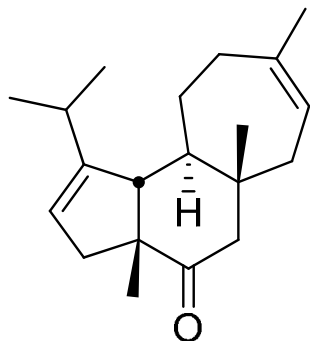


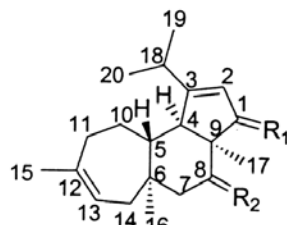
The Total Synthesis of (-)-Cyanthiwigin F by Means of Double Catalytic Enantioselective Alkylation



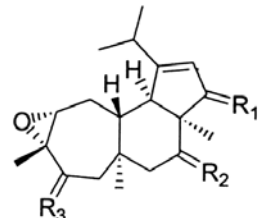
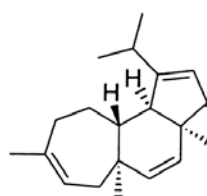
John A. Enquist Jr & Brian M. Stoltz
Nature **2008** *453* 1228-1231

Current Literature
Chenbo Wang @ Wipf Group
July 19th, 2008

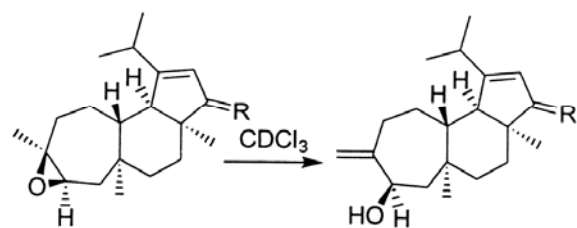
Cyanthiwigin F



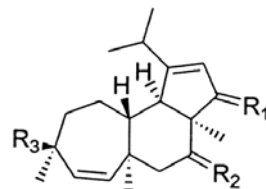
cyanthiwigin A (1): R₁=O R₂=H
 cyanthiwigin B (2): R₁=R₂=O
 cyanthiwigin C (3): R₁= α -OH R₂=H
 cyanthiwigin D (4): R₁= α -OH R₂= β -OH
 cyanthiwigin E (5): R₁=O R₂= β -OH
 cyanthiwigin F (6): R₁=H R₂=O



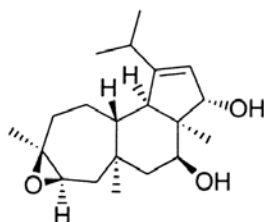
cyanthiwigin M (13): R₁= α -OH R₂=H R₃= β -OH
 cyanthiwigin N (14): R₁=O R₂=O R₃= β -OH
 cyanthiwigin O (15): R₁=O R₂=O R₃=O
 cyanthiwigin P (16): R₁=O R₂=H R₃= β -OH
 cyanthiwigin Q (17): R₁=O R₂=H R₃=O



cyanthiwigin H (8): R=O
 cyanthiwigin K (11): R= α -OH
 cyanthiwigin I (9): R=OH
 cyanthiwigin L (12): R= α -I



cyanthiwigin R (18): R₁=O R₂=O R₃=OOH
 cyanthiwigin S (19): R₁=O R₂=O R₃=OH
 cyanthiwigin T (20): R₁=O R₂= β -OH R₃=OOH
 cyanthiwigin U (21): R₁=O R₂=H R₃=OH
 cyanthiwigin V (22): R₁= α -OH R₂=O R₃=OH
 cyanthiwigin W (23): R₁= α -OH R₂=H R₃=OH
 cyanthiwigin X (24): R₁= α -OH R₂= β -OH R₃=OH

