Vibrational analysis of different crystalline phases of the organic electroluminescent material tris(8-hydroxyquinolinato)aluminum(III) (Alq3)

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ABSTRACT

Applying infrared (IR) spectroscopy on two different crystalline phases of tris(8hydroxyquinolinato)aluminum(III) (α -Alq₃ and δ -Alq₃) the isomerism of the Alq₃ molecule was investigated. Significant differences between the two phases were found in the IR spectra, which can be explained in terms of the different symmetries of the facial and meridional isomer. Additionally, intermolecular interactions of the Alq₃-molecules due to crystallinity have been taken into account. The results prove that α -Alq₃ consists of the meridional isomer while the recently discovered blue luminescent δ -phase is composed of the facial isomer of the Alq₃ molecule.

Introduction

After more than 10 years of intense research and development of organic light-emitting diodes (OLEDs), Alq₃ still continues to be the workhorse among the class of low-molecular weight materials for OLEDs.^{1,2} However, a still unresolved issue concerns the isomery of the Alq₃ molecule.^{3,4} It is well-known that octahedral complexes of the type MN₃O₃, where M is a trivalent metal and N and O stand for the nitrogen and oxygen atoms in the hydroxyquinoline ligands, can occur in two different geometrical isomers: meridional (*mer*) and facial (*fac*). Nevertheless only the meridional isomer has been clearly identified so far. Therefore, it is generally believed that the meridional isomer is predominant, both in amorphous films and crystals of Alq₃.^{5,6}

The existence and the properties of the facial isomer are intensely discussed in the literature.^{3,6-12} A key issue concerning sublimed Alq₃ films is the possible presence of the facial isomer, which is expected to cause structural defects in the film and to be an efficient electron trap. Its influence on trap density, charge carrier transport and thus on the characteristics and performance of OLEDs has been suggested,¹³⁻¹⁵ but until recently no direct experimental evidence for the facial isomer has been found. Thus the isolation of the facial isomer is of high interest, as it allows to examine its properties separately and to clarify its role in OLEDs.

In a previous publication we have reported a new crystalline phase of Alq₃ (δ -Alq₃), exhibiting strongly blue-shifted fluorescence.^{16,19} Structural investigations have proved that it consists of the facial isomer of the Alq₃ molecule.²¹ In this paper we present a comparison of the infrared (IR) spectra of the well-known α -phase of Alq₃ being known to contain the meridional isomer to the new δ -phase and demonstrate that their vibrational properties are consistent with the isomery of the Alq₃ molecule.

Due to the different molecular symmetry of the meridional and facial isomers (C1 versus C3) the

vibrational analysis using IR spectroscopy should be one possible way to differentiate them. Especially the central part of the molecule AlO₃N₃ should show characteristic vibrational properties for each isomer (Al-O and Al-N modes located below 600cm⁻¹), which were calculated by Kushto et al.⁶ Further, there is a weak coupling of the three ligands via the central part and motions around the central aluminum atom participate in most of the molecular vibrations below 1700cm⁻¹. This coupling depends on the relative positions of the oxygen atoms of the ligands. For the facial isomer each oxygen atom faces a nitrogen atom thus the coupling via the Al atom is identical for all ligands (see Figure 1). For the meridional isomer, however, the coupling mainly affects the ligand B and C, where the oxygen atoms face each other and to a lesser extend the A and B ligands, which have the oxygen and nitrogen atoms opposite. The coupling mechanism of ligand A and C is mainly characterized by the modes of the two opposite nitrogen atoms. This means, that due to a lower symmetry of the meridional molecule each vibrational mode has a slightly different energy for the three ligands.

Degli Esposti et al. published a detailed theoretical study of the vibrational properties of the meridional Alq₃-molecule.¹⁷ For the first time the coupling terms between the three ligands have explicitely been taken into account. This interaction of the ligands, mainly via the Al-bonds is expected to be different for the two isomers. An important result is a well-founded understanding of the IR spectra of *mer*-Alq₃ with the possibility to attribute lines to the three different ligands A, B and C. On the base of this detailed theoretical work we will first discuss our experimental results on polycrystalline Alq₃ for the entire spectrum up to 1700cm⁻¹. Thereafter we will focus on the stretching modes around the aluminum atom (below 600cm⁻¹) before investigating the influence of crystallinity on the vibrational spectra.

