

Synthetic Methods

Organoytterbium Ate Complexes Extend the Value of Cyclobutenediones as Isoprene Equivalents**

Emma Packard, David D. Pascoe, Jacques Maddaluno, Théo P. Gonçalves, and David C. Harrowven*

In memory of Margaret Harrowven

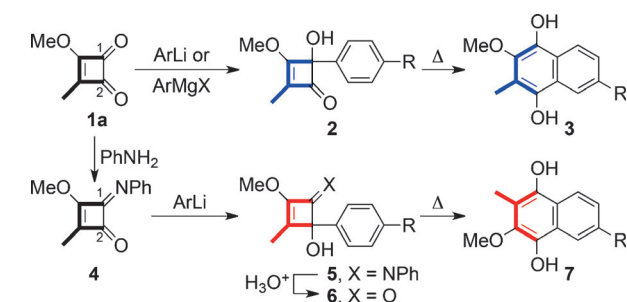
3-Methyl-4-methoxycyclobuten-1,2-dione (**1a**) has long been recognized as a valuable isoprene equivalent in natural products total synthesis, particularly for the preparation of hydroquinones, quinones, and their benzannulated analogues (e.g. Scheme 1).^[1,2a] It is readily introduced to a substrate as the electrophilic component in organolithium or Grignard addition reactions, for example, **1a**→**2**. Here, the differential reactivity of the C1 and C2 carbonyl groups in **1a** adds to its value as a synthon by providing a reliable and predictable means of achieving the ubiquitous head to tail connectivity of isoprene units.^[1,2a]

Though regiocontrol in the addition of Grignard and organolithium reagents to **1a** is of critical importance, it also

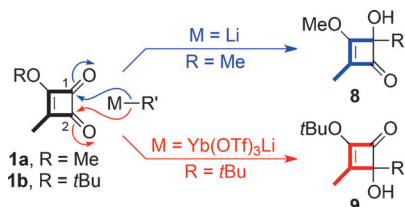
imposes a severe limitation. As evidenced by the example in Scheme 1, while the method is convenient for the synthesis of benzohydroquinones such as **3**, it proves cumbersome when targeting the regioisomeric series **7**. In this case addition of a carbon nucleophile to the vinylogous ester carbonyl (C2) of **1a** is required to achieve the desired outcome. Consequently, a protecting group strategy must be invoked to mask the more reactive C1 carbonyl (e.g. **1a**→**4**).^[2] Addition of the Grignard or organolithium reagent to C1 is then followed by deprotection (e.g. **5**→**6**), which can be difficult to achieve efficiently because of the presence of the acid sensitive tertiary alcohol and enol ether functions.

Herein we report an expedient solution to that long-standing problem and reveal some hitherto unknown facets of organoytterbium reactivity. In essence, while organolithium and Grignard reagents favor addition to the C1 carbonyl of cyclobutenedione (**1a**), the corresponding organoytterbium reagents give exclusive addition to the C2 carbonyl of **1b** (Scheme 2).^[3]

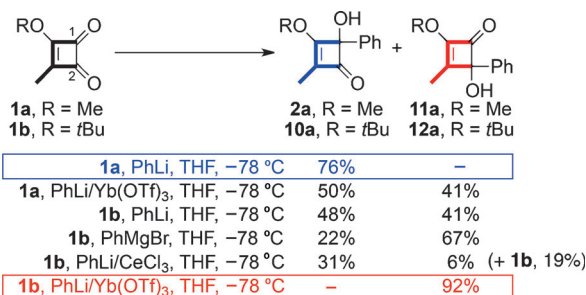
The discovery was made during an optimization study aimed at reducing side reactions resulting from deprotonation of the C3 methyl substituent in **1a**. In the addition of PhLi, for example (Scheme 3), these appeared to limit the yield of the



Scheme 1. Use of **1a** as an isoprene equivalent in the synthesis of benzohydroquinones.



Scheme 2. Dichotomous regioselectivity in the addition of organolithium and organoytterbium reagents to cyclobutenediones. Tf=tri-fluoromethanesulfonyl.



Scheme 3. Regioselectivity in the additions of various phenyl organometallics to **1a** and **1b**. THF=tetrahydrofuran.