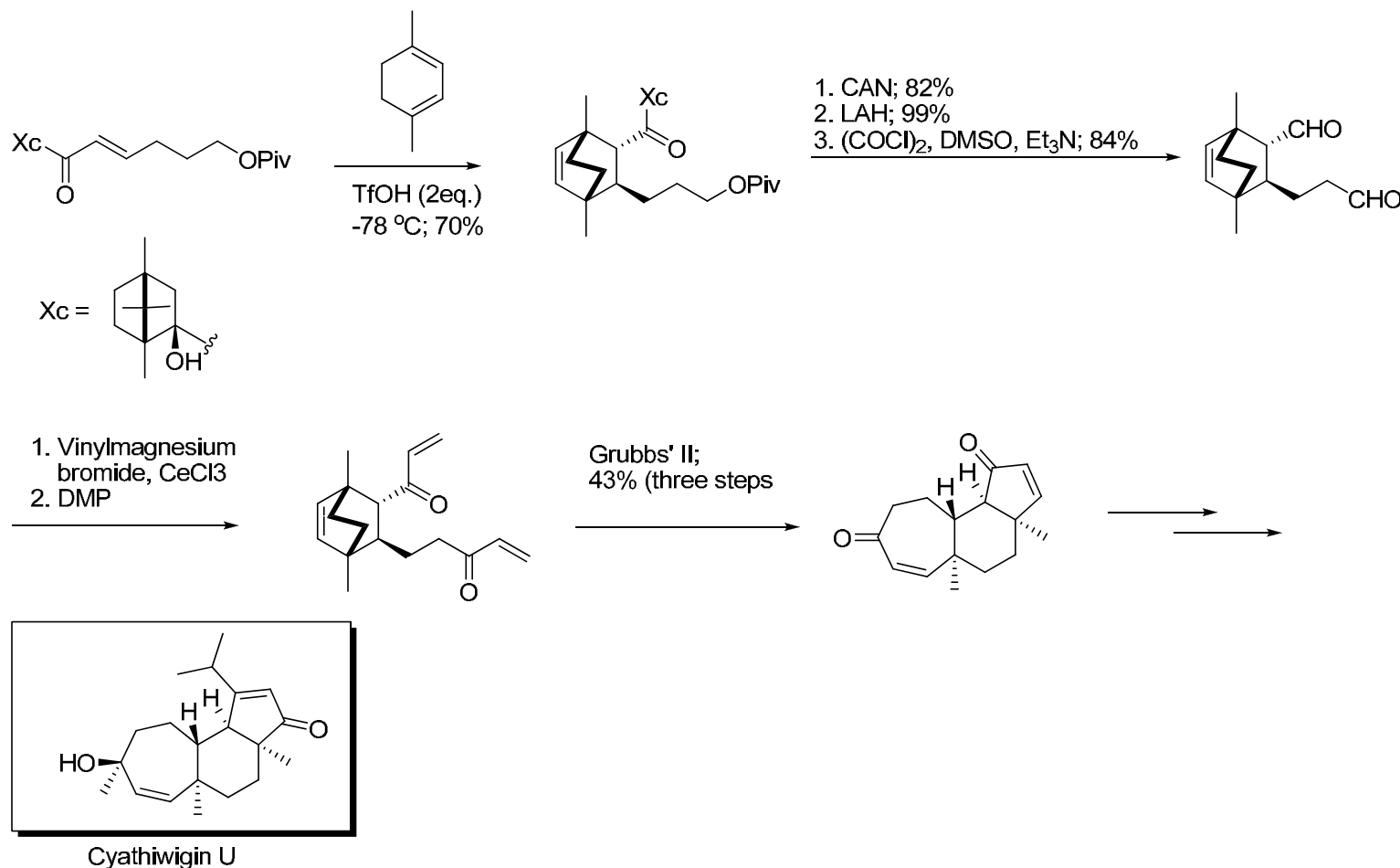


cyanthiwigin J (10)