Experimental

Polycrystalline samples of the α - and δ -phase of Alg₃ were fabricated in a temperature gradient sublimation apparatus as described elsewhere.¹⁶ Thereby we obtained the usual yellowish-green α -Alg₃ as the main phase together with small amounts of δ -Alg₃ growing in the hot zone of the sublimation tube. We note that δ -Alq₃ can also be obtained by annealing α -Alq₃ above 385°C.^{18,19} Measurements on samples fabricated by this annealing process gave the same spectra and led to the same results. The phases have been clearly identified by x-ray powder diffraction measurements.^{16,19,20} If not marked otherwise, polycrystalline samples as obtained from the sublimation tube have been measured. Powder samples were made by smoothly rubbing the polycrystalline samples mechanically. The Alq3-samples were measured on KBr pellets in a Bruker IFS55 Fourier transform infrared (FT-IR) spectrometer with KBr beamsplitters and a resolution of 2cm⁻¹. Additionally, to prove the interpretation below 420cm⁻¹, a Bruker IFS 66v spectrometer with PE-beamsplitters, CsI pellets and a resolution of 1cm⁻¹ was used for the low energy region between 370 and 480cm⁻¹. We note that all spectra shown here are raw data, without any further correction (smoothing, background, etc.). In the case of the polycrystalline samples scattering due to crystallinity is responsible for the broad background with sometimes asymmetric peaks. As will be seen from the comparison to the powder measurements this does not significantly influence the peak position.

Results and Discussion

The IR spectra of the polycrystalline α - and δ -phases of Alq₃ are shown in Figure 2 together with the spectra of the free-base ligand 8-hydroxyquinoline (8-Hq). 8-Hq shows intense lines above 700cm⁻¹. Thus it is obvious that for Alq₃ the region above 700cm⁻¹ is dominated by the ligand vibrations,

whereas below 600cm⁻¹ the central part around the Al-atom becomes more important. The spectra of the α - and δ -phases of Alq₃ show significant differences which will be discussed below following theoretical calculations by Degli Esposti et al.¹⁷ and Kushto et al.⁶ Table 1 summarizes the band positions and their relative intensities together with their preliminary assignments according to References 6 and 17 for the *mer*-Alq₃ molecule, which is the constituent of α -Alq₃.^{5,6} For the δ -phase of Alq₃ we know from structural investigations that it consists of the facial isomer. We can therefore use the IR spectra to identify characteristic differences in their vibrational properties.

As can be seen in Figure 2 most of the bands in δ -Alq₃ have slightly reduced line width. Furthermore, broad featureless bands in α-Alq₃ become narrower and exhibit pronounced structures in δ -Alq₃, in particular some adjacent bands in α -Alq₃ become one single peak in δ -Alq₃. Examples are found at wavenumbers 1470cm⁻¹, 1380cm⁻¹, 1333cm⁻¹, 864cm⁻¹, 750cm⁻¹, and 638cm⁻¹. According to Degli Esposti et al. the broad band of α -Alq₃ at 645cm⁻¹ consisting of peaks at 652cm⁻¹ ¹, 647cm⁻¹, 640cm⁻¹ and the small shoulder at about 625cm⁻¹ are due to the pyramidalization mode of the nitrogen atoms within the different ligands A, B and C under participation of Al-O stretching. Whereas the modes at 652cm⁻¹ to 640cm⁻¹ are mainly due to torsional and pyramidalization modes of ligand C, the shoulder at 625cm⁻¹ originates mainly from ligand A indicating the inequivalence among the ligands in mer-Alq3. In fac-Alq3 the ligands are equivalent and therefore the low frequency shoulder should not be observed in δ -Alq₃. We also note that the whole band is narrower for the δ -phase with a full width at half maximum (FWHM) of 28cm⁻¹ and 19cm⁻¹ for α - and δ -Alg₃, respectively. For α -Alg₃, the band centered at 747cm⁻¹ consists of two overlapping bands, one involving C-CABC stretching under participation of Al-O stretching on the low frequency side and a second one involving C-C^{ABC} deformations on the high frequency side. Each of the bands consists of contributions from ligands A, B and C. Taking all together A-vibrations are found at the flanks and C-vibrations in the center of the whole band. Contributions from B are in between the

