Transforming LiTMP Lithiation of Challenging Diazines through Gallium Alkyl Trans-Metal-Trapping

Marina Uzelac, Alan R. Kennedy, Eva Hevia,* and Robert E. Mulvey*

Abstract: This study establishes a new trans-metal-trapping (TMT) procedure based on a mixture of LiTMP (the base) and tris(trimethylsilylmethyl)gallium [Ga(CH3SiMe3)3, GaR3] (the trap) that, operating in a tandem manner, is effective for the regioselective deprotonation of sensitive diazines in hydrocarbon solution, as illustrated through reactions of pyrazine, pyridazine, and pyrimidine, as well as through the N-S heterocycle benzothiazole. The metallo-activated complexes of all of these compounds were isolated and structurally defined.

As one of the most used strategies in synthetic campaigns, metalation chemistry is currently enjoying a remarkable period of advancement.[1] Lithium alkyls and amides remain front-runners for base candidates in routine C–H to C–metal transformations, but for non-routine, more formidable substrates, base selection is generally far from straightforward.[2] One of nature’s most important class of heterocycles utilized widely in agrochemicals, foodstuffs, pharmaceuticals, and many other commercial commodities, diazines are firmly in the formidable category, especially parent naked diazines devoid of substituents capable of assisting the direction of the metalation.[3] To avoid competition from nucleophilic addition caused by a low-lying LUMO in the heteroaromatic ring, bulky LiTMP (TMP is 2,2,6,6-tetramethylpiperidide) is preferred to more basic alkyl lithium reagents as the base for these π-deficient diazines, as first noted by Quéguiner.[4] However, lithiated diazene intermediates are generally unstable, so Knochel,[5] Kondo,[6] Mongin,[7] and Hevia[8] have employed coalitions of components typically but not exclusively based on the softer metal zinc to improve stability and to perform metalation under milder conditions. Though excellent progress has been made, these coalition approaches still have their limitations, as for example in the zinccation of pyrazine using [(THF)LiZn(TMPrBu2)], where no stoichiometric control is possible because only 2,5-dizincation occurs in a 1:1 base:pyrazine stoichiometry.[9] Moreover, a lack of definitive structural information still impoverishes understanding of this area, which in the most extreme “black box” cases leads to a misidentification of the actual metalating base.[9]

This paper reports a new trans-metal-trapping (TMT) procedure based on a mixture of LiTMP and tris(trimethylsilylmethyl)gallium [Ga(CH3SiMe3)3, GaR3] that is effective for regioselective diazine and benzothiazole deprotonation in hydrocarbon solution. Whereas alkyl lithium reactions are generally irreversible, LiTMP reactions tend to be pH-dependent equilibria. In TMT, these equilibria are shifted towards the desired lithiated substrate product by its interception by a trapping agent (Scheme 1).[10]

![Scheme 1. Two-step lithiation and carbanion trapping process in equilibrium controlling Trans-Metal-Trapping (TMT).](image)

We have previously employed the diorganoaluminum amide (Bu2Al(TMP)) as a trapping agent.[10] For this challenging diazine work, we elected to test GaR3 as a trap because its bulkiness compromises its ability to form a weakly basic ate complex with LiTMP and it has the potential to sedate the sensitive incipient carbanions on account of gallium’s strong carbophilicity (stronger than that of aluminum). Moreover, this organometallic trap exhibits good hydrocarbon solubility and does not require low temperature procedures, giving it a decided advantage over salt traps (for example, MgCl2, ZnCl2),[11] which generally need the use of ethereal solvents and often require low temperatures to avoid competing salt metathesis reactions. This potential has been duly realized through isolation and characterization (crystallographic, elemental analyses, and NMR spectroscopic) of an unprecedented series of gallium diazide and related complexes. As well as launching the concept of gallium TMT, this study reports the first crystal structures of metallo-diazine complexes made by metalation (C–H to C–metal) reactions for gallium and indeed, bar one exception for zinc,[3] for any...
metal. Furthermore, the study highlights that two-metal synergistic reactions are not confined to concerted, synchronized processes where the metals belong within the same reagent, but can be extended to tandem, stepwise processes involving two separately added reagents that do not form a co-complex.

The study first established through NMR spectroscopy that LiTMP and GaR₃ remain separate in benzene solution (Supporting Information). Such separation is essential to the lithiation step of TMT because, whereas free LiTMP is a strong base, combining it with, for example, iBu₃Al to form aluminate LiAl(TMP)(iBu₃), greatly diminishes deprotonating power. TMT was then attempted on three classical naked diazines, pyrazine, pyridazine, and pyridimine, as well as the related nitrogen–sulfur ring compound benzothiazole.