Peng, J. et al. *Tetrahedron* 2002 58 7809

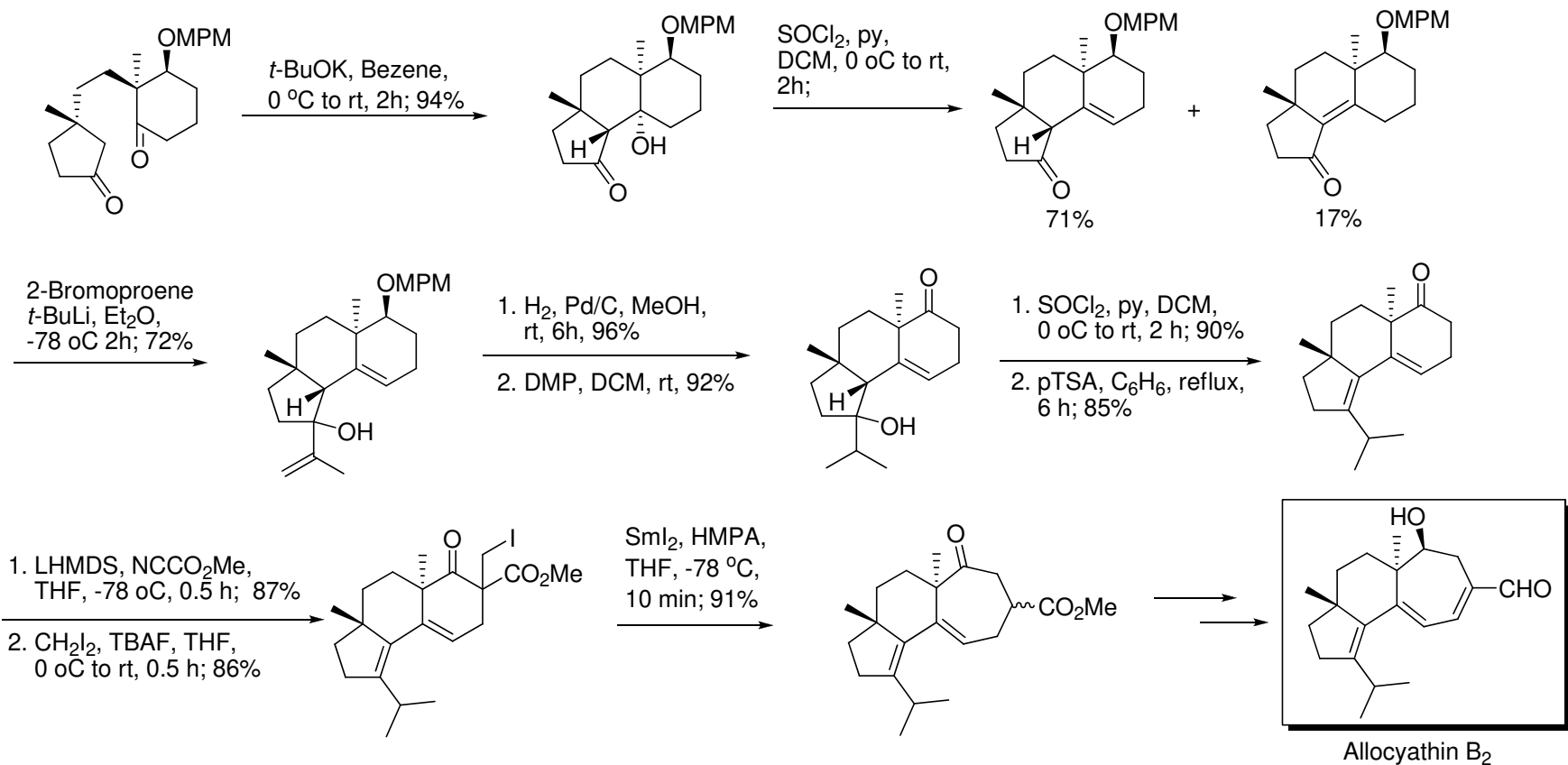
- Isolated from the Jamaican sponge *Myrmekioderma styx* in 2002.
- One of the 30 known cyanthiwigin natural products, all of which belong to cyathins.
- Cytotoxic against human primary tumour cells (with a half-maximal inhibitory concentration of 3.1 μ g ml⁻¹).

