

Table 1: calculated  $D_0$  vibrational frequencies (B3-LYP/6-311++G(d,p))for 2-methyl-indenyl radical (2MIR), inden-2-ylmethyl radical (I2MR), *trans*-1-phenylallyl radical (*t*-1PAR), and *cis*-1-phenylallyl radical (*c*-1PAR).

2MIR			I2MR			<i>c</i> -1PAR			<i>t</i> -1PAR		
$\nu$	$\Gamma$	$\omega$	$\nu$	$\Gamma$	$\omega$	$\nu$	$\Gamma$	$\omega$	$\nu$	$\Gamma$	$\omega$
34	$a'$	268	34	$a'$	251	48	$a$	63	33	$a'$	160
33	$a'$	450	33	$a'$	448	47	$a$	153	32	$a'$	361
32	$a'$	488	32	$a'$	484	46	$a$	172	31	$a'$	414
31	$a'$	600	31	$a'$	573	45	$a$	299	30	$a'$	624
30	$a'$	667	30	$a'$	653	44	$a$	328	29	$a'$	631
29	$a'$	806	29	$a'$	823	43	$a$	412	28	$a'$	835
28	$a'$	874	28	$a'$	870	42	$a$	466	27	$a'$	991
27	$a'$	888	27	$a'$	961	41	$a$	498	26	$a'$	1008
26	$a'$	981	26	$a'$	974	40	$a$	562	25	$a'$	1045
25	$a'$	1036	25	$a'$	1032	39	$a$	629	24	$a'$	1100
24	$a'$	1117	24	$a'$	1088	38	$a$	669	23	$a'$	1178
23	$a'$	1169	23	$a'$	1155	37	$a$	698	22	$a'$	1188
22	$a'$	1189	22	$a'$	1171	36	$a$	742	21	$a'$	1210
21	$a'$	1195	21	$a'$	1181	35	$a$	802	20	$a'$	1223
20	$a'$	1223	20	$a'$	1212	34	$a$	817	19	$a'$	1303
19	$a'$	1277	19	$a'$	1219	33	$a$	843	18	$a'$	1320
18	$a'$	1325	18	$a'$	1287	32	$a$	850	17	$a'$	1354
17	$a'$	1344	17	$a'$	1337	31	$a$	923	16	$a'$	1375
16	$a'$	1383	16	$a'$	1400	30	$a$	982	15	$a'$	1462
15	$a'$	1409	15	$a'$	1423	29	$a$	990	14	$a'$	1502
14	$a'$	1451	14	$a'$	1480	28	$a$	995	13	$a'$	1511
13	$a'$	1476	13	$a'$	1486	27	$a$	1004	12	$a'$	1542
12	$a'$	1500	12	$a'$	1496	26	$a$	1023	11	$a'$	1593
11	$a'$	1535	11	$a'$	1534	25	$a$	1047	10	$a'$	1618
10	$a'$	1602	10	$a'$	1614	24	$a$	1110	9	$a'$	3132
9	$a'$	1621	9	$a'$	1625	23	$a$	1122	8	$a'$	3138
8	$a'$	3030	8	$a'$	3021	22	$a$	1180	7	$a'$	3147
7	$a'$	3138	7	$a'$	3098	21	$a$	1198	6	$a'$	3158
6	$a'$	3157	6	$a'$	3159	20	$a$	1236	5	$a'$	3164
5	$a'$	3163	5	$a'$	3164	19	$a$	1281	4	$a'$	3175
4	$a'$	3175	4	$a'$	3175	18	$a$	1331	3	$a'$	3186
3	$a'$	3187	3	$a'$	3188	17	$a$	1354	2	$a'$	3193
2	$a'$	3198	2	$a'$	3197	16	$a$	1437	1	$a'$	3231
1	$a'$	3227	1	$a'$	3200	15	$a$	1453	48	$a''$	91
51	$a''$	106	51	$a''$	47	14	$a$	1491	47	$a''$	137
50	$a''$	197	50	$a''$	127	13	$a$	1508	46	$a''$	255
49	$a''$	273	49	$a''$	238	12	$a$	1556	45	$a''$	411
48	$a''$	422	48	$a''$	310	11	$a$	1596	44	$a''$	499
47	$a''$	463	47	$a''$	422	10	$a$	1620	43	$a''$	599
46	$a''$	541	46	$a''$	537	9	$a$	3120	42	$a''$	692
45	$a''$	592	45	$a''$	559	8	$a$	3146	41	$a''$	759
44	$a''$	719	44	$a''$	740	7	$a$	3154	40	$a''$	819
43	$a''$	751	43	$a''$	743	6	$a$	3159	39	$a''$	839
42	$a''$	791	42	$a''$	788	5	$a$	3166	38	$a''$	856
41	$a''$	817	41	$a''$	800	4	$a$	3176	37	$a''$	914
40	$a''$	865	40	$a''$	884	3	$a$	3189	36	$a''$	973
39	$a''$	930	39	$a''$	942	2	$a$	3212	35	$a''$	987
38	$a''$	962	38	$a''$	974	1	$a$	3243	34	$a''$	999
37	$a''$	980	37	$a''$	1055						
36	$a''$	1165	36	$a''$	1487						
35	$a''$	3058	35	$a''$	3066						

