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## **African Journal of Pure and Applied Chemistry**

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### **ARTICLE**

**Preparation, structural and thermal studies of boroxine adducts having aryl boronic acids and pyrazoles** Hezil Hassan

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### academic Journals

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Full Length Research Paper

## Preparation, structural and thermal studies of boroxine adducts having aryl boronic acids and pyrazoles

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Four new boroxine adducts ( $(B_3O_3(Ph)_3PzH)$  (1),  $(B_3O_3(Ph)_3(^{PztBu,iPr}H)_2)$  (2),  $(B_3O_3(PhF_2)_3PzH)$ . PzH (3) and  $(B_3O_3(PhF_2)_3(Pz^{tBu,iPr}H)_2)$  (4)) using phenylboronic acid, 3,5-difluorophenylboronic acid, 1H-pyrazole (PzH) and 3-tert-butyl-5-isopropyl pyrazole (Pz<sup>tBu,iPr</sup>H) were prepared and characterized by elemental analysis, IR, <sup>1</sup>H-NMR and X-ray diffraction. The crystallographic study reveals that PzH and Pz<sup>tBu,iPr</sup>H are bonded to boroxine molecule through B-N dative bond. It also demonstrates the different type of hydrogen bond interactions between adjacent molecules. The thermal stability of these adducts was investigated by TGA.

Key words: Boroxine, crystal structures, hydrogen bonding, thermal study.

#### INTRODUCTION

Because of importance in different synthetic reactions and significant applications in diverse areas, boronic acids are of great interest (Phillips and James, 2004; Davis and James, 2005; James, 2005; Striegler, 2003; Elfeky et al., 2010; Wimmer et al., 2009; Li et al., 2008). In recent years, boronic acids have also been used as

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Scheme 1. General method for the synthesis of 1-4.

exclusive building blocks in supramolecular chemistry (Fujita et al., 2008; Nishiyabu et al., 2011; Fossey and James, 2011). Boronic acids have  $-B(OH)_2$  group, and form the six-membered cyclic ring by simple dehydration of boronic acid. It is well established that boron in a cyclic ring  $(R_3B_3O_3, R = alkyl or aryl group)$  acts as a lewis acid and has a tendency to accept the lone pair of electrons from the N-donor ligands (lewis base), and involved in the formation of the B-N dative bond in adducts (Icli et al., 2011; Höpfi, 1999; Sheepwash et al., 2011, 2013; Jorge et al., 2012). N-donor ligands easily form 1:1 adducts with arylboronic acids even under mild reaction conditions, and thermodynamically favored over 1:2 or 1:3 adducts due to relief in boroxine ring strain. Adducts performing 1:2 and 1:3 boroxine-N-donor ligands stoichiometry are very limited (Beckett et al., 1995; Höpfi, 1999; Kua and lovine, 2005; Domingo et al., 2008; Saha et al., 2013; Jorge et al., 2016). In 1958, Synder et al. synthesized an adduct by using triphenylboroxine and pyridine by simple warming in anhydrous solvent (Snyder et al., 1958). In 2005, Cote et al. investigated highly stable and porous boronic acid derived covalent organic frameworks with large surface area (Cote et al., 2005). A large number of boroxine adducts with N-containing compounds have been studied due to wide commercial uses in various field like flame retardant materials, dopants, in Suzuki-Miyaura coupling reactions, non-linear optical materials, biosensors, covalent organic frameworks etc. (Bhat et al., 2011; lovine et al., 2008; Morgan et al., 2000; Mehta and Fujinami, 1997; Yang et al., 2002; Miyaura and Suzuki, 1995; Cote et al., 2005; Türker et al., 2009), but the structural characterization and thermal study of boroxine adducts with pyrazoles are not reported till now. This paper presents the synthesis, structural, and thermal study of four new boroxine adducts. The main purpose is to see the effect of substitution in phenyl boronic acids and pyrazoles ligands on the structure and crystal packing of these adducts.

#### MATERIALS AND METHODS

All synthesis was performed in air, and solvents were used as received. Phenylboronic acid, 3,5-difluorophenylboronic acid, and 1H-pyrazole were purchased from Aldrich Chemical Co. 3-tert-butyl-5-isopropyl pyrazole was synthesized by previously reported method (Imai et al., 1998). Elemental analysis was carried out on PerkinElmer Elemental analyzer. IR and <sup>1</sup>H-NMR spectra were recorded on Bruker ALPHA FT-IR and Bruker AM 400 MHz spectrometers, respectively. Thermal analysis was performed on PerkinElmer thermogravimetric analyzer.

#### Synthesis of adducts 1-4

Adducts 1-4, were synthesized according to scheme 1.

