 1000–700<br>850–700 (+ overtone 20 | )<br>000–1700) | gives excellent information on substitution patterns (o,m,p, etc) |  |
| C=N is not a common FG and C–N stretches are typically not useful. |   |                                    |                |   |  |

## DO NOT try to interpret every little peak in the spectrum, but DO recognize which ones are <u>significant</u> and are giving you clues to the sample's structure. It takes practice!