[*] E. Packard, Dr. D. D. Pascoe, T. P. Gonçalves, Prof. D. C. Harrowven Chemistry, University of Southampton Highfield, Southampton, SO17 1BJ (United Kingdom) E-mail: dch2@soton.ac.uk

Dr. J. Maddaluno
FR CNRS 3038, IRCOF, Université de Rouen
76821 Mont St. Aignan Cedex (France)

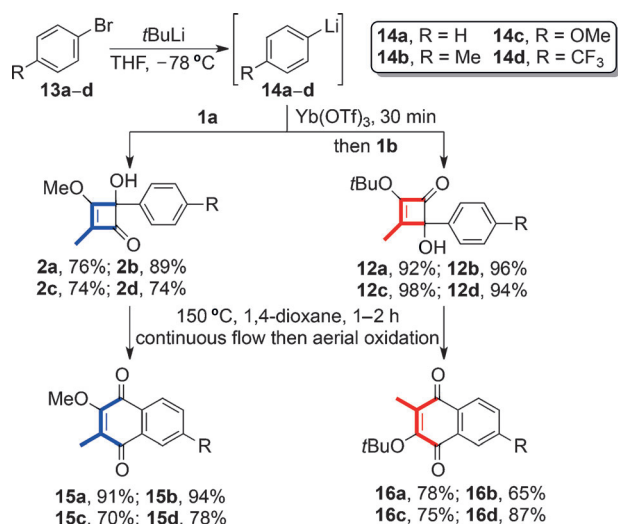
[**] Financial support from the EPSRC and ERDF (IS:CE-Chem & InterReg IVa program 4061) is gratefully acknowledged. We would also like to acknowledge Dr. Edmund Lee, the IRIDIS High Performance Computing Facility, the EPSRC National Service for Computational Chemistry Software (NSCCS), and their associated support staff for help in the completion of this work.

Supporting information for this article is available on the WWW under <http://dx.doi.org/10.1002/anie.201307193>.

© 2013 The Authors. Published by Wiley-VCH Verlag GmbH & Co. KGaA. This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

adduct **2a** to approximately 70–80% when employing standard protocols (e.g. THF at -78°C).^[4] Inspired by reports from the groups of Molander and Procter on the efficient addition of organoytterbium reagents to carbonyl compounds,^[5] we decided to examine the addition of phenyllytterbium triflate to **1a**.^[2] The result we attained was quite unexpected, as the reaction led to the formation of both C1 and C2 adducts, **2a** and **11a**, respectively, in the corresponding yields of 50% and 41% (Scheme 3). Having noted similar anomalies in the additions of PhLi and PhMgBr to **1b**, we were delighted to find that the addition of phenyllytterbium triflate to **1b** resulted in a complete reversal of the normal regiochemical course, thus giving the C2 adduct **12a** in a remarkable 92% yield upon isolation.^[5,6]

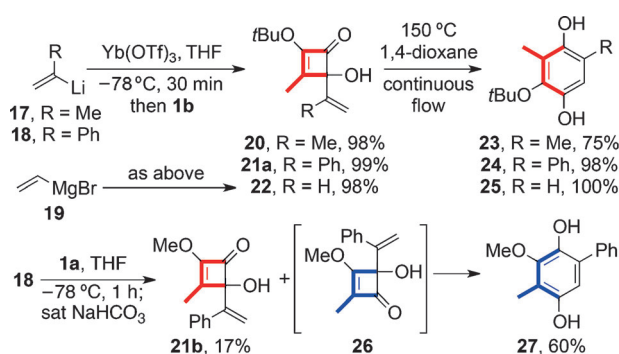
To demonstrate the value of the method, a series of aryllytterbium reagents were prepared and reacted with **1b**. These included *ortho*-,^[7] *meta*-,^[7] and *para*-substituted aromatics with a variety of substituents (+I, -I, and +M), and a heteroaromatic example^[7] (see Scheme 4 and the Supporting Information). In each case the C2 adducts **12** were formed



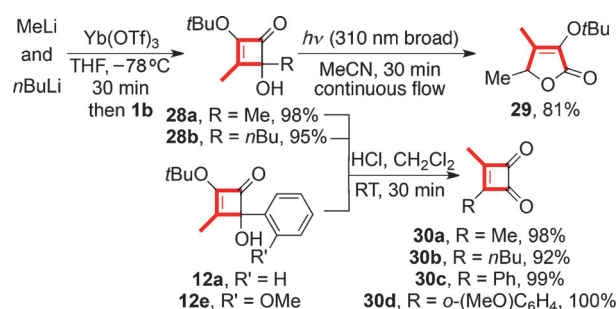
Scheme 4. Synthesis of regioisomeric series of benzoquinones using aryllithium and aryllytterbium additions to **1a** and **1b**.

in excellent yield. This result was in stark contrast to the corresponding organolithium additions to **1a** where the major product **2** resulted from addition to the C1 carbonyl.^[7] For completeness, all adducts were subjected to thermolysis and aerial oxidation, thus leading to the complementary series of benzoquinones **15** and **16**.