#### Synthesis of 1

A methanolic (10 ml) solution of phenylboronic acid (0.36 g, 3.00 mmol) and PzH (0.06 g, 1.00 mmol) was refluxed at 70°C for 4 h. Colorless crystals of 1 were obtained by the slow evaporation of solvent at room temperature in 0.24 g (62.5%) yield. Anal. Calcd for  $C_{21}H_{19}N_2O_3B_3$ : C, 66.40; H, 5.04; N, 7.37. Found: C, 65.12; H, 4.99; N, 7.17. IR (KBr, cm<sup>-1</sup>): 3381, 3196, 3012, 2889, 2626, 2317, 2029, 1921, 1814, 1709, 1627, 1461, 1235, 969, 699, 527. <sup>1</sup>H-NMR (400 MHz, CDCl<sub>3</sub>, *ppm*, 25°C): 6.21 (t, 1H, CH, Pz), 7.57 (d, 2H, CH, Pz), 12.61 (s, br, 1H, NH, Pz), 8.03 (dd, 6H, Ph), 7.35 (m, 9H, Ph).

#### Synthesis of 2

2 was obtained in 0.38 g (59%) yield by the same method as applied for 1 using  $Pz^{tBu,iPr}H$  (0.17 g, 1.00 mmol). Anal. Calcd for  $C_{38}H_{51}N_4O_3B_3$ : C, 70.84; H, 7.97; N, 8.69. Found: C, 70.19; H, 7.91; N, 8.49. IR (KBr, cm<sup>-1</sup>): 3418, 3143, 2956, 2865, 2247, 2139, 2069, 1971, 1829, 1786, 1609, 1573, 1437, 1296, 1049, 963, 827, 739,

Table 1. Crystal structures and	refinement parameters for 1-4
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Adducts	1	2	3	4
CCDC	1060378	1060379	1060376	1060377
Molecular formula	C <sub>21</sub> H <sub>19</sub> N <sub>2</sub> O <sub>3</sub> B <sub>3</sub>	C38H51N4O3B3	C24H17N4O3F6B3	C38H45N4O3F6B3
Mr	379.81	644.26	555.85	752.21
Crystal size (mm <sup>3</sup> )	0.31 × 0.26 × 0.19	0.28 × 0.23 × 0.17	0.23 × 0.17 × 0.11	0.33 × 0.26 × 0.19
Crystal system	triclinic	monoclinic	triclinic	triclinic
Space group	<i>P</i> -1	C 2/c	<i>P</i> -1	<i>P</i> -1
a (Å)	10.392(4)	19.887(18)	7.6511(6)	11.821(5)
b (Å)	11.741(6)	11.415(11)	11.5133(8)	13.078(5)
<i>c</i> (Å)	16.744(8)	18.432(18)	15.4005(11)	14.193(6)
a (deg)	84.17(3)	90.00	78.922(4)	84.43(2)
$\beta$ (deg)	89.84(3)	115.16(2)	75.632(5)	68.97(2)
γ (deg)	89.88(3)	90.00	77.296(4)	77.50(2)
<i>V</i> (Å <sup>3</sup> )	2032.4(16)	3787(6)	1268.36(16)	1999.1(15)
Z	4	4	2	2
$ ho_{Calc}$ (gcm <sup>-3</sup> )	1.241	1.130	1.455	1.250
μ(Mo Ka) (cm <sup>-1</sup> )	0.081	0.070	0.125	0.097
F(000)	792	1384	564	788
T(K)	296	296	296	296
Theta range for data collection (°)	1.74-28.36	3.56-26.00	1.38-25.00	1.54-25.00
Range of indices	-13, 12; -15, 11; -22, 16	-24, 24; -13, 14; -20, 13	-9, 9; -13, 13; -16, 18	-14, 12; -15, 14; -16, 10
No. of reflections collected	9692	2744	4442	6813
Unique reflections	2683	1737	2655	2132
Data/restraints/parameters	9692/0/523	2744/0/224	4442/0/362	6813/0/498
Goodness-of-fit	0.821	0.928	1.135	0.943
Final R indices(I > 2(I) aR1	0.065	0.065	0.061	0.118
<sup>b</sup> wR₂	0.142	0.186	0.174	0.245

 ${}^{a}R_{1} = \sum \left\| \mathsf{F}_{o} \right\| - \left\| \mathsf{F}_{c} \right\| / \sum \left\| \mathsf{F}_{o} \right\|, {}^{b}wR_{2} = \left\{ \sum \left( w \left( \mathsf{F}_{o}^{2} - \mathsf{F}_{c}^{2} \right)^{2} \right) / \sum w \left( \mathsf{F}_{o}^{2} \right)^{2} \right\}^{1/2}, \text{ where } w = 1 / \left( \sigma^{2}(\mathsf{F}_{o}^{2}) + (bP) \right) \text{ and } P = \left( \max \left( 0, \mathsf{F}_{o}^{2} \right) + 2\mathsf{F}_{c}^{2} \right) / 2$ 

