

THE OPTICAL STARK SPECTRUM OF THE $A^3\Phi_4 - X^3\Phi_4$ BAND SYSTEM OF IRIDIUM MONOFLUORIDE, IrF

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Recently the New Brunswick group^a reported on the field-free detection and analysis of the $A^3\Phi_4 - X^3\Phi_4$ band system of IrF. Here we report on the analysis Q(4)(15922 cm^{-1}) branch feature of the (1,0) band of the ^{191}IrF isotopologue of that system recorded at field strengths of up to 3000 V/cm. The spectra are surprisingly complex at the achieved resolution of 40 MHz due to the presence of both the $^{191}\text{Ir}(I=3/2)$ and $^{191}\text{Ir}(I=1/2)$ magnetic hyperfine splitting. The determined permanent electric dipole moment, μ_{el} , for the $X^3\Phi_4$ state is compared with that recently determined^b for the $X^3\Phi_4$ state of isovalent CoF. The trend in μ_{el} amongst the ground states of IrF, IrC and IrN^c will be discussed. Finally, a simple molecular orbital correlation diagram will be used to rationalize the change in μ_{el} upon excitation from the $X^3\Phi_4$ to $A^3\Phi_4$ state.

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