The complementary reactivity extended to vinyllytterbium and vinyllytterbium triflate additions to **1a** and **1b** as demonstrated by the syntheses of the hydroquinones **23–25** and **27** (Scheme 5).^[1] Notably, vinyllytterbium triflate derived from vinylmagnesium bromide displayed the same regiochemical preference as those derived from vinyllytterbium intermediates. Interestingly, the addition of the vinyllytterbium **18** to **1a** proved unusual in that it gave **27** directly after a bicarbonate quench at -78°C and warming to ambient temperature. The minor product of the reaction was the C2 adduct **21b**,^[7] rather than the corresponding hydroquinone.



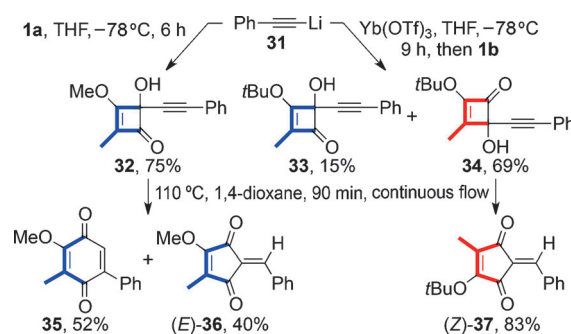
Scheme 5. Synthesis of regioisomeric series of hydroquinones by vinyllytterbium and vinyllytterbium additions to **1a** and **1b**.



Scheme 6. Alkyllytterbium additions to **1b** for the synthesis of 5H-furanones and diorganylcyclobutenediones.

The addition of methyllytterbium triflate to **1b** also proved facile (Scheme 6). In contrast, the analogous sequence with *n*BuLi required a 2:1 equivalence with ytterbium(III) triflate to achieve a high yield of the C2 adduct **28b**.^[8] The photochemical rearrangement of the adduct **28a** into the furanone **29**,^[9] and the facile generation of the diorganylcyclobutenediones **30** by exposure of these adducts to acid, provide further illustrations of the method's utility.

For completeness, the addition of phenylalkynyllytterbium triflate to **1b** was examined (Scheme 7). The reaction proved sluggish (9 h at -78°C) in comparison with other additions and regioselectivity was attenuated, with the formation of cyclobutenones **34** and **33** in 69% and 15% yield, respectively. In contrast, the addition of the organolithium **31** to **1a** gave the cyclobutenone **32** as the sole isolated product.



Scheme 7. Phenylalkynyllithium and ytterbium additions to **1a** and **1b**, respectively.