645, 593, 497. <sup>1</sup>H-NMR (400 MHz, CDCl<sub>3</sub>, *ppm*, 25°C): 6.51 (s, 2H, CH, Pz), 4.49 (m, 2H, CH, Pz), 1.39 (d, 12H, CH<sub>3</sub>, Pz), 1.54 (s, 18H, CH<sub>3</sub>, Pz), 12.64 (s, br, 2H, NH, Pz), 8.03 (dd, 6H, Ph), 7.37 (m, 9H, Ph).

#### Synthesis of 3

3 was prepared in 0.53 g (64%) yield by using the method as outlined for 1 using 3,5-difluorophenylboronic acid (0.48 g, 3.00 mmol) and PzH (0.06 g, 1.0 mmol). Anal. Calcd for  $C_{24}H_{17}N_4O_3F_6B_3$ : C, 51.86; H, 2.91; N, 10.08. Found: C, 50.91; H, 2.89; N, 9.97. IR (KBr, cm<sup>-1</sup>): 3467, 3139, 3089, 2971, 2849, 2597, 2257, 2091, 1969, 1781, 1624, 1579, 1446, 1223, 1321, 1199, 1163, 1077, 929, 829, 644, 547. <sup>1</sup>H-NMR (400 MHz, CDCl<sub>3</sub>, *ppm*, 25°C): 6.27 (t, 2H, CH, Pz), 7.64 (d, 4H, CH, Pz), 12.67 (s, br, 2H, NH, Pz), 8.04 (dd, 6H, Ph), 7.40 (m, 3H, Ph).

#### Synthesis of 4

4 was synthesized in 0.45 g (59.8%) yield by the same method as described for 3 using Pz<sup>tBu,IPr</sup>H (0.17 g, 1.00 mmol). Anal. Calcd for  $C_{38}H_{45}N_4O_3F_6B_3$ : C, 60.84; H, 5.77; N, 7.46. Found: C, 60.13; H, 5.68; N, 7.07. IR (KBr, cm<sup>-1</sup>): 3418, 3129, 3018, 2917, 2755, 2601,

2239, 2069, 1911, 1837, 1755, 1629, 1457, 1213, 913, 729, 653, 547, 499. <sup>1</sup>H-NMR (400 MHz, CDCl<sub>3</sub>, *ppm*, 25°C): 6.53 (s, 2H, CH, Pz), 4.51 (m, 2H, CH, Pz), 1.42 (d, 12H, CH<sub>3</sub>, Pz), 1.57 (s, 18H, CH<sub>3</sub>, Pz), 12.69 (s, br, 2H, NH, Pz), 8.07 (dd, 6H, Ph), 7.39 (m, 3H, Ph).

#### X-ray diffraction analysis

All crystals were obtained from the slow evaporation of methanolic solutions, and mounted on glass capillaries. All data were collected on a Bruker Kappa four circle-CCD diffractometer with graphitemonochromated MoKa radiation, operated at 50 kV and 40 mA at 25°C. Data were corrected for Lorentz and polarization effects (Sheldrick, 1996), and the SHELXTL program package was used for the structure solution and refinement (Sheldrick 1990, 2000). The hydrogen atoms were placed in geometrically calculated positions by using a riding model, and non-hydrogen atoms were refined anisotropically. Diamond and Mercury softwares were used for the formation of images and hydrogen bonding interactions (Brandenburg, 2000). 1, 3 and 4 are crystallized in triclinic system with P-1 space group, while 2 in monoclinic system with C2/c space group. The crystallographic data, hydrogen bond distances, selected bond lengths and angles are shown in Tables 1, 2, Appendix S1 and S2, respectively.

 Table 2. Selected H-Bond parameters for 1-4.

