

# Anion Translocation in Organolithiums: A Mechanism for the Lithiation and Cyclisation of Tertiary Naphthamides

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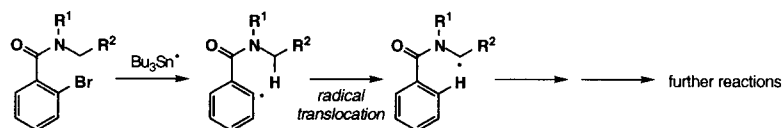
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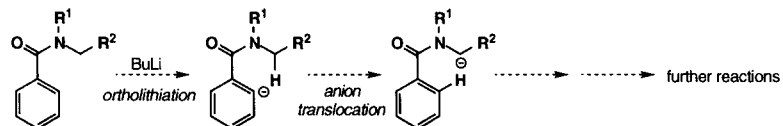
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**Abstract:** Deuterium labelling shows that an intramolecular proton transfer ("anion translocation") is a key step in the mechanism leading to an  $\alpha$ -lithiated tertiary naphthamide and thence to the products of anionic cyclisation. The kinetic isotope effect means that proton transfer from the *ortho* position can become the sole mechanism for  $\alpha$ -lithiation, though for undeuterated amides a parallel mechanism also operates in which lithiation occurs directly at the position  $\alpha$  to nitrogen. © 1998 Elsevier Science Ltd. All rights reserved.

Radical translocation – the intramolecular radical abstraction of a hydrogen atom – is a key step in some important radical reactions, and ingenious reaction sequences involving radical translocation from an amide protecting group have been used to generate  $\alpha$ -nitrogen substituted radicals.<sup>1</sup> In principle, the anionic equivalent of these radical translocation reactions, in which an anion (organolithium) formed under kinetic control undergoes an intramolecular proton transfer, could provide new routes to  $\alpha$ -nitrogen substituted organolithiums (Scheme 1b). Such "anion translocation" reaction sequences have not been generalised in synthetic chemistry,<sup>2</sup> but in this Letter we describe a reaction which proceeds, at least in part, by this process, and points to the potential development of anion translocation as a new synthetic strategy.

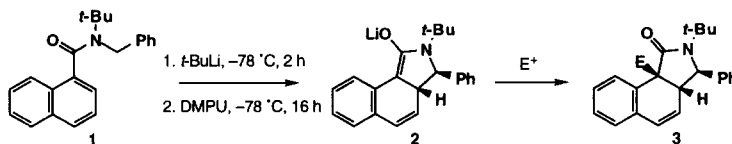


Scheme 1a: Radical translocation from an amide



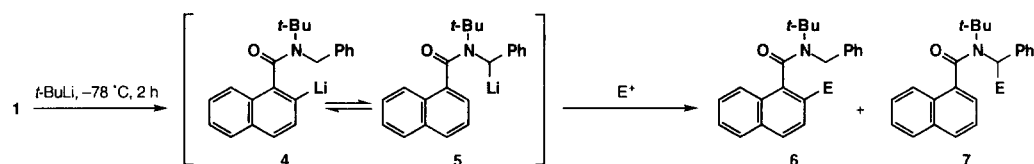
Scheme 1b: Anion translocation from an amide

We recently described an anionic cyclisation reaction of *N*-benzyl naphthamides (Scheme 2).<sup>3</sup> Treatment of *N*-*t*-butyl-*N*-benzyl-1-naphthamide **1** with *t*-BuLi and then DMPU<sup>4</sup> yields a tricyclic enolate **2** which can be quenched stereoselectively with electrophiles to give functionalised benzo[*e*]isoindolinones **3**.



Scheme 2: Anionic cyclisation of *N*-*t*-butyl-*N*-benzyl-1-naphthamide **1**

The mechanism of the cyclisation is intriguing: lithiation occurs in the absence of DMPU, but the organolithium intermediate does not cyclise until DMPU is added. Table 1 shows the results of trapping the uncyclised intermediate organolithium, prior to addition of DMPU: amide **1** was lithiated with *t*-BuLi at  $-78^{\circ}\text{C}$ , left for a period of 2 h and then quenched with an electrophile (Scheme 3)