Cyanthin Synthesis: Previous Works



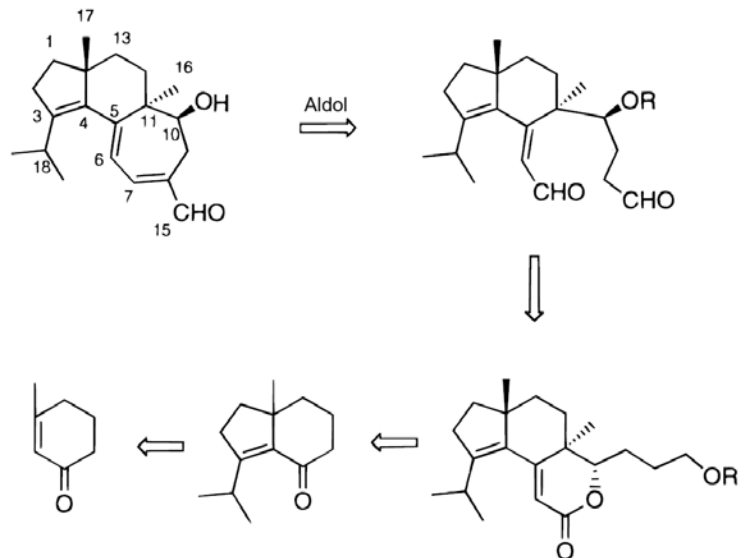
Philips, A. J. et al *J. Am. Chem. Soc.*, 2005 127, 5334

Cyanthin Synthesis: Previous Works

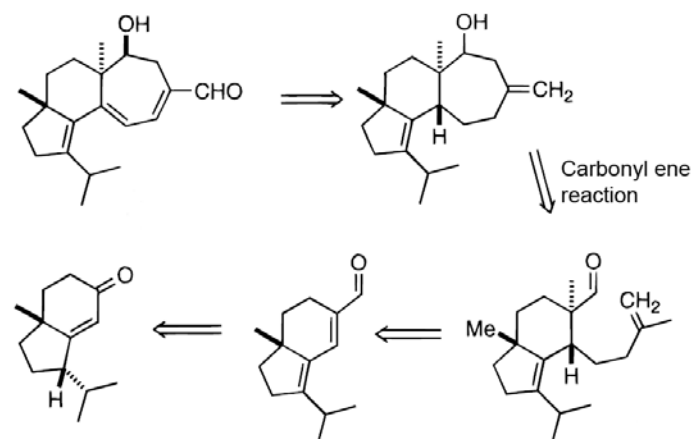


Nakada, M. et al *Org. Lett.* **2004**, 6, 4897.

Cyanthin Synthesis: Previous Works



- Allocyathin B2



- Allocyathin B2

Tori, M. et al. *J. Org. Chem.* **1998**, *63*, 306.
Snider, B. B et al *J. Am. Chem. Soc.* **1996**, *118*, 7644.