Bond (symmetry)	<i>d</i> <sub>D-H</sub> (Å)	<i>d</i> <sub>H…A</sub> (Å) <i>d</i> <sub>D…A</sub> (Å)		<dh…a (°)<="" th=""></dh…a>
1				
C8-H8B5	0.931	3.153 (58)	3.987	150.0
C16-H16B5	0.930	3.058 (46)	3.983	173.5
C35-H35B3	0.930	3.154 (58)	3.988	150.1
N2-H2A····π	0.861	3.021 (46)	3.691	136.2
N4-H4A····π	0.861	2.646 (36)	3.360	141.1
С19-Н19п	0.930	2.572 (35)	3.338	139.9
С20-Н20п	0.930	3.502 (54)	3.767	119.9
C21-H21π	0.929	2.897 (43)	3.614	135.0
С40-Н40п	0.931	2.973 (45)	3.696	135.5
C41-H41…π	0.930	2.889 (44)	3.604	134.6
С42-Н42п	0.929	3.510 (37)	3.907	108.5
2				
C17-H17BB2	0.960	3.141 (15)	3.704	119.1
C17-H17CB2	0.959	3.358 (8)	3.704	103.7
С18-Н18В	0.961	3.005 (14)	3.863	149.4
3				
C20-H20B1	0.931	3,403 (7)	3,998	124.0
N4-H4AF5	0.860	2.337 (7)	3.054	141.1
C4-H4F3	0.930	2.850 (1)	3.734	159.1
C6-H6F1	0.930	2.685 (6)	3.272	117.2
C16-H16F2	0.930	2.470 (6)	3.394	172.6
C19-H19F6	0.930	2.728 (4)	3.396	129.4
C22-H22F2	0.930	2.722 (9)	3.351	125.6
C22-H22F3	0.930	2.558 (7)	3.179	124.6
4				
C26-H26CB3	0.960	3.469 (60)	4.100	125.3
C34-H34BB3	0.960	3.213 (40)	3.704	113.6
C34-H34CB3	0.961	3.327 (50)	3.704	105.6
C8-H8F6	0.929	2.731 (26)	3.540	146.1
C10-H10F1	0.930	2.818 (18)	3.567	138.3
C12-H12-F2	0.930	2.565 (26)	3.461	161.9
C27-H27CF4	0.960	2.804 (25)	3.483	128.5
C28-H28BF6	0.959	2.832 (32)	3.380	117.1
C33-H33BF5	0.960	2.661 (36)	3.493	145.3
C37-H37AF3	0.961	2.662 (33)	3.356	129.5

#### **RESULTS AND DISCUSSION**

All adducts  $(B_3O_3(Ph)_3PzH)$  (1),  $(B_3O_3(Ph)_3(Pz^{tBu,iPr}H)_2)$ (2),  $(B_3O_3(PhF_2)_3PzH).PzH$  (3) and  $(B_3O_3(PhF_2)_3(Pz^{tBu,iPr}H)_2)$  (4) have been prepared by using phenylboronic acid, 3,5-difluorophenylboronic acid and corresponding pyrazoles  $(PzH/Pz^{tBu,iPr}H)$  in methanol, and the different formulations were confirmed by elemental analysis, IR, NMR and crystallographic structure analysis. 2 and 4 are rare 1:2 adducts of 3,5difluorotriphenylboroxine and Pz<sup>tBu,iPr</sup>H, whereas 1 and 3 are 1:1 adducts of triphenylboroxine and PzH but 3 is crystallized with free pyrazole as solvate.

#### Infrared and NMR spectroscopy

1-4 show strong bands in the region 1460-1250 cm<sup>-1</sup>, and at 1255 cm<sup>-1</sup> due to B-O and B-N stretching bands, respectively (Smith and Northrop, 2014). NH stretching



Figure 1. (a) Molecular structure of 1. (b) 2-D sheet like framework. Color code: B, green; C, gray; H, purple; O, red; N, blue.

bands (3500-3400 cm<sup>-1</sup>) are shifted at 3100-3055 cm<sup>-1</sup> due to formation of adjacent B-N dative bond. The IR spectra do not show any O-H stretching vibration in the region of 3300-3200 cm<sup>-1</sup> that suggests the absence of O-H bands (Faniran and Shurvell, 1968). The formation of adducts have also been confirmed by the <sup>1</sup>H-NMR spectra, showing the prominent downfield shift in each case with respect to free compounds as shown in Table S3.