Pyrazine previously required four molar equivalents of LiTMP in THF at −75°C, but only to afford modest yields of 2-substituted derivatives (39–65%, depending on the electrophile) mixed in some cases with 2,5-disubstituted species (16%). When performed at room temperature in hexane solution, our new LiTMP-trialkylgallium approach in a 1:1:1 stoichiometry with pyrazine selectively afforded the 2-monogalllated pyrazine manifested in the crystalline complex [1-(PMDETA)Li-3-(GaR₃)]C₃H₇N₆ (1) isolated yield 61%: note NMR monitoring of reaction showed 1 is obtained quantitatively, see the Supporting Information).

The role of PMDETA is to aid crystallization and stabilization of the sensitive metallo species by chelating to lithium, but it is added at a later stage to avoid undergoing a competing TMT deprotonation itself. Stoichiometric control was also evident when the base:trap:substrate ratio was doubled to 2:2:1, giving the 2,5-digallated pyrazine [1,4-[(PMDETA)Li]-2,5-[(GaR₃)]C₃H₇N₆] 2 cleanly as crystals isolated in a 44% yield (Scheme 2). However, NMR monitoring of the reaction revealed 2 forms in a 55% yield, though a second regiosomer is formed in 33%, which appears to be the analogous product of 2,6-digallation. This stoichiometric control contrasts with the performance of zincate ([THF)LiZn(TMP)]Bu₃, which operates through a synchronised bimetallic synergy distinct to that of stepwise TMT, as it affords only the 2,5-disubstituted pyrazine even with a 1:1 base:substrate stoichiometry. Previously, excess LiTMP (1.5 equivalents) dispersed as 0.5ZnCl₂:TMEDA/1.5LiTMP in THF produced 59% of the isolated 2-iodopyrazine that reportedly decomposes at room temperature, but in hexane a significant amount of coupled dimer product was also seen. Note that a control reaction between pyrazine and gallate LiGaR₃ did not produce any gallation, but only R group addition, with concomitant dearamortization of the heterocycle (Supporting Information).

It may seem surprising that sensitive pyrazinyl mono- and di-carbanions can be trapped in crystalline form at room temperature and structurally defined, but this is where the structures become informative as they show that the heterocyclic units of 1 (Figure 1) and 2 (Figure 2) are cooperatively stabilized through coordination by both the Li and Ga centers that tie up the lone pairs of the N and C atoms, respectively. Both structures are monomeric with aggregation blocked at Li by tridentate PMDETA, though centrosymmetric 2 is tetranuclear having two Ga and two Li centers. Notably, 2 is the more congested structure with its GaR₃ units having proximal dispositions to the (PMDETA)Li units; whereas in 1 these units have a 1,3-separation. The Ga–sp³C(diazide) bond lengths show little variation with each other or with the Ga–sp³C(R group) bonds (see the Supporting Information for full crystallographic details and supporting NMR characterization).

With its 1,2-placement of N atoms, pyridazine offers a choice of metalation sites. Site selectivity in its metalation is exceptionally challenging, as evident from previous work using excess LiTMP in 0.5ZnCl₂:TMEDA/1.5LiTMP/I₃, which in hexane at room temperature achieved only 27% of the 3-iodo product mixed with the 4-iodo, and 3,6-diiodo derivatives as well as 54% unreacted pyridazine (note yields determined from NMR data). In THF, the yield of the 3-iodo product rises to 66% but only under extreme reflux conditions. On its own, LiTMP (4 equivalents) in THF at −75°C produced only 16–32% yields of 3-substituted pyrazidines following quenching with different electrophiles. When run in hexane solution at room temperature, our TMT reaction afforded a 51% yield of the isolated product [2-(PMDETA)Li-3-(GaR₃)]C₃H₇N₆ 3. Interestingly, ¹H NMR
monitoring of the reaction revealed an important effect of the order of metal reagent addition on reaction regioselectivity. Thus, when LiTMP is added as a solid to the hexane solution of GaR₃ and pyridazine, 3 is obtained in a 78% yield, along with small amounts of the C₄-gallated regioisomer (16% yield). Contrastingly, if the substrate is added to the hexane suspension of LiTMP and GaR₃, the yield of 3 decreases to 50%, and more C₄ metalated product is seen (36%). These contrasting results suggest an activating effect of the GaR₃ component, which perhaps can initially coordinate to the Lewis basic N atoms of the heterocycle, facilitating its lithiation at C3. The crystal structure of 3 (Figure 3) shows GaR₃ elects to sit at the most acidic 3-position adjacent to one N (confirmed in solution by NMR spectra; Supporting Information). A novel feature is the Li(PMDETA) unit bridging the two diazine N atoms [Li–N bond lengths 2.093(5) and 2.043(5) Å for the non-disordered molecule of the Z′ = 2 structure] leading to a 1, “2.5”-separation. Consequently, the spiro Li center, connecting the 3- and 2 × 5-atom rings, has a 5-coordinate geometry.