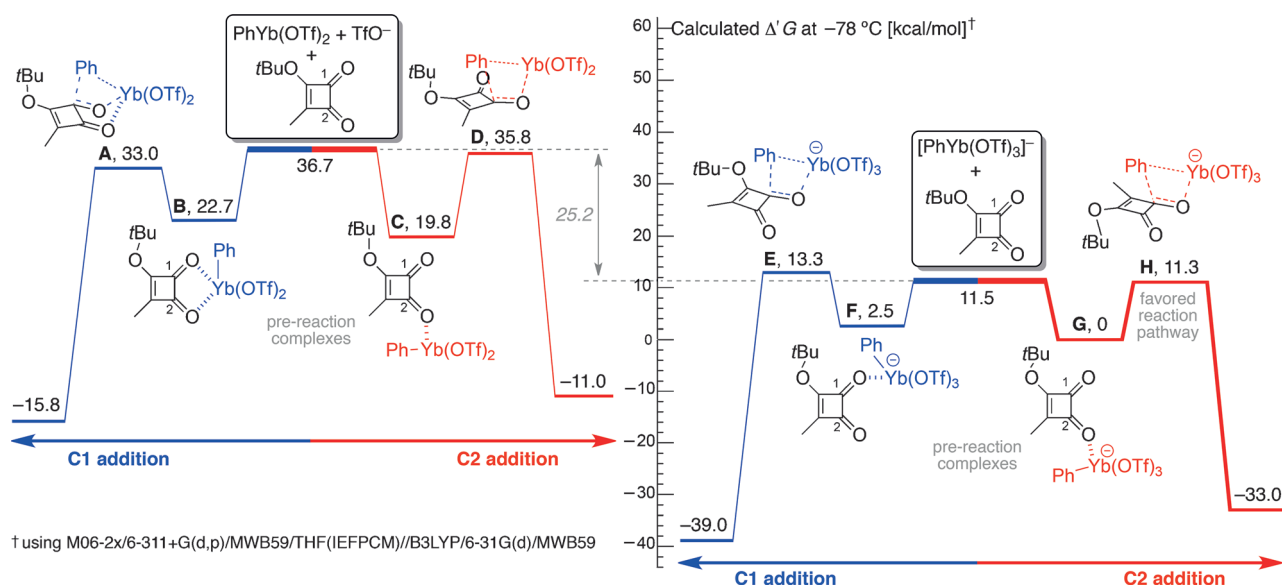


Figure 1. Calculated free energy barriers for C1 and C2 addition in the reactions of **1b** with PhYb(OTf)₂ and [PhYb(OTf)₃]⁻ in THF.

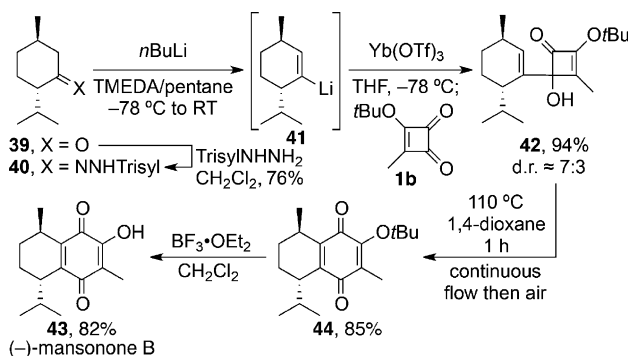
Thermolyses of the adduct **32** and **34** exposed a further anomaly, with the former giving a 5:4 mixture of the quinone **35** and cyclopentenedione (*E*)-**36**, while the latter gave only the cyclopentenedione (*Z*)-**37**.

To gain insights into the mechanistic course of organoytterbium addition reactions we decided to initiate a computational study of the reaction of PhYb(OTf)₂ with **1b**. However, these investigations proved fruitless, with DFT calculations predicting a preference for reaction at the ketonic C1 carbonyl (i.e. $\Delta G^\ddagger_A < \Delta G^\ddagger_D$, Figure 1).^[10] The results led us to question the nature of the organometallic reagent. Based on limited literature precedent,^[11] we had assumed that the reaction of PhLi with Yb(OTf)₃ would first give the ate complex PhYb(OTf)₃Li, which in turn would collapse to PhYb(OTf)₂ and LiOTf. When this premise was tested computationally, our calculations indicated that loss of a triflate ligand from the ate complex would cost approximately 25 kcal mol⁻¹ at -78 °C! Moreover, when the reactivity of [PhYb(OTf)₃]⁻ towards **1b** was examined, the predicted outcome mirrored our experimental findings by revealing a kinetic preference for phenyl addition to the C2 carbonyl (i.e. $\Delta G^\ddagger_H < \Delta G^\ddagger_E$, Figure 1).

The modeling also exposed some seminal facets of organoytterbium reactivity (Figure 1). In particular, the pre-reaction complexes of [PhYb(OTf)₃]⁻·**1b** (**F** and **G**) each showed single ytterbium–carbonyl interactions leading to trapezoidal transition states. The strength of those interactions was a key factor in determining the regiochemical course of addition, with $\Delta G^\ddagger_G$ less than $\Delta G^\ddagger_F$ by about 2.5 kcal mol⁻¹. Additionally, steric interactions in the addition phase play an important role, as evidenced by 1) the need to rotate the *t*Bu ether during C1 addition (**F**→**E**, Figure 1) at a cost of about 2.6 kcal mol⁻¹^[12] and 2) the low selectivity observed for addition of [PhYb(OTf)₃]⁻ to **1a** compared to that for **1b** (Scheme 3).