Scheme 3: Electrophilic trapping of the organolithium intermediate

Table 1: Trapping the organolithium intermediate

Entry	Electrophile	E =	Yield <b>6</b> (%) <sup>a</sup>	Yield <b>7</b> (%) <sup>a</sup>	Ratio <b>6:7</b> <sup>b</sup>
1	D <sub>2</sub> O	D		59	1:2
2	MeI	Me	28	56	1:2
3	EtI	Et	33	13	2:1
4	Me <sub>3</sub> SiCl	Me <sub>3</sub> Si	n/d	n/d	>10:1
5	DMPU (2 min) then MeI	Me	4	62 <sup>c</sup>	1:10

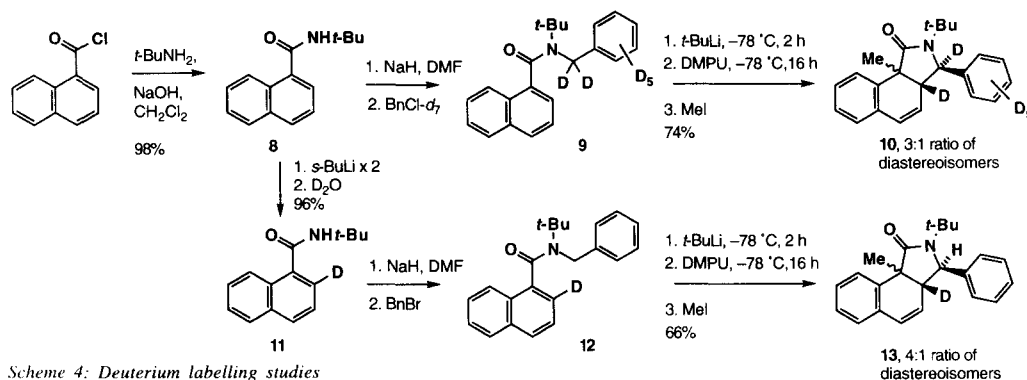
<sup>a</sup>Isolated yield. Remainder of the crude product is largely made up of some cyclisation product **3** and small amounts of remaining **1**. <sup>b</sup>Ratio from <sup>1</sup>H NMR of crude product. <sup>c</sup>Plus 23% of a mixture of diastereoisomers of **3**.

The trap never gave entirely one product, and indicated that the organolithium from which the enolate **2** forms in the cyclisation is in fact a mixture of *ortho*- and  $\alpha$ -lithiated compounds **4** and **5**. The electrophile-dependent outcome furthermore suggests that the organolithiums **4** and **5** are in equilibrium. Quenching with reactive electrophiles such as MeI or D<sub>2</sub>O gives a 1:2 mixture of *ortho* and  $\alpha$  products **6** and **7**, and we view this ratio as the nearest estimate we have of the actual ratio of **4:5** in solution. Less reactive electrophiles gave a greater proportion of the *ortho* product **6** – for EtI the ratio reverses to 2:1 **6:7**, while with Me<sub>3</sub>SiCl, only the *ortho* product **6** was detectable (albeit in low yield) in the product mixture. It seems that equilibration of **4** and **5** is rapid, with **5** slightly more stable but the **4** more reactive. MeI and D<sub>2</sub>O provide a snapshot view of this equilibrium; EtI and TMSiCl react selectively with **4** at a rate comparable with or slower than the rate at which the two species interchange. Table 1 gives us our first glimpse of an anion translocation, since this must be the mechanism by which **4** and **5** interconvert.

Adding DMPU to the mixture makes it cyclise, and how it does this is at least partly revealed by entry 5 of Table 1, which shows the result of adding DMPU but allowing insufficient time for the organolithiums to cyclise: DMPU forces the *ortho* organolithium **4** to undergo anion translocation to give the  $\alpha$  organolithium **5**. We know that this anion translocation must take place during the cyclisation, because only **5** can cyclise, yet yields for **1**→**3** (Scheme 2) are often near quantitative.