The Enantioselective Tsuji Allylation

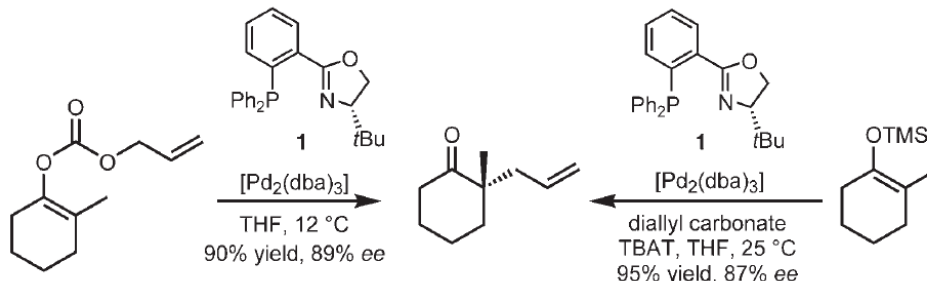


Table 2. Enantioselective Tsuji Enol Carbonate Allylation^a

entry	substrate	product	time (h)	% yield ^b	% ee ^c
1	OCO ₂ allyl		2	85	87
2 ^d			5	85	88 (96) ^e
3'			9	90	89
4	OCO ₂ allyl		2	96	92
5 ^g			10	55 ^h	82
6			2	96	85
7			2	87	88
8 ^g	OCO ₂ 2'-Me-allyl		8	89	91
9	OCO ₂ allyl		1	94	92
10			1	87	86
11			1	91	89
12'			2	87	91
13'			8	94	91
14			n = 1	81	87
15			n = 2	90	79

Table 3. Enantioselective Tsuji Enol Silane Allylation^a

entry	substrate	product	time (h)	% yield ^b	% ee ^c	
1			R = CH ₃	2	95	87
2			R = CH ₂ CH ₃	3	96	92
3 ^d				4	79	91
4				2	99	81
5			n = 1	2	94	86
6			n = 2	3	96	79

Behenna, D. C. & Stoltz, B. M. *J. Am. Chem. Soc.* **2004** 126, 15044

Pd-Catalyzed Decarboxylative Allylation of β -Ketoesters

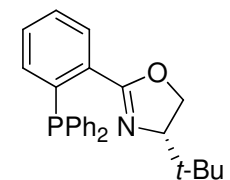
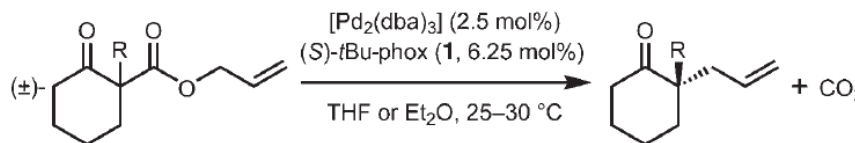
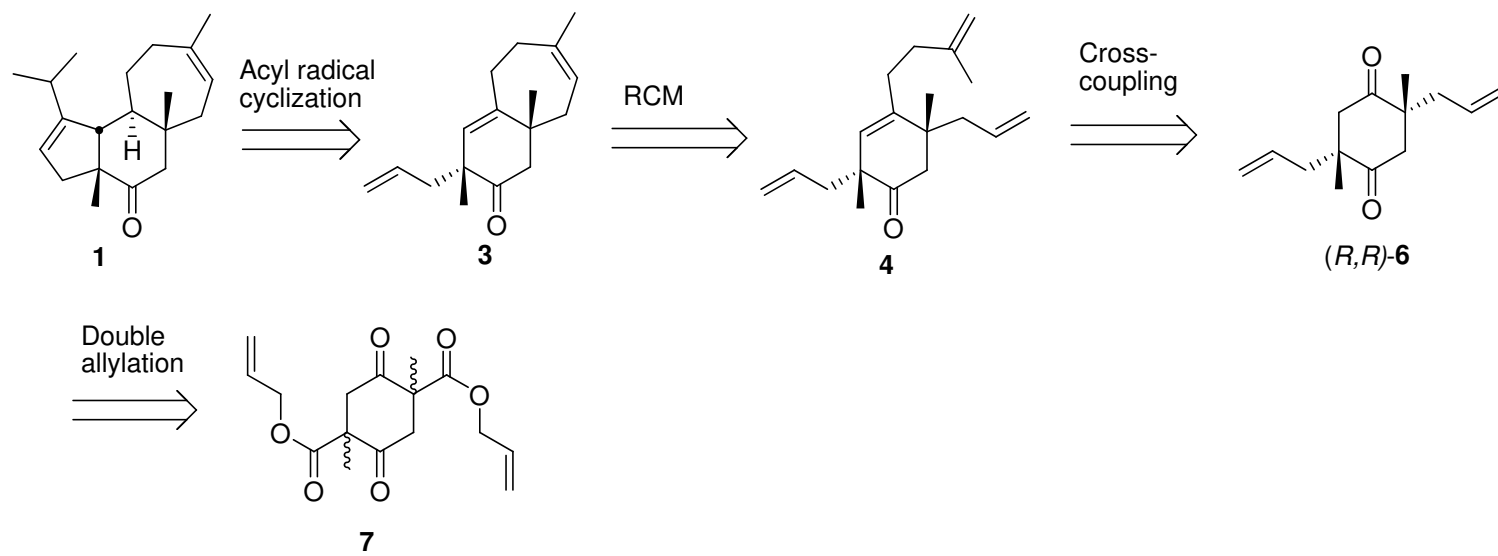