#### Structure description of 1-4

According to Figure 1a, the crystal structure of 1 shows two molecular units. In each unit one boron atom (B(1)) has tetrahedral geometry, while other two boron atoms (B(2) and B(3)) shows trigonal planar geometry. B(1) has Sp<sup>3</sup> hybridization due to the additional B-N dative bond. In one molecular unit, the B(2)-O(1), B(2)-O(3), B(3)-O(2)and B(3)-O(3) bond distances are in the range of 1.347(22) to 1.399(21) Å, and these are much smaller than that of B(1)-O(1) (1.468(20) Å) and B(1)-O(2) (1.456(23) Å) in B<sub>3</sub>O<sub>3</sub> ring. In the same manner, B(2)-C(7) (1.542(21) Å) and B(3)-C(13) (1.548(24) Å) bond lengths are also shorter than the B(1)-C(1) (1.595(25) Å). The B(1)-N(1) bond length is (1.617(23) Å) which is nearly matched with the reported literature (Wu et al., 1999). Another unit also follows the same pattern as previous one. The crystal packing shows that the one unit is noncovalently hydrogen bonded to neighboring unit through the various weak C-H...B (C8-H8...B5, 3.153 (58) Å; C16-H16...B5, 3.058 (46) Å; C35-H35...B3, 3.154 (58) Å), N-HA…π (N2-H2A…π, 3.021 (46) Å; N4-H4A…π, 2.646 (36) Å) and C-H···π (C19-H19···π, 2.572 (35) Å; C20-H20····π, 3.502 (54) Å; C21-H21···π, 2.897 (43) Å; C40-H40···π, 2.973 (45) Å; C41-H41···π, 2.889 (44) Å; C42-H42···π, 3.510 (37) Å) intermolecular interactions (Sarma and Baruah, 2009) (Appendix Figure S1). The angles between weak C-H···B bonds are 150.0 and 173.5°. All these interactions help to create the two dimensional sheet like framework (Figure 1(b)).

The asymmetric unit of 2 shows that two boron atoms B(1) and B(1') have tetrahedral geometry (Sp<sup>3</sup>), while other boron atom B(2) has a trigonal planar geometry (Sp<sup>2</sup>). In this molecular structure, B1<sup>1</sup>, O2<sup>1</sup>, C1<sup>1</sup>, N1<sup>1</sup> lie on inversion centers with symmetry code (i = 2-x, y, 0.5-z) (Figure 2(a)). From the X-ray structure, it is clear that both PztBuilPrH ligands in boroxine adduct are anti to each other which is more stable than syn configuration (lovine et al., 2008). The B(1)-O(1) (1.437(13) Å) and B(1)-O(2) (1.446(4) Å) bond distances are greater than the B(2)-O(2) (1.362(10) Å). Similarly, the B(1)-C(1) (1.621(8) Å) bond length is slightly higher than that of B(1)-C(7)(1.579(6)Å). The B(1)-N(1) bond length is (1.655(11) Å), which is greater than the B-N bond of 1. The crystal structure analysis describes that one molecular unit is hydrogen bonded with other units through C17-H17B…B2, 3.141(15) Å; C17-H17C…B2, 3.358 (8) Å and C18-H18B···· $\pi$ , 3.005 (14) Å non covalent interactions (Melikova et al., 2002) (Appendix Figure S2), and the C-H...B bond angles in 2 are 119.1 and 103.7°C which are lesser than the C-H···B bond angles of 1 (Table 2). Three dimensional zig-zag layered network is obtained by all C-H···B and C-H··· $\pi$  non covalent interactions (Figure 2b). 3 contains an adduct having 3,5-difluorophenylboronic acid and PzH with one free PzH in lattice (Figure 3a). In this structure, B(1) has tetrahedral geometry (Sp<sup>3</sup>), while B(2) and B(3) both atoms present trigonal planar



Figure 2. (a) Molecular structure of 2. Color code: B, green; C, gray; H, purple; O, red; N, blue. (b) 3-D zig-zag layered network. Color code: color by atomic displacement.



Figure 3. (a) Molecular structure of 3. Color code: B, green; C, gray; H, purple; O, red; N, blue; F, orange. (b) 3-D ladder like framework. Color code: adduct, green; free Pz, blue.

geometry (Sp<sup>2</sup>). The B(1)-O(1) (1.447(5) Å) and B(1)-O(2) (1.447(4) Å) bond distances are greater than the B(2)-O(2) (1.346(6) Å), B(2)-O(3) (1.378(6) Å), B(3)-O(1) (1.349(5) Å), B(3)-O(3) (1.381(4) Å). B(1)-C(1) bond length is 1.608(6) Å, which is slightly higher than the B(2)-C(7) (1.566(5) Å) and B(3)-C(13) (1.549(5) Å). B(1)-N(1) bond distance is 1.632(5) Å, which is higher than 1 but less than 2 (Appendix Table S1). The packing of crystal shows that the molecular units are interconnected to each other via various intermolecular hydrogen bond interactions, that is,  $C_{20}$ -H<sub>2</sub>0···B1, 3.403(7) Å;  $C_4$ -H<sub>4</sub>···F3,

2.850(1) Å; C<sub>6</sub>-H<sub>6</sub>...F1, 2.685(6) Å; C<sub>16</sub>-H<sub>16</sub>...F2, 2.470(6) Å; C<sub>19</sub>-H<sub>19</sub>...F6, 2.728(4) Å; C<sub>22</sub>-H<sub>22</sub>...F3, 2.558(7) Å (Appendix Figures S3 and S4(a)). On the other side, the free PzH in lattice also shows the N4-H4A...F5, 2.337(7) Å and C22-H22...F2, 2.722(9) Å noncovalent interactions with adducts (Appendix Figure S4(b)). The C-H...B bond angle in 3 is 124.0° which are lesser than the C-H...B bond angles of 1 and greater than 2. All these interactions design a three dimensional ladder like framework (Figure 3b).