contributions of A and C. If one also takes into account a possible splitting due to the crystal field it is clear why this band is so broad. In contrast for δ -Alq₃ the FWHM again is smaller (α : 40cm⁻¹, δ : 17cm⁻¹) and two peaks are clearly resolved. As for *fac*-Alq₃ the ligands are equivalent and only one peak is expected; therefore the observed splitting can be attributed to a crystalline effect (this will be discussed in more detail later on).

Weak but interesting bands are observed at 867cm^{-1} and 859cm^{-1} for α -Alq₃ and assigned to torsional and pyramidalization modes of the different ligands. The peak at 867cm^{-1} is due to vibrations of ligand B, while the peak at 859cm^{-1} originates from vibrations of ligands A and C. For δ -Alq₃, however, instead of this splitting only one single peak at 864cm^{-1} is observed. The difference between the ligands seems indeed to be repealed in δ -Alq₃, as expected for the facial isomer. The superposition of two peaks at 1332cm^{-1} and 1327cm^{-1} results in a broad band for α -Alq₃. This band is a result of CCH^{ABC} bending modes of the different ligands and C-O^{ABC} stretching modes. Theoretically the C-O stretching bands of ligand A and C are separated by more than 6cm^{-1} . The C-O stretching mode of ligand A is located at the high frequency side and the C-O stretching mode of ligand C at the low frequency side of this band. Consequently a broad band is observed in the IR-spectrum. Instead of this broad feature there is only one peak for δ -Alq₃ located at 1333cm⁻¹. This is another clear hint for the equivalence of vibrational modes in the three ligands of the facial isomer in the δ -phase.

For the doubled band observed at about 1380cm^{-1} in α -Alq₃ splittings of 2-3 cm⁻¹ are predicted due to C-C^{ABC} and C-N^{ABC} stretch vibrations in the different ligands which are not resolvable. The clear doublets in both crystalline phases (α and δ) are thus attributed again to a crystalline effect as will be discussed later. Another intense and broad feature is observed for α -Alq₃ at 1468 cm⁻¹, being the superposition of C-O^{ABC} and C-C^{ABC} stretching vibrations in the ligands A, B and C of the meridional molecule. Theoretical energetic distances between these contributions are 4cm^{-1} and 8cm^{-1} ¹ resulting in one broad band. For δ -Alq₃ again only one peak is expected and observed at 1473cm⁻¹. The weak low frequency shoulder in both phases is assigned to C-H bending.

Kushto et al. calculated the IR-spectra of the facial and the meridional isomer.⁶ Characteristic differences were predicted at about 400cm⁻¹ and 550cm⁻¹. Theoretically, a much more intense peak at 406cm⁻¹ is expected for the *fac*-Alq₃ compared to *mer*-Alq₃. Indeed this is observed experimentally for the band at 397cm⁻¹. Further they found the characteristic feature of two neighbored peaks at 550cm⁻¹ for the meridional molecule and only one peak for the facial molecule to be most significant to distinguish between the two isomers. As shown in Figure 2 one observes exactly this characteristic difference between α - and δ -Alq₃.