Table 2: Optimized geometries for 2-methyl-indenyl radical (2MIR), inden-2-ylmethyl radical (I2MR).

**2MIR**

C	0.763194	-1.436407	0.000000
C	-0.433958	-0.611229	0.000000
C	-1.785716	-0.914811	0.000000
C	-2.719748	0.139018	0.000000
C	-2.297531	1.464628	0.000000
C	-0.926944	1.781920	0.000000
C	0.000000	0.749520	0.000000
C	1.448886	0.730672	0.000000
C	1.890733	-0.608453	0.000000
C	3.323397	-1.061144	0.000000
H	0.777494	-2.519078	0.000000
H	-2.127962	-1.944553	0.000000
H	-3.779989	-0.087354	0.000000
H	-3.031345	2.262608	0.000000
H	-0.608213	2.819177	0.000000
H	2.086115	1.606294	0.000000
H	3.859299	-0.694148	0.881160
H	3.391417	-2.151075	0.000000
H	3.859299	-0.694148	-0.881160

**I2MR**

C	3.285589	-1.002305	0.000000
C	1.977535	-0.603212	0.000000
C	0.832116	-1.433442	0.000000
C	-0.358185	-0.641051	0.000000
C	-1.714065	-1.013072	0.000000
C	-2.684201	-0.016292	0.000000
C	-2.324130	1.336882	0.000000
C	-0.974328	1.715732	0.000000
C	0.000000	0.733349	0.000000
C	1.505869	0.850453	0.000000
H	3.549434	-2.053408	0.000000
H	4.098271	-0.285763	0.000000
H	0.859902	-2.515838	0.000000
H	-1.998736	-2.059650	0.000000
H	-3.733998	-0.288410	0.000000
H	-3.096510	2.097581	0.000000
H	-0.705134	2.767173	0.000000
H	1.874779	1.388038	0.880130
H	1.874779	1.388038	-0.880130

Table 3: Optimized geometries for *trans*-1-phenylallyl radical (*t*-1PAR), and *cis*-1-phenylallyl radical (*c*-1PAR).

<b><i>t</i>-1PAR</b>			
C	-1.712257	-1.509950	0.000000
C	-2.722644	-0.543430	0.000000
C	-2.375847	0.809672	0.000000
C	-1.041562	1.188845	0.000000
C	0.000000	0.228952	0.000000
C	-0.376630	-1.136871	0.000000
C	1.363945	0.677812	0.000000
C	2.521724	-0.121020	0.000000
C	3.797364	0.371731	0.000000
H	4.657032	-0.286223	0.000000
H	3.989180	1.439544	0.000000
H	2.402286	-1.201257	0.000000
H	1.512021	1.755167	0.000000
H	-0.781117	2.242227	0.000000
H	-3.150768	1.568265	0.000000
H	-3.764548	-0.842009	0.000000
H	-1.973180	-2.562669	0.000000
H	0.384538	-1.907481	0.000000

<b><i>c</i>-1PAR</b>			
C	-1.512988	1.422474	-0.138841
C	-2.571946	0.539691	0.081634
C	-2.311521	-0.828568	0.188558
C	-1.011314	-1.300755	0.088682
C	0.083181	-0.423027	-0.103995
C	-0.208273	0.955344	-0.232069
C	1.405926	-0.990658	-0.195482
C	2.675283	-0.384024	-0.084833
C	3.014819	0.873788	0.338125
H	4.055718	1.171111	0.373198
H	2.291993	1.592365	0.699408
H	3.505412	-1.041578	-0.335981
H	1.422479	-2.064606	-0.359748
H	-0.818547	-2.365446	0.170765
H	-3.126934	-1.525767	0.346546
H	-3.587537	0.911173	0.155654
H	-1.708959	2.483346	-0.250418
H	0.587377	1.653824	-0.450098