The molecular structure of 4 has same structural



Figure 4. (a) Molecular structure of 4. Color code: B, green; C, gray; H, purple; O, red; N, blue; F, orange. (b) 3-D perspective view. Color code: color by atomic displacement.



Figure 5. TGA plot for 1-4.

dimension and geometry as 2, presented in Figure 4a. The B(1)-O(2) (1.438(18) Å), B(1)-O(3) (1.456(17) Å), B(2)-O(1) (1.461(18) Å) and B(2)-(O(3) (1.407(19) Å) bond distances are much greater than B(3)-O(1) (1.380(17) Å) and B(3)-O(2) (1.363(19) Å). Similarly, the B(1)-C(1) (1.607(20) Å) and B(2)- C(7) (1.619(17) Å) bond lengths are higher than that of B(3)-C(13) (1.569(20) Å). The B(1)-N(1) and B(2)-N(2) bond lengths are 1.644(25) and 1.658(28) Å, respectively, which are greater than that of 2. The X-ray crystal structural analysis shows that the molecule is intermolecular hydrogen bonded to adjacent molecules through C<sub>26</sub>-H<sub>26</sub>C···B3, 3.469 (60) Å; C<sub>34</sub>-H<sub>34</sub>B···B3, 3.213 (40) Å; C<sub>34</sub>-H<sub>34</sub>C···B3, 3.327(50) Å; C<sub>8</sub>-H<sub>8</sub>···F6, 2.731 (26) Å; C<sub>10</sub>-H<sub>10</sub>···F1, 2.818 (18) Å; C<sub>12</sub>-H<sub>12</sub>···F2, 2.565 (26) Å; C<sub>27</sub>-H<sub>27</sub>C···F4, 2.804 (25) Å; C<sub>28</sub>-H<sub>28</sub>B···F6, 2.832 (32) Å; C<sub>33</sub>-  $H_{33}B$ ···F5, 2.661 (36) Å and  $C_{37}$ - $H_{37}A$ ···F3, 2.662 (33) Å interactions (Appendix Figures S5 and S6). C-H···B bond angles in 4 are 125.3, 113.6 and 105.6° which are nearly matched with 2 and 3, but lesser than the C-H···B bond angles of 1. Three dimensional perspective view is created by involving all type of intermolecular interactions (Figure 4b).

#### Thermal study

All adducts 1-4 are stable at room temperature and their TGA plots are given in Figure 5. 1 and 2 show the one step decomposition. Adduct 1 decomposes completely in the temperature range of 169-236°C (~ 92.7% mass loss), while 2 shows the 91.2% mass loss in the

temperature range of 221-377°C. 3 decompose in two steps. In the first step free PzH releases between 103-123°C temperature range with 11.4% mass loss, while the second step corresponds to the removal of adduct in the 127-223°C temperature range with 76.7% mass loss. 4 follows the same pattern as 2 with 93.4% weight loss in the temperature range of 305-389°C.

#### Conclusions

This study has synthesized and structurally characterized four new boroxine adducts with different stoichiometric compositions (1:1, and 1:2) having phenyl boronic acid, 3,5-difluorophenylboronic acid, 1H-pyrazole and 3-tertbutyl-5-isopropyl pyrazole. The X-ray crystal structure studies conclude that on increasing the substituents of phenyl boronic acids and pyrazoles, the stoichiometry, number of non-covalent interactions varies from 1 to 4. and the C-H...B bond angles of 2-4 are lesser than 1. The molecules are intermolecularly hydrogen bonded to each other through various noncovalent interactions, and gives two/three dimensional frameworks. From this it is clear that the packing changes with the substitution in aryl boronic acids, pyrazoles, and diversity in adduct stoichiometry. Thermal study shows that all adducts are stable at room temperature and decompose at high temperature.

#### **Conflict of Interests**

The authors have not declared any conflict of interests.

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#### Supplementary information

The crystallographic data are available free of charge at <u>deposit@ccdc.cam.ac.uk</u> or <u>http://www.ccdc.cam.ac.uk</u>). Bond lengths and bond angles tables (Table S1, S2 and S3), hydrogen bond interaction Figures (Figure S1-S6) are also available.