Further arguments for different isomers in α - and δ -Alq come from symmetry considerations of the central fragment around the Al-atom. As discussed above the differences in the IR spectra of the two phases, in particular the higher number of IR-active vibrational modes for the α -phase in comparison to the δ -phase of Alq₃, can be ascribed to the different symmetry of the constituting meridional and facial isomers, namely C₁ and C₃, respectively. Focusing on the central fragment AlO₃N₃, the local symmetry for both isomers is C_{2v} and C_{3v}, respectively, as shown in Figure 3. The separation of the central part from the ligands is justified by the different and well separated vibrational energies belonging to these groups, as observed in the comparison of Alq₃ with the hydroxyquinoline parent of the ligands (see Ref. 22). Thus IR-modes that belong to the AlO₃N₃ group are found predominantly in the region below 600cm⁻¹. In particular we observe in our experiment the stretching vibrations. For α -Alq₃, which consists of the *mer*-Alq₃ isomer (C_{2v}) six stretching vibrations are expected, three involving Al-N and three involving Al-O modes (see figure 3). As they are all dipole allowed, they are observable by IR-spectroscopy. According to Ref. 6 we make the following assignments for α -Alq₃: Al-N stretching: 396cm⁻¹, 418cm⁻¹, Al-O

stretching: 522cm⁻¹, 542cm⁻¹, 549cm⁻¹. By contrast δ -Alq₃ shows a total of only four bands in this region (397cm⁻¹, 423cm⁻¹, 531cm⁻¹, 548cm⁻¹). As the AlO₃N₃ fragment of the facial isomer belongs to symmetry C_{3v} also six stretching vibrations are expected but four of them belong to two degenerate vibrational states and therefore only four bands should be observed in IR-spectroscopy as one does for δ -Alq₃. The Al-N stretching is found at 397cm⁻¹ and 423cm⁻¹, the Al-O stretching at 531cm⁻¹ and 548cm⁻¹. The degeneracy of the first and last band are lifted in the α -phase of Alq₃ (see Figure 3 and Figure 2 incl. inset), which consists of the meridional isomer, in agreement with theoretical calculations of Kushto et al.

A further point to consider is the fact, that vibrational studies of the polycrystalline phases of Alq₃ do not only involve the intramolecular characteristics but also the effect of crystallinity of the sample. Therefore, in Figure 4 we compare IR-spectra of polycrystalline samples to very fine powders of the α - and δ -phases. The upper trace shows the polycrystalline samples as prepared and the lower trace the same samples after smoothly rubbing them directly on the KBr pellets, giving a very fine powder. We note that less crystallinity of the samples is also reducing the background due to scattering, especially in the higher energy region above 700cm⁻¹ resulting in symmetric peaks in the measured spectrum. Nevertheless, the energetic positions of the peaks are essentially not affected by the sample crystallinity.

We observe two distinct effects: At first the bands above 600cm^{-1} and therefore the vibrational modes originating mainly from the hydroxyquinoline ligands become sharper and the lines at 750 and 1380cm⁻¹ lose their doublet character after rubbing. This is attributed to the intermolecular loss of geometric correlation. Secondly we note that the Al-N and Al-O stretching vibrations found in the α - and δ -crystals stay, apart from slight shifts, at the same frequency after disintegration of the crystals. The degenerated bands of the facial isomer at 397cm⁻¹ and 548cm⁻¹ still are one single peak for the δ -Alq₃ and split into two distinct peaks for the meridional α -Alq₃ respectively. Thus the

differences between the δ -Alq₃ and the α -Alq₃ are not related to the crystallinity or packing but due to the difference in the symmetry of the molecules.

One significant change between the crystalline and the powder spectra of δ -Alq₃ is the large loss of intensity of the Al-O band at 531cm⁻¹, which corresponds to the redshifted mode at 522cm⁻¹ in α -Alq₃. From theoretical considerations the intensity of the Al-O streching mode for the facial isomer at 531cm⁻¹ is expected to be very small.⁶ This is indeed observed in the spectrum of the very fine powder in Figure 4a. It is, however, well known, that molecular ordering in a crystalline environment can change the relative intensities of related vibronic bands, e.g. due to dipole-dipole interaction.²³⁻²⁵ The strength of this effect depends on the relative orientation of the dipoles, their distance r (~r⁻³), the difference of the squares of their transition energies and their oscillator strength. Unfortunately the absolute values for the oscillator strength can not simply be obtained from the IR-spectra²⁶ and thus it is not possible to make quantitative calculations. But considering the relative positions of the molecules in the crystal known from structural analysis^{5,21} one can roughly estimate that this effect should be more pronounced for the facial isomer in the δ -phase. Thus the band at 531cm⁻¹ can be viewed as a kind of crystallinity band of δ -Alq3.