Because pyrimidine was found to be totally inert to LiTMP from 0°C to reflux temperatures, it seemed the greatest challenge to TMT. However, as in the case of 1–3, TMT was demonstrated tangibly through isolation and crystallographic characterization of a metalated derivative, here [1-(PMDETA)Li-6-(GaR₃)-C₆H₄N₃]·THF, 4. This was made in a 27% isolated crystalline yield, though NMR monitoring of the reaction shows that under the conditions studied 4 forms in a higher 59% yield. Its structure (Figure S13) exhibits many of the features in 1–3 with the proximal 1,6-separation of its GaR₃ and (PMDETA)Li akin to that of those in dimetalated pyrazine 2.

In view of the fact that the 2-lithiated derivative of the related nitrogen–sulfur heterocycle benzothiazole is known to exist simultaneously in ring-closed and ring-open forms, as best evidenced by Boche’s ¹³C NMR studies in D₂O-THF at −75°C, we extended the TMT study to this fused heterocycle.[¹⁴] Run at room temperature in hexane solution (Scheme 3), an equilibrium TMT reaction produced the crystalline complex [2-(GaR₃)-3-Li(PMDETA)]C₅H₄N₃S.

![Figure 3. Molecular structure of [2-(PMDETA)Li-3-(GaR₃)-C₆H₄N₃] (3) with 50% probability displacement ellipsoids. All H atoms except those in the C₆H₄N₃ ring have been omitted for clarity.](image)

5 (Scheme 3) in a remarkably high isolated yield of 84%. Significantly, 5 is quantitative in solution with no ring-opened metallo(2-isocyno)thiophenolate isomer detected. Deprotonative gallation of the C2 center was evident from its downfield resonance at 209.5 ppm in the ¹³C NMR spectrum. In the crystal, 5 follows the pattern in the TMT diazine series with the GaR₃ and Li(PMDETA) units adjacent on deprotonated C and N atoms, respectively, with a Ga1-C13-N1-Li1 torsion angle of −13.9(4°) (Figure 4). This first Ga TMT reaction of a N-S heterocycle is competitive with Mønigin’s LiTMP/CdCl₂-TMEDA in THF solution approach, which used excess (1.5) base equivalents for a 97% yield of 2-iodobenzothiazole after I₂ quenching, though no metallo intermediate was identified.[⁷,⁶⁵]

For synthetic campaigns, it is important that the new C–Ga bonds in these systems are accessible to electrophiles. In preliminary NMR spectroscopic experiments (see the Supporting Information for full details), on subjecting the gallated benzothiazole 5 to methyl trifluoromethanesulfonate (methyl triflate, MeOTf) we obtained a yield of 62% of the desired methylated product and did not observe any opening of the azole ring. In contrast, though this strong electrophile did quench the monogallated pyrazine 1, it proved too aggressive and decomposition ensued. Therefore, we turned to a second electrophile in trimethylsilyl chloride, which offers another potential pitfall in being too bulky. Reassur-
ingyly, this pitfall proved unfounded and the heterocyclic rings of both 1 and 5 were successfully converted to Me₂Si derivatives at their C–Ga sites (59 and 88% yields, respectively). These promising results demand that a full systematic, optimized study of all of these new gallated compounds is now undertaken with a series of electrophiles.

In summary, because LiTMP trans-metal-trapping via a gallium alkyl has been substantiated here with challenging sensitive unactivated diazines and benzothiazoles, this can potentially open the floodgates to a general improvement in many other mettallation reactions with various sensitive and non-sensitive substrates where LiTMP and related bulky bases give only low-to-moderate yields of products.

**Experimental Section**

Full experimental details and copies of NMR spectra are included in the Supporting Information. CCDC 1494979 (1), 1494980 (2), 1494981 (3), 1494982 (4), and 1494983 (5) contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre.

**Acknowledgements**

We are grateful to the European Research Council ((ERC-StG, MixMetApps) and the EPSRC (EP/N011384/1) for their generous sponsorship of this research and thank Dr. Donna Ramsay for her contributions to the development of TMT chemistry. Data supporting this research are openly available from http://dx.doi.org/10.15129/fce92db-989f-4430-b18a-d88478481c7f.