Table 2: Enantioconvergent decarboxylative allylation of β -ketoesters.

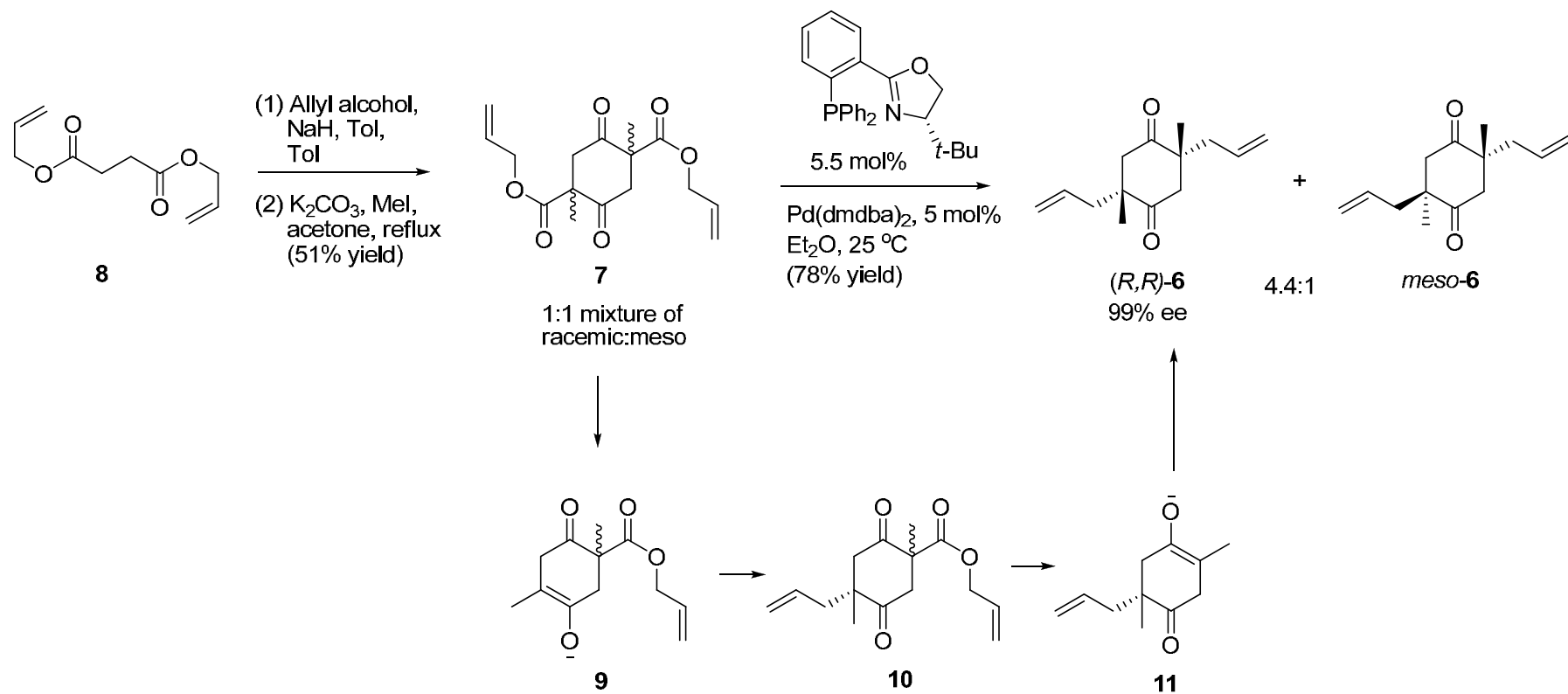
Entry	Substrate	Product	T [°C]	t [h]	Yield [%] ^a	ee [%] ^b
1 ^c			25 25	1.5 24	94 94	85 86
3			30	9	89	90
4			25	5	90	85
5 ^{d,e}			30	4	77	90
6 ^d			25	10	97	92
7			25	9.5	83	87
8 ^d			35	6.5	87	92
9 ^{d,e}			35	2.5	87	91
10			25	2.5	91	92