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#### APPENDIX

#### Preparation, structural and thermal studies of boroxine adducts having aryl boronic acids and pyrazoles

erea seria ierigini		
1.468(20)	O6—B5	1.355(22)
1.347(22)	O6—B4	1.457(22)
1.456(23)	N1—B1	1.617(23)
1.351(21)	N3—B4	1.612(22)
1.399(21)	C1—B1	1.595(25)
1.374(18)	C7—B2	1.542(21)
1.457(20)	C13—B3	1.548(24)
1.342(22)	C22—B4	1.599(25)
1.378(19)	C28—B5	1.552(24)
1.389(21)	C34—B6	1.550(21)
	;	
1.437(13)	B2—O2'	1.362(10)
1.437(13)	N1—B1	1.655(11)
1.446(4)	C1—B1	1.621(8)
1.362(10)	C7—B2	1.579(6)
1 117(5)	02 P2	1 201(4)
1.447(3)	U3-B3	1.301(4)
1.349(3)		1.032(3)
1.447(4)		1.606(6)
1.340(0)	C/B2	1.566(5)
1.378(6)	C13—B3	1.549(5)
1.461(18)	N1—B1	1.644(25)
1.380(17)	N2—B2	1.658(28)
1.438(18)	C1—B1	1.607(20)
1.363(19)	C7—B2	1.619(17)
1.456(17)	C13—B3	1.569(20)
1.407(19)		()
	1.468(20) 1.347(22) 1.456(23) 1.351(21) 1.399(21) 1.374(18) 1.457(20) 1.342(22) 1.378(19) 1.389(21) 1.437(13) 1.437(13) 1.446(4) 1.362(10) 1.447(5) 1.349(5) 1.447(4) 1.346(6) 1.378(6) 1.461(18) 1.380(17) 1.438(18) 1.363(19) 1.456(17) 1.407(19)	1.468(20)       06B5         1.347(22)       06B4         1.456(23)       N1B1         1.351(21)       N3B4         1.399(21)       C1B1         1.374(18)       C7B2         1.457(20)       C13B3         1.342(22)       C22B4         1.378(19)       C28B5         1.389(21)       C34B6         1.437(13)       B2O2 <sup>i</sup> 1.437(13)       N1B1         1.446(4)       C1B1         1.362(10)       C7B2         1.447(5)       O3B3         1.349(5)       N1B1         1.349(5)       N1B1         1.349(5)       N1B1         1.346(6)       C7B2         1.378(6)       C13B3         1.461(18)       N1B1         1.380(17)       N2B2         1.438(18)       C1B1         1.363(19)       C7B2         1.438(18)       C1B1         1.363(19)       C7B2         1.456(17)       C13B3         1.407(19)       C1B1

Table S1. Selected bond lengths (Å) for 1-4.

Table S2. Selected bond angles (deg) for 1-4.

1			
B2—O1—B1	122.90(34)	O2—B1—C1	113.05(34)
B3—O2—B1	121.98(31)	O1—B1—C1	111.37(35)
B3—O3—B2	121.07(35)	O2—B1—N1	104.37(29)
B6—O4—B4	122.18(34)	O1—B1—N1	103.62(31)
B5—O5—B6	120.74(33)	C1—B1—N1	110.32(32)
B5—O6—B4	121.75(31)	O1—B2—O3	119.20(35)
C19—N1—B1	123.49(32)	O1—B2—C7	121.46(40)
N2—N1—B1	130.28(33)	O3—B2—C7	119.34(38)
N4—N3—B4	121.64(32)	O2—B3—O3	120.88(39)
C40—N3—B4	131.93(33)	O2—B3—C13	120.51(36)
C6—C1—B1	122.00(34)	O3—B3—C13	118.61(38)

Table S2. Contd.