**Keywords:** diazines · gallium · heterocyclic chemistry · lithiation · structure elucidation

**How to cite:** Angew. Chem. Int. Ed. 2016, 55, 13147–13150

Angew. Chem. 2016, 128, 13341–13344

[1] a) J. Clayden, *Organolithiums: Selectivity for Synthesis*, Elsevier, Oxford, 2002; b) M. Schlosser, *Organometallics in Synthesis Third Manual*, Wiley, Hoboken, 2013.

[2] “The Chemistry of Organolithium Compounds”: *Patai’s Chemistry of Functional Groups, Vol. 1 and 2* (Ed.: Z. Rappoport, I. Marek), Wiley, New York, 2004.

[3] F. Chevallier, F. Mongin, *Chem. Soc. Rev.* 2008, 37, 595 –609.

[4] N. Plé, A. Turck, K. Couture, G. Quéguiner, *J. Org. Chem.* 1995, 60, 3781 –3786.

[5] a) M. Morin, P. Knochel, *Org. Lett.* 2009, 11, 1837 –1840; b) K. Groll, S. M. Manolikakes, X. M. du Jourdin, M. Jaric, A. Bredihhin, K. Karaghiosoff, T. Carell, P. Knochel, *Angew. Chem. Int. Ed.* 2013, 52, 6776 –6780; *Angew. Chem.* 2013, 125, 6909 –6913; c) Z. Dong, G. C. Clososki, S. H. Wunderlich, A. Unsinn, J. Li, P. Knochel, *Chem. Eur. J.* 2009, 15, 457 –468.

[6] T. Imahori, Y. Kondo, *J. Am. Chem. Soc.* 2003, 125, 8082 –8083.

[7] a) A. Seggio, F. Chevallier, M. Vaultier, F. Mongin, *J. Org. Chem.* 2007, 72, 6602 –6605; b) J.-M. L’Helgoualc’h, G. Bentabed-Abalsa, F. Chevallier, M. Yonechara, M. Uchiyama, A. Derdour, F. Mongin, *Chem. Commun.* 2008, 42, 5375 –5377.

[8] a) V. L. Blair, D. C. Blakemore, D. Hay, E. Hevia, D. C. Pryde, *Tetrahedron Lett.* 2011, 52, 4590 –4594; b) S. E. Baille, V. L. Blair, D. C. Blakemore, D. Hay, A. R. Kennedy, D. C. Pryde, E. Hevia, *Chem. Commun.* 2012, 48, 1985 –1987.

[9] D. R. Armstrong, A. R. Kennedy, R. E. Mulvey, J. A. Parkinson, S. D. Robertson, *Chem. Sci.* 2012, 3, 2700 –2707.

[10] a) D. R. Armstrong, E. Crosbie, E. Hevia, R. E. Mulvey, D. L. Ramsay, S. D. Robertson, *Chem. Sci.* 2014, 5, 3031 –3045; b) W. Clegg, E. Crosbie, S. H. Dale-Black, E. Hevia, G. Honeyman, A. R. Kennedy, R. E. Mulvey, D. L. Ramsay, S. D. Robertson, *Organometallics* 2015, 34, 2580 –2589.

[11] A. Frischmuth, M. Fernández, N. M. Bártil, E. Rarchainer, H. Zipse, G. Berioni, H. Mayr, K. Karaghiosoff, P. Knochel, *Angew. Chem. Int. Ed.* 2014, 53, 7928 –7932; *Angew. Chem.* 2014, 126, 8062 –8066.

[12] The limited solubility of LiTMPs in arene solvents such as benzene or toluene precluded DOSY NMR studies. Comparison of the ‘H NMR spectrum of a LiTMP/GaR₃ mixture with those of the individual components evidenced the lack of co-complexation between these reagents.

[13] D. R. Armstrong, E. Branner, T. Cadenbach, E. Hevia, A. R. Kennedy, *Organometallics* 2013, 32, 480 –489.

[14] C. Hilt, F. Bosold, K. Harms, M. Marsch, G. Boche, *Chem. Ber.* 1997, 130, 1213 –1221.

[15] K. Snégarroff, J.-M. L’Helgoualc’h, G. Bentabed-Ababas, T. T. Nguyen, F. Chevallier, M. Yonechara, M. Uchiyama, A. Derdour, F. Mongin, *Chem. Eur. J.* 2009, 15, 10280 –10290.