Stoltz, B. M. et al *Angew. Chem. Int. Edn Engl.* 2005 44, 6924

Title Paper: Retrosynthesis



Double Catalytic Enantioselective Stereoab ablative Allylation



Catalytic Enantioselective Stereoab ablative Reactions

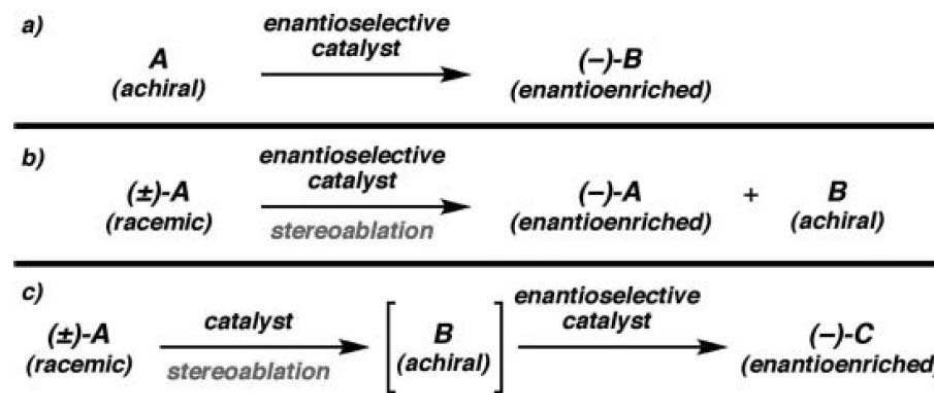
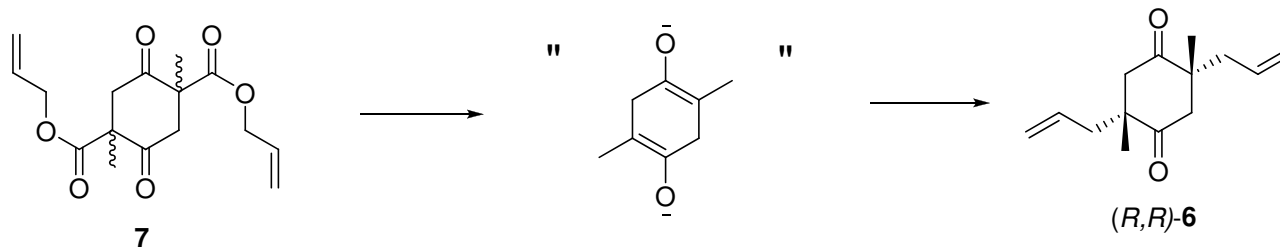
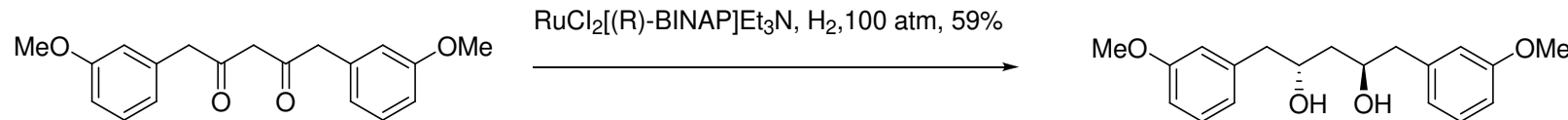


Fig. 1 Strategies for enantioselective catalysis.