Further it is clearly observed that the doubling of the band centered at 750cm⁻¹ in δ -Alq₃ disappears after destroying the crystallinity of the sample, confirming the assignment made in the first part of this work. The same holds for the double band observed for both phases at about 1380cm⁻¹. It also becomes a single mode after powdering. We note that although all vibrational peaks of both isomers become narrower after mechanical treatment, essential bands in α -Alq₃ remain broader than in δ -Alq₃, accounting for the inequivalence of the A, B and C ligands in *mer*-Alq₃. Examples are the lines at the region of 640cm⁻¹, 1333cm⁻¹ and 1470cm⁻¹ with FWHM of 17cm⁻¹, 12cm⁻¹, 13cm⁻¹ and 20cm⁻¹ , 15cm⁻¹, 19cm⁻¹ for the δ - and the α -Alq₃ respectively. As the intramolecular geometry is retained after making the fine powder we note that the removal of crystallinity (seen above 600cm⁻¹) permits us to clearly identify the differences in the central AlO₃N₃ part of the Alq₃ molecules below 600cm⁻¹. Therefore it is evident that the meridional molecule constitutes the α -phase and the facial molecule the δ -phase of Alq₃.

Conclusion

In conclusion we presented the first IR-measurements on the blue luminescent δ -phase of Alq₃. From a thorough comparison with theoretical work by Kushto et al.⁶ and Degli Esposti et al.¹⁷, together with an explicit consideration of the symmetry of the central fragment AlO₃N₃ as well as the influence of crystallinity we have convincing evidence that the δ -phase of Alq₃ contains the facial isomer, which is also consistent with structural investigations. Both isomers (*mer* and *fac*-Alq₃) show characteristic differences in their vibrational spectra that can be used like fingerprints to distinguish and to identify them. From these findings it should now be possible to examine the importance of the isomerism of the Alq₃ molecule, e.g. for the growth and structural properties of thin films, and to specify its role in OLEDs.

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TABLE 1: Comparison of the experimental frequencies of the δ -phase (E(δ)), the α -phase (E(α)) and 8-hydroxyquinoline (E_{quin}) together with the assignments according to Ref. 6 and 17. A, B and C refer to the corresponding ligands in *mer*-Alq₃. Theoretical frequencies for the *mer*-Alq₃ are listed under (E_{cale}). In the last column the normal modes of 8-hydroxyquinoline are labeled by the symmetry within the C_s group. Relative intensities for the δ - and the α -phase are given in column I(δ) and I(α). Asterisks mark peaks related to crystallinity.

