Table 5. Observed frequencies (cm⁻¹), isotopic shifts, and band assignments for ZnBr₂ · (4,4'-DTDP)

4,4'-DTDP	$^{\rm NA}{\rm ZnBr}_2{\cdot}(4,4'{\text{-}}{\rm DTDP})$	⁶⁴ ZnBr ₂ ·(4,4'-DTDP)	68ZnBr ₂ ·(4,4'-DTDP)	$\tilde{v}(^{64}\mathrm{Zn}) - \tilde{v}(^{68}\mathrm{Zn})$	Assignments
533(s, sp)	550(vw)	547	547	0,	
500(sp)	499(s, sp)	498	497	1)	Ligand and ligand induced
488(s, sp)	486(s, sp)	484	484	0	
438(m)	446(m)	445	445	0	
414(m)	409(vw)	409	409	0 /	
379(w)	389(vvw)	386	386	0	
343(vw)	363(vw)	363	363	0	
281(vvw)	•	284	284	0)	
	264(s)	267	261	6	ν ZnBr
	• •	261(sh)	201	. 0	VZIIDI
	233(vvw)	233	233	0	Ligand induced
	212(s)	212	210	2	vZnN
183(vvw)	192(vvw)	192	192	0	Ligand
	167(m)	168	166	2	Liganu
	103(w, br)	104	104	0	Lattice
	92(w, br)	92	92	0	33000100

Abbreviations: s = strong; sp = sharp; m = medium; w = weak; v = very; sh = shoulder; br = broad.

The observed i.r. frequencies for the NAZnI₂(4,4'-DTDP) complex from 650-80 cm⁻¹ are 548(vvw), 498(s, sp), 485(s, sp) 446(m), 408(vvw), 385(vvw), 362(vvw), 282(vvw), 261(vvw), 243(m), 223(m), 204(m), 173(m), 163(w), 104(vvw). No band assignments were made for this compound since no isotopic studies were conducted for it.

High-pressure studies of the 341 and 299 cm⁻¹ bands in the chloride indicate that the 299 cm⁻¹ band is more intensity sensitive to pressure than the 341 cm⁻¹ band, and is the most intense of the Raman bands, and therefore, may be assigned as the $\nu_{\rm ZnCl_{sym}}$ mode. The mode at 113 cm⁻¹ in the chloride shifts to lower frequency in the bromide and may be assigned as a lattice mode. Other assignments are listed in Tables 3 and 4.

B. Raman studies. Raman scattering spectra of the 4,4'-DTDP complexes, obtained from pellets of powders, showed a lack of coincidence with infrared frequency position indicating a possible centrosymmetric configuration existing for the complexes (see Table 3).

III. Stereochemistry of complexes

The spectroscopic results indicate a high degree of asymmetry in the 2,2'-DTDP complexes. The local environment around the central zinc atom is tetrahedral, as indicated by the position of the zinc-halogen stretching vibrations. Only nitrogen atoms of the ligand are involved in the bonding to zinc. Since one ligand molecule is coordinated per molecule of zinc halide, the ligand could be bidentate (2 nitrogen atoms of a ligand molecule linked to a zinc atom), but this would involve a strained, seven-membered ring which is unlikely. Additionally, such a structure is monomeric and would confer a higher degree of solubility on the complex than is observed. A twisted, polymeric structure of bridging 2,2'-DTDP molecules, held together by zinc halide molecules, is plausible.

The 4,4'-DTDP complexes appear to demonstrate a lack of coincidences between i.r. and Raman frequencies, and thus the complexes are judged to possess a higher degree of symmetry than the 2,2'-DTDP complexes. The local symmetry is tetrahedral and is determined from the far i.r. data. The stoichiometry involves one ligand molecule per molecule of zinc halide, and a linear, polymeric structure involving bridged 4,4'-DTDP molecules, held together by zinc halide molecules, is plausible.