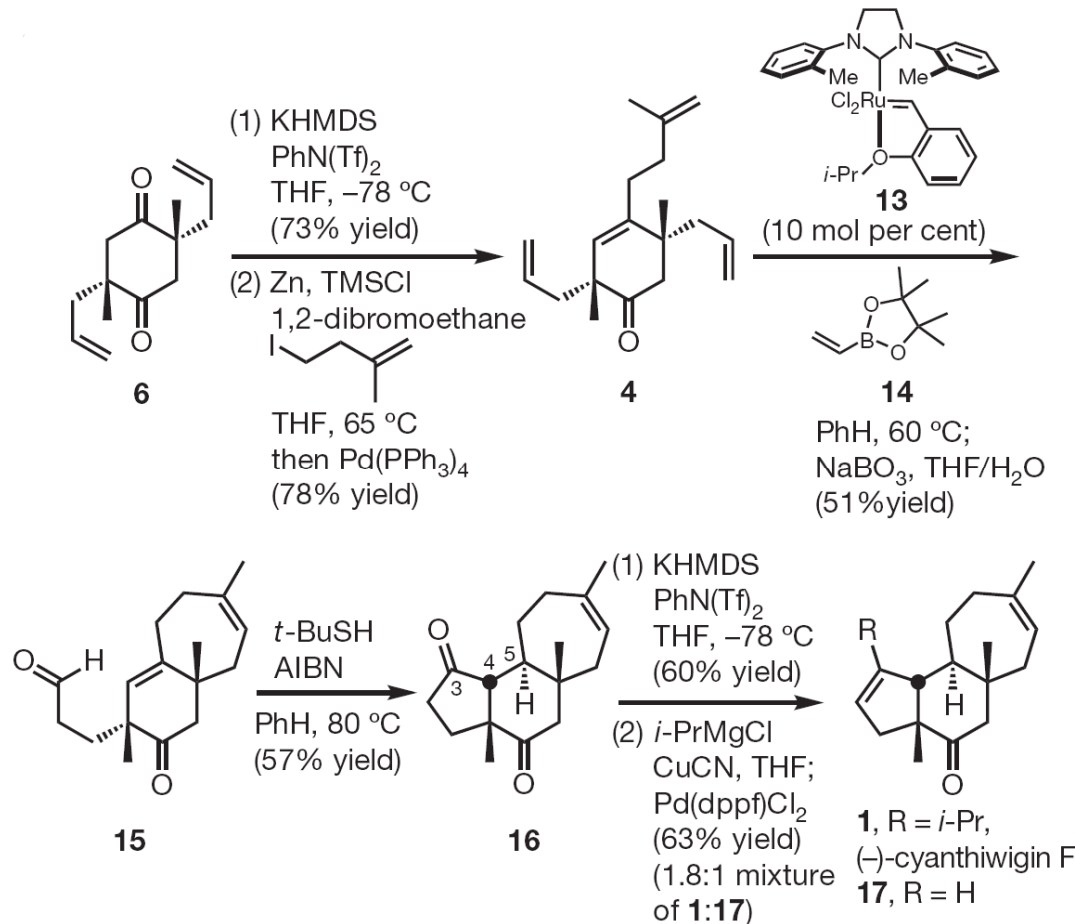


vs.



Stoltz, B. M. et al *Org. Biomol. Chem.*, 2007, 5, 3571
Schreiber, S. L. et al *J. Am. Chem. Soc.* 1993, 115, 3360

Completion of The Synthesis



Summary

- The total synthesis of (-)-cyanthiwigin F was accomplished in 10 steps, 1.2% overall yield
- Key steps include a double catalytic enantioselective allylation, RCM and acyl radical cyclization.
- The synthesis is protection-group-free.