No.	Ε (δ)	Ι(δ)	Mode	Ε (α)	I(α)	E_{calc}	Mode	E_{quin}	Sym-
	[cm ⁻¹]	[a.u.]		[cm ⁻¹]	[a.u.]	[cm ⁻¹]		[cm ⁻¹]	metry
1	1604	57	CC-str., CCC-bend.	1603	59	1606.5	CC-str. ^A , CC-str. ^B		
						1605.4	CC-str. ^C , CCC-bend. ^C		
2	1594	23	CCC-bend., CC-str.	1594sh	27	1601.7	CCC-bend. ^B , CC-str. ^C		
								1592	9A'
						1583.4	NC-str. ^A , CC-str. ^A	1580	10A'
3	1580	89	NC-str., CC-str.	1578	72	1582.4	NC-str. ^C , CC-str. ^C		
						1575.7	NC-str. ^B , CC-str. ^B		
4	1509	25	11A'					1508	11 A'
						1503.2	CC-str. ^B		
5	1498	183	CC-str., CCH-bend.	1497	150	1500.6	CC-str. ^C , CCH-bend. ^C	1503	
						1499.7	CC-str. ^C , CCH-bend. ^B		
6						1476.1	CO-str. ^A , CC-str. ^A		
7	1473	141	CO-str., CC-str.	1471	96	1468.0	CO-str. ^C , CC-str. ^A	1471	12 A'
8	1463	76	CC-str., CO-str.			1462.7	CC-str. ^B , CO-str. ^B	1471	12 A'
						1426.9	CCH-bend. ^A	1434	13 A'
9	1421	39	CCH-bend.	1421	17	1426.2	CCH-bend. ^C		
						1425.0	CCH-bend. ^B		
								1410	14 A'
10	1403	11							
						1392.3	NC-str. ^B , NCH-bend. ^B		
11	1384	194	NC-str., NCH-bend.	1385	112	1391.4	NC-str. ^C , NCH-bend. ^C		
						1389.8	NC-str. ^A , NCH-bend. ^A		
						1376.7	CC-str. ^A , CC-str. ^B	1381	15 A'
12	1377	143	CC-str.	1375	117	1375.8	CC-str. ^C , CC-str. ^A		
						1372.9	CC-str. ^B , CC-str. ^A	1372s	
						1337.2	CCH-bend. ^A , CO-str. ^A		
13	1333	97	CCH-bend., CO-str.	1327	86	1334.6	CCH-bend. ^B , CCH-bend. ^C	1355	16 A'

						1331.2	CCH-bend. ^C , CO-str. ^C		
								1285	17 A'
						1294.3	NC-str. ^A , CCC-bend. ^A		
14	1282	55	CO-str., NC-str. CCH-bend.	1282	48	1291.7	NC-str. ^B , CO-str. ^B	1276	18 A'
						1290.1	CO-str. ^C , CCH-bend. ^C		
15	1244	25	CCH-bend.	1242	10		CCH-bend.	1244	
						1229.1	NC-str. ^A , NCH-bend. ^A		
16	1230	52	NC-str., NCH-bend.	1227	54	1226.9	NCH-bend. ^C , NC-str. ^C		
						1221.6	NCH-bend. ^B , NC-str. ^B	1223	19A'
						1216.9	CC-str. ^B	1206	20 A'
17	1209	10	CC-str.	1211	16	1216.3	CC-str. ^C		
						1216.2	CC-str. ^A		
								1187	
						1169.3	CCH-bend. ^C , CCH-bend. ^A	1173	
18	1173	20	CCH-bend.	1173	13	1168.6	CCH-bend. ^B	1166	21 A'
						1168.3	CCH-bend. ^A		
						1134.2	CCH-bend. ^C	1140	22 A'
19	1133	10	CCH-bend.	1133	3	1135.5	CCH-bend. ^A		
						1133.1	CCH-bend. ^B		
						1105.5	CCH-bend. ^A , CNC-bend. ^A	1113	
20	1111	143	CCH-bend., CNC-bend.	1112	120	1104.3	CCH-bend. ^C , CNC-bend. ^C	1099	23 A'
						1102.5	CCH-bend. ^B , CNC-bend. ^B		
						1056.1	CC-str. ^A		
21	1060	22	CC-str.	1058	20	1052.0	CC-str. ^C	1060	24 A'
						1050.1	CC-str. ^B		
						1034.8	CC-str. ^A		
22	1036	70	CC-str.	1033	47	1030.3	CC-str. ^C	1029	
						1029.1	CC-str. ^B		
								974	1 A"
								959	2 A"
								950	3 A"
								897	26A"
23				867	12	868.5	CC-tors. ^B , C-pyr. ^B		
24	864	31	CC-tors., C-pyr.	859	19	865.6	CC-tors. ^C , C-pyr. ^C	866	4 A''
						864.8	CC-tors. ^A , C-pyr. ^A		
						826.2	C-pyr. ^B	818	5 A"
25	823	227	C-pyr.	823	197	824.3	C-pyr. ^C		
						820.2	C-pyr. ^A		
						793.2	CCC-bend. ^C		
26	804	168	CCC-bend.	803	152	792.4	CCC-bend. ^B	807	27 A"