C2-C1-B1	122.64(35)	O6—B4—O4	113.92(34)
C8—C7—B2	120.77(42)	O6—B4—C22	112.10(33)
C12—C7—B2	122.15(43)	O4—B4—C22	111.12(35)
C18—C13—B3	121.17(41)	O6—B4—N3	105.64(29)
C14—C13—B3	121.48(40)	O4—B4—N3	102.85(31)
C27—C22—B4	121.58(34)	C22—B4—N3	110.66(32)
C23—C22—B4	122.36(34)	O6—B5—O5	120.44(39)
C33—C28—B5	121.18(40)	O6—B5—C28	120.18(36)
C29—C28—B5	121.53(40)	O5—B5—C28	119.36(38)
C39—C34—B6	122.06(43)	O4—B6—O5	120.30(36)
C35—C34—B6	121.15(41)	O4—B6—C34	120.94(40)
O2—B1—O1	113.42(34)	O5—B6—C34	118.75(38)
2			
B1—O1—B1'	124.27(21)	O1—B1—C1	111.42(26)
B2—O2—B1	121.0(2)	O2—B1—C1	113.10(25)
C12—N1—B1	135.98(30)	O1—B1—N1	102.82(23)
N2—N1—B1	116.88(27)	O2—B1—N1	105.84(24)
C2—C1—B1	123.57(33)	C1—B1—N1	108.06(25)
C6—C1—B1	120.15(27)	O2—B2—O2'	123.17(15)
C8—C7—B2	121.49(13)	O2—B2—C7	118.41(11)
C8 <sup>i</sup> —C7—B2	121.49(13)	O2 <sup>i</sup> —B2—C7	118.41(11)
O1—B1—O2	114.69(26)		
2			
3	404 50(00)	00 04 04	440.07(04)
B3-01-B1	121.52(23)	02—B1—C1	112.07(24)
B2	121.57(24)		112.09(23)
B2	120.06(25)	02—B1—N1	104.64(25)
CZI-NI-BI	123.15(23)		104.29(25)
N2—N1—B1	130.35(27)	C1—B1—N1	106.95(24)
C6-C1-B1	121.42(26)	02—B2—03	120.90(34)
C2-C1-B1	121.51(24)	02—B2—C7	120.33(28)
C12—C7—B2	121.05(30)	03—B2—C7	118.77(29)
C8—C7—B2	119.79(30)	01—B3—03	120.89(27)
C18—C13—B3	121.78(27)	01—B3—C13	119.55(25)
C14—C13—B3	119.79(28)	O3—B3—C13	119.55(27)
02—B1—01	114.58(24)		
4			
B3—O1—B2	120.41(78)	O2—B1—C1	112.90(66)
B3—O2—B1	121.12(68)	O3—B1—C1	111.93(62)
B2—O3—B1	123.96(63)	02—B1—N1	105.85(62)
C21—N1—B1	134.00(69)	03—B1—N1	104.24(63)
N3—N1—B1	119 30(55)	C1—B1—N1	107 28(57)
C31—N2—B2	135 80(71)	03 - B2 - 01	115 41(74)
N4—N2—B2	117,25(63)	03 - B2 - C7	111.77(67)
C2-C1-B1	120.64(68)	01 - B2 - C7	111.58(77)
C6-C1-B1	122 18(70)	03 - B2 - N2	104 27(69)
C12—C7—B2	118 91(76)	01 - B2 - N2	106 56(72)
C8_C7_R2	125 66(65)	C7 = B2 = N2	106 44(64)
C18_C13_B3	122.00(00)	02 = R3 = 01	122 01(78)
C14 - C13 - B3	119 48(84)	02 B3 - C13	118 45(78)
0203	113 8/(7/)	01 - B3 - C13	119 50(87)
52 51-05	110.0+(74)		113.00(07)

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0/11	Pyrazoles/Adducts	Chemical Shifts (δ ppm)					
S/NO.		CH(Pz)	CH-N(Pz)	CH (CHMe <sub>2</sub> )	Me <sub>2</sub> (CHMe <sub>2</sub> )	Me <sub>3</sub> (CMe <sub>3</sub> )	NH
1.	PzH	6.13	7.15	_	_	_	9.81
2.	Pz <sup>tBu,iPr</sup> H	5.89	_	2.96	1.28	1.32	9.05
3.	Adduct 1	6.21	7.57	—	—	_	12.61
4.	Adduct 2	6.51	_	4.49	1.39	1.54	12.64
5.	Adduct 3	6.27	7.64	_	_	_	12.67
6.	Adduct 4	6.53	_	4.51	1.42	1.57	12.69

Table S3. Comparison of <sup>1</sup>H NMR spectra.



**Figure S1. 1** shows various C-H---B, C-H--- $\pi$  and N-H--- $\pi$  noncovalent interactions. Color code: B, green; C, gray; H, purple; O, red; N, blue.



Figure S2. 2 shows various C-H····B and C-H····π non covalent interactions. Color code: B, green; C, gray; H, purple; O, red; N, blue.



Figure S3. 3 shows various C-H---F non covalent interactions. Color code: B, green; C, gray; H, purple; O, red; N, blue; F, orange.



Figure S4 (a) and (b). 3 shows various C-H---B and N-H---F non covalent interactions. Color code: B, green; C, gray; H, purple; O, red; N, blue; F, orange.



**Figure S5. 4** shows various C-H---B non covalent interactions. Color code: B, green; C, gray; H, purple; O, red; N, blue; F, orange.



Figure S6. 4 shows various C-H···F non covalent interactions. Color code: B, green; C, gray; H, purple; O, red; N, blue; F, orange.

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