Finally, to demonstrate the method's potential in terpenoid synthesis we decided to target mansonone B, a natural

product whose identity has yet to be determined with rigor but seemed likely to be as depicted in **43**.^[13] Our synthesis began with (–)-menthone (**39**; Scheme 8), which was readily



Scheme 8. First total synthesis of (–)-mansonone B. TMEDA = *N,N,N',N'*-tetramethylethylenediamine, Trisyl = 2,4,6-trisopropylphenyl-sulfonyl.

transformed into the required vinyl lithium reagent **41** by means of a Shapiro reaction.^[14] Transmetalation with ytterbium triflate, and subsequent addition of the resulting vinylytterbium intermediate to **1b**, gave a 7:3 diastereomeric mixture of the C2 adduct **42** in 94% yield. Thermolysis of that mixture under continuous flow at 110 °C with concomitant aerial oxidation, gave the quinone **44**, from which our target **43** was readily derived. Pleasingly, the physical and spectral characteristics displayed by our synthetic sample (–)-**43** matched those reported for mansonone B in its isolation paper,^[13a] thus confirming the regiochemical and relative stereochemical identity of the natural product. Its absolute configuration remains uncertain because of a lack of available optical rotation data for comparison.

In conclusion, we have shown that organoytterbium additions to **1b** are easy to effect, proceed in excellent

yield, and follow a complementary regiochemical course to related organolithium additions to **1a**. Alkyl, aryl, hetaryl, vinyl, and alkynyl ytterbium reagents participate in the reaction, greatly extending the utility of cyclobutenone rearrangements by providing a predictable means of introducing this isoprene unit in a regioselective manner. High-level computational studies have provided seminal insights into the nature of organoytterbium intermediates and their reactivity. Of particular note is the realization that reactions proceed via ytterbium ate complexes.

Received: August 15, 2013

Published online: October 23, 2013

Keywords: density functional calculations · flow chemistry · natural products · rearrangement · ytterbium

