the compound containing the metal in natural abundance to determine if the complex had formed.

B. Materials

The natural abundance zinc halide hydrates used were C.P. grade. The isotopes were obtained from Oak Ridge National Laboratory, Oak Ridge, Tennessee, in the form of oxide. The ligands were obtained from Aldrich Chemical Co., Inc., Milwaukee, Wisconsin.

C. Analyses

The analyses for carbon, nitrogen, and hydrogen were made at Argonne National Laboratory, using micro-analytical techniques. The elemental analysis follow: Anal. Calc. for $(\text{ZnCl}_2 \cdot 2, 2' - \text{DTDP})$ or $(\text{ZnCl}_2 \cdot 4, 4' - \text{DTDP})$: C, 33.68%; N, 7.86%; H, 2.26%; S, 17.96%. Found for $(\text{ZnCl}_2 \cdot 2, 2' - \text{DTDP})$: C, 33.60%; N, 7.81%; H, 2,18%; Found for $(\text{ZnCl}_2 \cdot 4, 4' - \text{DTDP})$: C, 33.85%; N, 7.83%; H, 2.24%; Calc. for $(\text{ZnBr}_2 \cdot 2, 2' - \text{DTDP})$ or $(\text{ZnBr}_2 \cdot 4, 4' - \text{DTDP})$: C, 26.95%; N, 6.29%; H, 1.80%; S, 14.38%; Found for $(\text{ZnBr}_2 \cdot 4, 4' - \text{DTDP})$: C, 26.89%; N, 6.26%; H, 1.82%; S, 14.23%; Found for $(\text{ZnBr}_2 \cdot 4, 4' - \text{DTDP})$: C, 27.22%; N, 6.33%; H, 1.85%; S, 14.44%. Calc. for $(\text{ZnI}_2 \cdot 2, 2' - \text{DTDP})$ or $(\text{ZnI}_2 \cdot 4, 4' - \text{DTDP})$: C, 22.26%; N, 5.19%; H, 1.48%. Found for $(\text{ZnI}_2 \cdot 2, 2' - \text{DTDP})$: C, 22.10%; N, 4.97%; H, 1.46%; Found for $\text{ZnI}_2 \cdot 4, 4' - \text{DTDP}$: C, 22.48%; N, 4.92%; H, 1.54%.

D. Infrared and Raman measurements

Infrared measurements from 4000–650 cm⁻¹ were made with KBr disks of the solids using a Beckman IR-12. Measurements in the region from 650–80 cm⁻¹ were obtained with a Beckman IR-11 or a Perkin–Elmer Model No. 301, using polyethylene disks. High-pressure measurements in the far i.r. (up to ~ 24 kbar), were obtained with an opposed diamond-anvil cell using the Model 301 equipped with a $6 \times$ beam condenser. The techniques used and the method of pressure calibration have been previously reported [23, 24]. The Raman spectra were obtained on the powdered solids using a Cary 81 spectrophotometer with a helium–neon laser.

Results and Discussion

I. Complexes with 2,2'-DTDP

A. Infrared studies. The mid-i.r. spectra from $4000-650 \text{ cm}^{-1}$ confirmed the analytical results showing no water absorptions in the complexes. The carbonnitrogen ring vibration at about ~1570 cm⁻¹ in 2,2'-DTDP shifted toward higher frequencies in the complexes, and the results were indicative of bonding occurring to the nitrogen atom of the ligand [25, 26]. Little change occurred in the C—S stretching

^[23] J. R. FERRARO, S. S. MITRA and C. POSTMUS, Inorg. Nucl. Chem. Letters 2, 269 (1966); 4, 55 (1968).

^[24] L. J. BASILE, C. POSTMUS and J. R. FERRARO, Spectry Letters 1, 189 (1968).

^[25] J. R. FERRARO, J. ZIPPER and W. WOZNIAK, Appl. Spectry 23, 160 (1969).

^[26] J. R. FERRARO and K. C. DAVIS, Inorg. Chim. Acta 3, 685 (1969).

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vibration region at $700-800 \text{ cm}^{-1}$ from the uncomplexed ligand, indicating that no bonding occurred to the sulfur atom [27].

