Corrigenda

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Page 78, Fig. 1.A.

The experimental modified molecular intensity points have been plotted with the wrong sign from s = 6.750 to $8.375 \, \text{Å}^{-1}$ inclusive. The experimental as well as the theoretical intensities are *positive* in this range. The difference curve, Fig. 1.B., is correct.

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In Table 3 the correct values of $|J(^{119}Sn-^{13}C)|$ for ring carbons of the two fluoro derivatives should read:

(1)	C(2) (C(3)	C(4)	C(5)	C(6)
		_			42.0 32.5
	67.0 4	67.0 42.0	67.0 42.0 50.2	67.0 42.0 50.2 11.8	67.0 42.0 50.2 11.8 50.2

Regression analysis parameters remain unaffected by these changes.

Page 108, The caption to Fig. 3 should read:

Correlation of $|J(Sn-CH_3)|$ with |J(Sn-C(1))| (averaged tin-117,119 values).

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