Radiofrequency Microwave Double Resonance Experiments in Fourier Transform Technique

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Radiofrequency microwave double resonance has proved as a valuable method in microwave spectroscopy in the frequency domain. We present comparable experiments in the time domain Fourier transform spectroscopy.

Microwave Fourier Transform (MWFT) spectroscopy [1, 2] developed into a very useful method in rotational spectroscopy. The sensitivity and resolution could be improved in comparison with Stark spectroscopy. The double resonance (DR) technique was also introduced [3] and extended in different versions [4–8]. The investigation of Stark effect became also feasible [9–12]. In 1983 Stahl performed radio-frequency microwave double resonance (RFMWDR) experiments which were not published. Similar experiments were also performed by Vogelsanger and Bauder [13]. We now report on three types of experiments:

a) RFMWDR with continuous pump radiation,
b) RFMWDR with modulated pump radiation, and
c) RFMWDR with simultaneous application of a DC-Stark field and modulated RF pump.

RFMW Double Resonance with Continuous RF Pump Radiation

In this experiment we used a MWFT spectrometer described in [12]. The Stark cell has a septum 1 mm thick and one vacuum tight N connector at each end. By one connector the RF is fed into the cell, by the second the RF is monitored after attenuation and DC-blocking with a power meter. In Fig. 1 we present an internal rotation doublet of ethyl fluoride, CH₃CH₂F, Jₖ₋ₖ₊ = 61₄₋₇₂₅. Under the influence of continuous RF pump radiation, each component splits again according to the theory given in [3].

RFMW Double Resonance with Modulated RF Pump Radiation

In this experiment we used the MWFT spectrometer described above but with the RF being on/off modulated. The 40 dB amplitude modulation of the RF source (SMPD, Rhode & Schwarz) proved to be insufficient. Therefore we used PIN modulation (PIN switch, HP 33144A, SPST, 0.1–18 GHz, isolation 60 dB above 1 GHz, or Rohr ES-351, SPDT 1–200 MHz, isolation 60 dB lower for the frequency range) and the technique described in [7]. This experiment was very conclusive in the investigation of the bromine nuclear quadrupole coupling of cyclopropyl bromide, C₃H₃Br. This molecule was first investigated by Lam and Dailey [14]. By these authors the quadrupole coupling constants were determined using an approximative model adequate to the lower resolving power of the Stark spectrometer used. The problem for this nearly symmetric top molecule was, that the transitions 2₀₂₁₋₁₁₀₁, 2₁₁₁₋₁₁₀₀, and 2₁₂₋₁₁₁₂ of the ⁸¹Br and ⁹¹Br isotopomers overlap severely in their hfs-patterns. Other, yet unassigned lines (probably vibrational satellites and high-J-transitions) complicate the spectrum even more. By experiments of the type given in Fig. 2 we were able to get an unambiguous assignment of most of the hyperfine components. The analysis is still in progress, the results will be published.

RFMW Double Resonance with Modulated RF Pump and DC Stark Field

In Fig. 3 we demonstrate the Stark effect with a DC field of approximately 100 V/cm of the rotational transition

Fig. 1. a) Transition J₋K = 61₄₋₇₂₅ of ethyl fluoride, CH₃CH₂F, showing internal rotation splitting. Polarizing frequency 9938 MHz, 20 ns sampling interval, 0.32–10⁶ experiment cycles, pressure 2.3 mTorr, temperature 25 °C, 1024 data points. Zero filling to 4096 points prior to Fourier transformation. — b) Same transition with continuous radio-frequency pump of 62.45 MHz and 1 mW power at the end of the cell pumping the J₋K = 6₃₋₄₃₄ transition. 10⁷ experiment cycles. The cell has an attenuation of less than 3 dB at this frequency. The two double resonance doublets are clearly seen but not completely resolved. The amplitude spectra are approximately to scale.

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Fig. 2. a) A range of 16 MHz of the amplitude spectrum from 10147 to 10163 MHz of cyclopropyl bromide. In this region the hfs patterns of the $J = 2 - 1$ lines overlap. Polarizing frequency 10155 MHz, sampling interval 10 ns, $10^7$ experiment cycles, pressure 2.2 mTorr, temperature $-40^\circ$C, 1024 data points supplemented by 3072 zeros before Fourier transformation. – b) The same range with a modulated RF pump of 116 MHz and 20 mW RF pump power at the cell entrance. Conditions as in Figure 2 a. – the hfs component $F = 5/2 - 5/2$ of the $J_{K-K} = 2_{11} - 1_{10}$ transition of $^{81}$Br remains under pumping the $J_{K-K} = 1_{10} - 1_{11}$, $F = 5/2 - 5/2$ transition. Cell attenuation below 6 dB. – c) The same range with a modulated RF pump of 223 MHz and 20 mW, thus pumping the hfs component $J_{K-K} = 1_{10} - 1_{11}$, $F = 1/2 - 3/2$ of the $^{79}$Br species. Conditions as in Figure 2 a. The remaining line is assigned to the $J_{K-K} = 2_{11} - 1_{10}$, $F = 3/2 - 1/2$ hfs component. Cell attenuation below 9 dB. Intensity scale of b) and c) corrected for modulation loss (factor of 2).

Fig. 3. a) Transition $J_{K-K} = 2_{02} - 1_{11}$ of ethyl fluoride, CH$_3$CH$_2$F with a DC Stark field of approximately 100 V/cm. The $M$ quantum numbers are given. Internal rotation splitting of 84 kHz not resolved. Polarizing frequency 8388 MHz, sampling interval 10 ns, $10^6$ experiment cycles. Pressure 5 mTorr, temperature 25°C. – b) Same range with the same DC Stark field. The transition $J_{K-K} = 1_{10} - 1_{11}$, $M = 1 - 1$ was pumped with modulated RF pump of 1164 MHz. Pump power 10 mW. The $M = \pm 1$ component is relatively enlarged by the double resonance. The intensity scale was corrected for the loss by modulation (factor of 2).

Superimposition of a modulated RF pump radiation via a bias-T (Microlab/FXR HW-15N) results in a remarkable change of the intensity ratio, since the $J_{K-K} = 1_{10} - 1_{11}$, $M = 0 - 0$ pump transition is forbidden, while the $M = 1 - 1$ (and, perhaps, $M = 0 - 1$ and $M = 1 - 0$) is not.

Conclusion

As in the case of MMMW double resonance we show that RFMW double resonance has a great potential in simplifying complicated rotational spectra. Especially the DR modulation technique appears to compensate for the lack of Stark modulation, which up to now hindered assignment of yet unknown spectra.

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[8] B. Vogelsanger, M. Andrist, and A. Bauder to be published.