Tables 1 and 2 tabulate the low-frequency absorptions for 2,2'-DTDP, ^{NA}ZnCl₂·(2,2'-DTDP), ^{NA}ZnBr₂·(2,2'-DTDP), and for the zinc complexes containing the zinc isotopes of mass 64 and 68. Figure 1 depicts the spectra of isotopic zinc halide complexes from 325-100 cm⁻¹.

2,2'-DTDP	$^{\texttt{NA}}ZnCl_2\textbf{\cdot}(2,2'\text{-}DTDP)$	$^{64}\mathrm{ZnCl}_2{\cdot}(2,2'{\text{-}\mathrm{DTDP}})$	68 ZnCl ₂ ·(2,2'-DTDP)	$\tilde{v}(^{64}\mathrm{Zn}) - \tilde{v}(^{68}\mathrm{Zn})$	Assignments	
	648(m, sp)	648	648	0	A COLORADOR OF	
	648(m, sp)	641	641	0)		
622(s, p)				o non territo non		
	499(m, sp)	500	501	-1	Ligand and ligand	
	487(m, sp)	488	487	1)	induced	
471(m, sp)				and the second second second		
429(m, sp)	429(m, sp)	429	429	0		
	417(s, sp)	417	417	0/		
402(sh)						
345(s, sp)	345(w)					
	321(vs)	322	321	1} 3	vZn-Clasym + ligand	
	293(vs)	294	291	3	vZn-Clsym	
254(vvw)						
	242(m)	242	242	07	Times	
	231(m)	231	231	0}	Ligand	
	222(m)	224	220	4	vZn—N	
158(w)	158(m)		to a detter	-)	Linned Str. Cland	
	130(s), 121(sh)	130	129	1}	Ligand, δZn —Cl and lattice vibrations	
	108(m)	108	108	0)	lattice vibrations	

Table 1. Observed frequencies (cm⁻¹), isotopic shifts, and band assignments for $ZnCl_{o} \cdot (2,2'-DTDP)$

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

2,2'-DTDP ^{NA} ZnBr ₂ ·(2,2'-DTDP)* ⁶⁴ ZnBr ₂ ·(2,2'-DTDP)			⁶⁸ ZnBr ₂ ·(2,2'-DTDP)	$\tilde{\nu}(^{64}\mathrm{Zn}) - \tilde{\nu}(^{68}\mathrm{Zn})$	Assignments	
- point	646(m, sp) 639(m, sp)	646	646	0)	i a leas stille - o	
622(s, sp)	000(m, 5p)			and the second second		
	499(m, sp)	501	500	1		
	486(m, sp)	488	488	0	Times days d Room d	
471(m, sp)				>	Ligand and ligand	
129(m, sp)	429(m, sp)	429	429	0	induced	
	417(s, sp)	418	417	1		
102(sh),						
845(s, sp)				ALT THE R. DOCK M. L. C. DOCK		
	320(m)	320	320	0)		
254(vvw)						
	247(vs)	248	244	4	vZnBrasym	
	223(s)	226	221	5	vZn—N	
	200(s)	201	197	4	vZn—Br _{sym}	
158(w)	152(vw)	-		-)		
	133(vw), 120(vw)	133	133	0	Ligand and lattice	
	113(vw)	113(vw) —			vibrations	
	98(m)	100	99	1)		

Table 2.	Observed	frequencies	$(cm^{-1}),$	isotopic shifts,	and band	l assignments fo	or ZnBr ₂	\cdot (2,2'-DTDP)
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Abbreviations: s = strong; sp = sharp; m = medium; w = weak; v = very; sh = shoulder. * The observed i.r. frequencies for the ^{NA}ZnI₂(2,2'-DTDP) complex from 650-80 cm⁻¹ are 648(m), 528(vvw), 487(s, sp), 438(w), 433(w), 421(s, sp), 417(s, sp), 348(w), 314(s), 240(m), 231(m), 213(m), 194(s), 185(s), 162(m), 140(vw), 115(vvw), 100(vvw), 84(m). No band assignments were made for this compound since no isotopic studies were conducted for it.

[27] P. C. H. MITCHELL and R. J. P. WILLIAMS, J. Chem. Soc. 1912 (1960).