						791.7	CCC-bend. ^A		
						787.6	CC-tors. ^B , C-pyr. ^B		
27	788	182	N-pyr., CC-tors. C-pyr.	786	150	786.2	N-pyr. ^C , CC-tors. ^C	781	6 A''
						784.2	C-pyr. ^A , CC-tors. ^A		
						751.8	C-pyr. ^B		
28	753*	250	C-pyr. *	747	250	750.3	C-pyr. ^C		
						749.1	C-pyr. ^A		
						746.0	OAl-str. ^A , CC-str. ^A		
29	747*	239	OAl-str., CC-str. CCC-	747		743.2	OAl-str. ^C , CCC-bend. ^C	741	7 A''
						741.0	OAl-str. ^B , CC-str. ^B	711	28 A"
30	652	98	N-pyr., CC-tors.	652	111	653.4	N-pyr. ^C , CC-tors. ^C		
						649.5	N-pyr. ^C , CC-tors. ^C		
31	645	179	N-pyr., C-pyr.	647	145	648.8	N-pyr. ^C , C-pyr. ^A		
32	638	150	N-pyr.	640	127	646.8	N-pyr. ^B , N-pyr. ^C	637	
33				625sh	60	639.8	N-pyr. ^A , COAl-bend. ^A		
						570.5	CCC-bend. ^C , COAl-bend. ^C		
34	574	56	COAl-bend, CCC-bend.	575	46	568.8	COAl-bend. ^A , CCC-bend. ^A	575	10 A"
						566.9	COAl-bend. ^B , COAl-bend. ^C		
35	548	174	OAl-str.	549	144	540.4	OAl-str. ^B , OAl-str. ^C	547	
36				542	141	535.7	OAl-str. ^A , CCC-bend. ^A	543	30 A''
37	531*	77	OAl-str., CCC-bend.*	522	54	512.0	OAl-str. ^B , CCC-bend. ^B		
						495.6	CCC-bend. ^B , CCC-bend. ^C		
38	501	31	CCC-bend.	503	33	493.0	CCC-bend. ^B , CCC-bend. ^C		
						490.4	CCC-bend. ^A	491	31 A''
						476.0	C-pyr. ^B , NAl-tors. ^C		
39	472	21	NAl-tors., C-pyr., CC-tors.,	471	59	474.6	NAl-tors. ^C , N-pyr. ^C	471	11 A"
						473.5	C-pyr. ^A , CC-tors. ^A		
40	469	23	N-pyr., NAl-tors.	465sh	35	464.3	N-pyr. ^C , NAl-tors. ^C	465	32 A'
41	452	96	N-pyr.	454	145	462.5	N-pyr. ^C , N-pyr. ^A		
42	443	26	N-pyr., NAl-tors.	442	69	450.3	N-pyr. ^C , NAl-tors. ^C		
43	423	260	Al-N-str.	418	206	433	Al-N-str.		
44				405	49	415	Al-N-str.		
45	397	117	Al-N-str., COAl-bend	396	45	407	Al-N-str.		

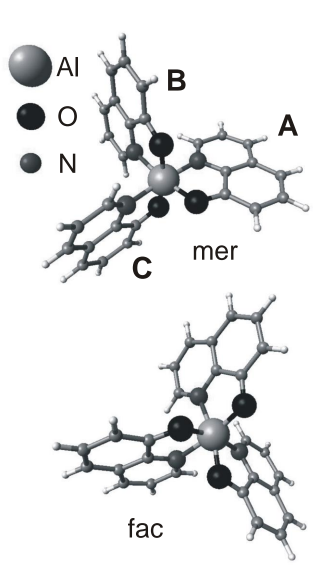


Figure 1. Chemical structure of the two geometrical Alq₃ isomers (*mer*: meridional, *fac*: facial) and labeling of the three ligands in *mer*-Alq₃.