- [1] a) M. Mohamed, T. P. Gonçalves, R. J. Whitby, D. C. Harrowven, H. F. Sneddon, *Chem. Eur. J.* **2011**, *17*, 13698–13705; b) D. Knueppel, S. F. Martin, *Tetrahedron* **2011**, *67*, 9765–9770; c) E. Peña-Cabrera, L. S. Liebeskind, *J. Org. Chem.* **2002**, *67*, 1689–1691. See also d) A. Padwa, M. J. Chughtai, J. Boonsombat, P. Rashatasakhon, *Tetrahedron* **2008**, *64*, 4758–4767; e) D. C. Harrowven, D. D. Pascoe, I. L. Guy, *Angew. Chem.* **2007**, *119*, 429–432; *Angew. Chem. Int. Ed.* **2007**, *46*, 425–428; f) D. C. Harrowven, D. D. Pascoe, D. Demurtas, H. O. Bourne, *Angew. Chem.* **2005**, *117*, 1247–1248; *Angew. Chem. Int. Ed.* **2005**, *44*, 1221–1222; g) S. K. Verma, E. B. Fleischer, H. W. Moore, *J. Org. Chem.* **2000**, *65*, 8564–8573; see the Supporting Information for additional references.
- [2] a) B. M. Trost, O. R. Thiel, H.-C. Tsui, *J. Am. Chem. Soc.* **2003**, *125*, 13155–13164; b) B. M. Trost, O. R. Thiel, H.-C. Tsui, *J. Am. Chem. Soc.* **2002**, *124*, 11616–11617; See also c) G. Oppermann, E. Schaumann, M. Stranberg, H. W. Moore, *Synthesis* **2010**, 2027–2038; d) E. Schaumann, G. Oppermann, M. Stranberg, H. W. Moore, *Aust. J. Chem.* **2010**, *63*, 1656–1664; e) M. P. Winters, M. Stranberg, H. W. Moore, *J. Org. Chem.* **1994**, *59*, 7572–7574.
- [3] The cyclobutenediones **1a** and **1b** were each prepared in two steps from squaric acid using literature methods: a) M. W. Reed, D. J. Pollart, S. T. Perri, L. D. Foland, H. W. Moore, *J. Org. Chem.* **1988**, *53*, 2477–2482; b) A. Enhsen, K. Karabelas, J. M. Heerding, H. W. Moore, *J. Org. Chem.* **1990**, *55*, 1177–1185. Yb(OTf)₃·xH₂O was dried by heating it at 240 °C in a Kugelrohr oven for 4 d under high vacuum.
- [4] a) S. L. Xu, H. W. Moore, *J. Org. Chem.* **1992**, *57*, 326–338; b) L. S. Liebeskind, S. Iyer, C. F. Jewell, *J. Org. Chem.* **1986**, *51*, 3065–3067.
- [5] a) G. A. Molander, E. R. Burkhardt, P. Weinig, *J. Org. Chem.* **1990**, *55*, 4990–4991; b) G. A. Molander, A. M. Estévez-Braun, *Bull. Soc. Chim. Fr.* **1997**, *134*, 275–282; c) G. A. Molander, C. Köllner, *J. Org. Chem.* **2000**, *65*, 8333–8339; d) D. Johnston, N. Francon, J. J. Edmonds, D. J. Procter, *Org. Lett.* **2001**, *3*, 2001–2004; e) D. Johnston, E. Couché, D. J. Edmonds, K. W. Muir, D. J. Procter, *Org. Biomol. Chem.* **2003**, *1*, 328–337; f) G. A. Molander, D. J. St Jean, J. Haas, *J. Am. Chem. Soc.* **2004**, *126*, 1642–1643. See the Supporting Information for additional references.
- [6] At the request of a reviewer, the reactivity of PhLi/CeCl₃ towards **1b** was examined and shown to favor C1 addition (Scheme 3).
- [7] Additions of *ortho*-, *meta*-, and hetero-arylytterbium reagents to **1b** are detailed in the Supporting Information along with further examples of organolithium additions to **1a** and **1b** and the thermal rearrangements of **21b** and **33**.
- [8] For a possible explanation of 2:1 stoichiometry required with alkyl Yb reagents see: S.-i. Fukuzawa, K. Mutoh, T. Tsuchimoto, T. Hiyama, *J. Org. Chem.* **1996**, *61*, 5400–5405, and references therein.
- [9] D. C. Harrowven, M. Mohamed, T. P. Gonçalves, R. J. Whitby, D. Bolien, H. F. Sneddon, *Angew. Chem.* **2012**, *124*, 4481–4484; *Angew. Chem. Int. Ed.* **2012**, *51*, 4405–4408.
- [10] DFT calculations were performed using B3LYP with a mixed basis set of MWB59 for Yb. Pre-optimization was carried out with MOPAC PM7/SPARKLE [a) J. J. P. Stewart, *J. Mol. Model.* **2013**, *19*, 1–32; b) J. D. L. Dutra, M. A. M. Filho, G. B. Rocha, R. O. Freire, A. M. Simas, J. J. P. Stewart, *J. Chem. Theory Comput.* **2013**, *9*, 3333–3334]. Single-point energies were calculated at the M06-2x/6-311+G(d,p)/MWB59 level [c) Y. Zhao, D. Truhlar, *Theor. Chem. Acc.* **2008**, *120*, 215–241] using the IEFPCM solvation model (THF) [d) S. Miertuš, E. Scrocco, J. Tomasi, *Chem. Phys.* **1981**, *55*, 117–129] on optimized geometries [e) M. J. Frisch, et al., Gaussian09 A. 02, see Supporting Information for full reference]. ChemCraft was used for visualization: G. A. Zhurko, <http://www.chemcraftprog.com>
- [11] A. Dzudza, T. J. Marks, *J. Org. Chem.* **2008**, *73*, 4004–4016.
- [12] This estimate is based on the analogous rotamers of the vinyl ether **1b**, which differ in energy by 2.6 kcal mol^{−1} (see the Supporting Information).
- [13] a) G. B. Marini Bettolo, C. G. Casinovi, C. Galeffi, *Tetrahedron Lett.* **1965**, *6*, 4857–4864; b) E. Garcia, V. Mendoza, J. A. Guzman, *Nat. Prod. Lett.* **1997**, *11*, 67–72.
- [14] R. H. Shapiro, M. F. Lipton, K. J. Kolonko, R. L. Buswell, L. A. Capuano, *Tetrahedron Lett.* **1975**, *16*, 1811–1814. For a review see A. R. Chamberlin, S. H. Bloom, *Org. React.* **1990**, *39*, 1–83.