How does the mixture of organolithiums **4** + **5** arise? Unlike many other tertiary amides, *N*-*t*-butyl amides are conformationally uniform, with the *t*-butyl group *syn* to oxygen.<sup>5</sup> This appears to prevent a "complex-induced proximity effect"<sup>6</sup> operating in the lithiation  $\alpha$  to nitrogen (as is preceded for other tertiary amides).<sup>7</sup> Complex-induced ortholithiation of the naphthamide is however possible,<sup>8</sup> and the presence of the ortholithiated species **4** in the mixture suggested to us that *all* of the lithiated material might originate by

ortholithiation followed by rapid formation of an equilibrium mixture of **4** + **5** through anion translocation.<sup>9</sup> We tested this theory with some deuterium-labelled compounds. Alkylation of the sodium anion of *N*-*t*-butyl-1-naphthamide **8** with *d*<sub>7</sub>-benzyl chloride gave the Bn-*d*<sub>7</sub>-labelled amide **9**, and ortholithiation of the same secondary amide **8**, followed by D<sub>2</sub>O quench to give **11** and then amide alkylation with BnBr, gave the *ortho*-*d*-labelled amide **12**.



Scheme 4: Deuterium labelling studies

When **9** was treated with *t*-BuLi and then DMPU in THF at  $-78\text{ }^{\circ}\text{C}$ , followed by methyl iodide, it gave a good yield of a mixture of stereoisomers of **10**, with 100% deuterium labelling at the 3*a* (ring junction) position. This confirms a mechanism in which the *ortho* proton is removed to give Bn-*d*<sub>7</sub>-**4** which subsequently undergoes anion translocation by deuterium transfer to give an *ortho*-*d* organolithium **5** which then cyclises. Cyclisation of a mixture of **9** and **1** gave only unlabelled **3** (*E* = Me) and fully labelled (<1% *d*<sub>6</sub>) **10**, proving that the transfer is intramolecular.

When we carried out the same reaction on **12**, with the deuterium label at the *ortho* position, we got another result entirely. Cyclisation occurred to **13** with the deuterium label intact at the *ortho* position, indicating a mechanism in which *t*-BuLi has lithiated **12**  $\alpha$  to nitrogen directly, by-passing the ortholithiated species **4**.

The two reaction courses must be a consequence of the kinetic isotope effect, which can have very high values of  $k_{\text{H}}/k_{\text{D}}$  (>20) at  $-78\text{ }^{\circ}\text{C}$ .<sup>10</sup> We found that it was fully possible to direct alkylation to the *ortho* or the  $\alpha$  position purely by using deuterium substitution (Scheme 5): when **9** was lithiated and methylated (in the absence of DMPU, to prevent cyclisation) **14** (and <10%  $\alpha$ -methylated product) was obtained; when **12** was lithiated and methylated only **15** (and no *ortho*-methylated product) was obtained.<sup>11,12</sup>



Scheme 5: Deuterium directs the regioselectivity of alkylation

The answer to how much of the equilibrating mixture of unlabelled organolithiums **4** and **5** arises through each of the two possible routes for deprotonation – and hence which is the true mechanism of our cyclisation – was finally provided by competition experiments in which a mixture of **1** and **9** or **1** and **12** was lithiated with a deficit of *t*-BuLi (0.5 eq) and quenched with MeI, as summarised in Table 2. Undeuterated **1**

lithiates faster than either **9** or **12**, and the degree to which the rate of lithiation of each falls short of the rate of lithiation of **1** gives a crude estimate of the contribution that lithiation at their respective deuterated positions makes to the overall rate of lithiation. The results suggest that only one third of the **5** formed from **1** by *t*-BuLi is generated by ortholithiation–anion translocation, and the rest by deprotonation at the  $\alpha$  position.

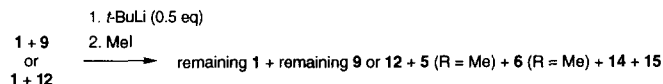


Table 2: Isotopic labelling and relative rates of reaction

Entry	Starting material contains: <sup>a</sup>			Product contains: <sup>a</sup>			Relative rate of lithiation of:		
	<b>1</b> (%)	<b>9</b> (%)	<b>12</b> (%)	<b>1</b> (%)	<b>9</b> (%)	<b>12</b> (%)	<b>1</b>	<b>9</b>	<b>12</b>
1	54	46	0	14	32	0	1	0.41	–
2	55	0	45	26	0	27	1	–	0.76

<sup>a</sup>by mass spectroscopy

Whatever the preferred mechanism of the cyclisation itself, intramolecular anion translocation is a viable route to organolithiums at sites of lower kinetic acidity, and we hope to apply this idea more generally.

#### Footnotes and References

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