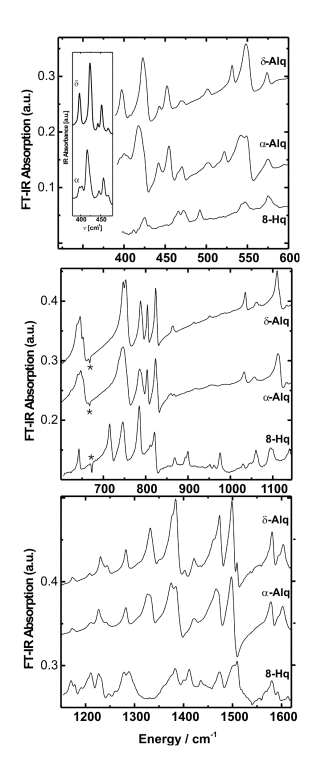


Figure 2: Comparison of the FTIR-spectra of δ -Alq₃ (upper trace), α -Alq₃ (middle trace) and hydoxiquinoline (8-Hq, lower trace) in the range from 350cm⁻¹ to 1650cm⁻¹. The inset shows α -Alq₃ and δ -Alq₃ in the region between 370 and 480cm⁻¹ measured with a better resolution below 420cm⁻¹ due to other beamsplitters *, CsI windows and samples on CsI pellets. The peak marked with an asterisk (*) is an artifact of the setup and is not related to the sample.

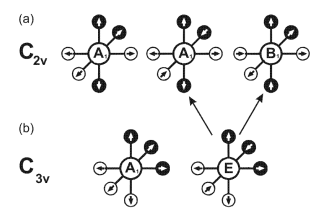


Figure 3: Schematic picture of the central part of the meridional (a) and the facial (b) isomer of Alq₃. Hollow and filled circles around the central Al-atom stand for oxygen and nitrogen atoms, respectively. The three stretching modes of the meridional molecule (C_{2v} -symmetry) and the two for the facial molecule (C_{3v} -symmetry, one is degenerated) are marked with arrows at the O and N atoms.

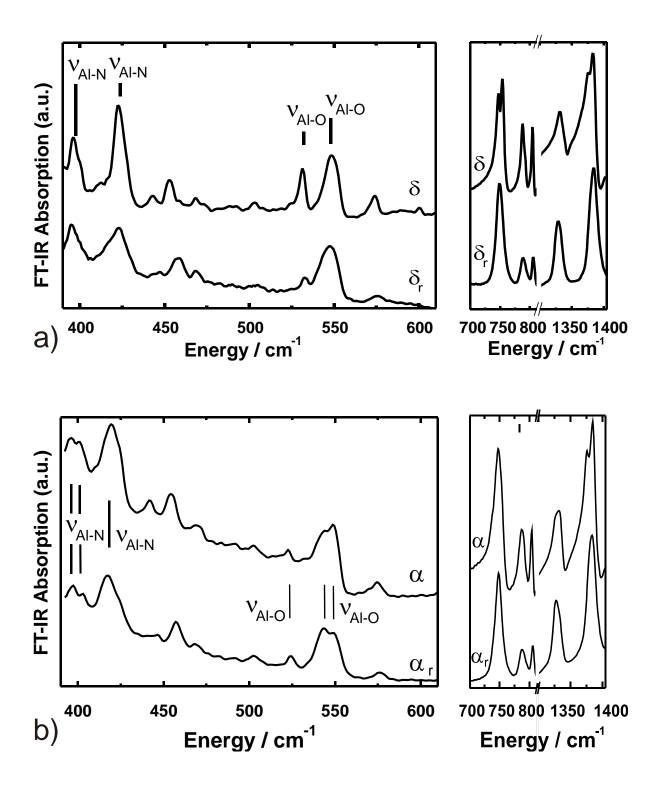


Figure 4: Comparisons of polycrystalline Alq₃ as obtained from the sublimation tube and very fine powders of the δ - and α -phases are shown in (a) and (b) respectively. The upper traces shows the polycrystalline samples as prepared and the lower trace the same samples after smoothly rubbing them mechanically, giving a very fine powder.