

Total Synthesis of the Hsp90 Inhibitor
Geldanamycin

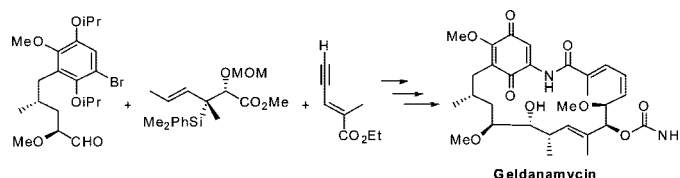
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ABSTRACT



An enantioselective synthesis of the Hsp90 inhibitor geldanamycin was achieved in 20 linear steps and 2.0% overall yield from 2-methoxyhydroquinone. The synthesis is highlighted by a regio- and stereoselective hydroboration reaction; a $\text{Sc}(\text{OTf})_3/\text{Et}_3\text{SiH}$ -mediated pyran ring-opening reaction; an enantioselective crotylation to simultaneously install the C8–C9 (*E*)-trisubstituted olefin, the C10 and C11 stereocenters; a chelation-controlled asymmetric metallated acetylide addition; and an intramolecular copper(I)-mediated aryl amidation reaction to close the 19-membered macrolactam.

Heat shock protein 90 (Hsp90) is a molecular chaperone that regulates folding, transport, and degradation of client proteins. It also plays a key role in the conformational maturation of oncogenic signaling proteins.¹ Hsp90 ATPase binding region's

role in cancer and protein maintenance as well as its broad range of functions make it a significant therapeutic target for anticancer drug development.^{1d,e}

Geldanamycin, a natural product isolated from *Streptomyces hygroscopicus* var. *geldanus* var. *nova* in 1970,² is the first reported Hsp90 inhibitor. It is currently under development as a therapeutic agent for cancers associated with abnormally elevated levels of receptor tyrosine kinase activity.^{3a} Studies have shown that the Hsp90 client proteins can be destabilized when geldanamycin binds to the ATP-binding site of Hsp90 and inhibits the chaperone activity of the protein.³ Accordingly, several geldanamycin analogues are in various stages of

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development as novel antitumor agents and as chemotherapeutic agents in a number of diseases.⁴

Geldanamycin, together with herbimycin A,⁵ macbecin I,⁶ and reblastatin,⁷ are members of the ansamycin class of natural products (Figure 1). Geldanamycin differs at C6, C11, C15,

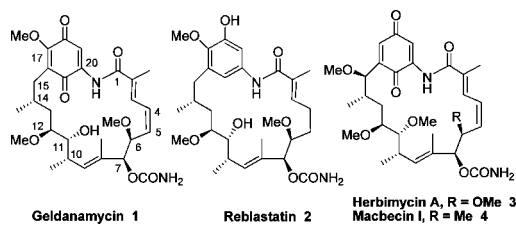
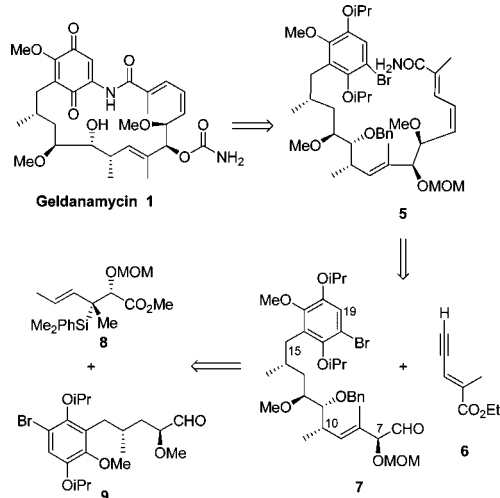


Figure 1. Geldanamycin and related ansamycin antibiotics.

and C17 from macbecin I and herbimycin A. Presently, one total synthesis of geldanamycin has been described.⁸ Herein, we report the total synthesis of geldanamycin as part of our ongoing studies of ansamycins.

Our strategy for the synthesis of geldanamycin (**1**) is shown in Scheme 1. We envisioned the 19-membered macrocycle

Scheme 1. Retrosynthetic Analysis of Geldanamycin



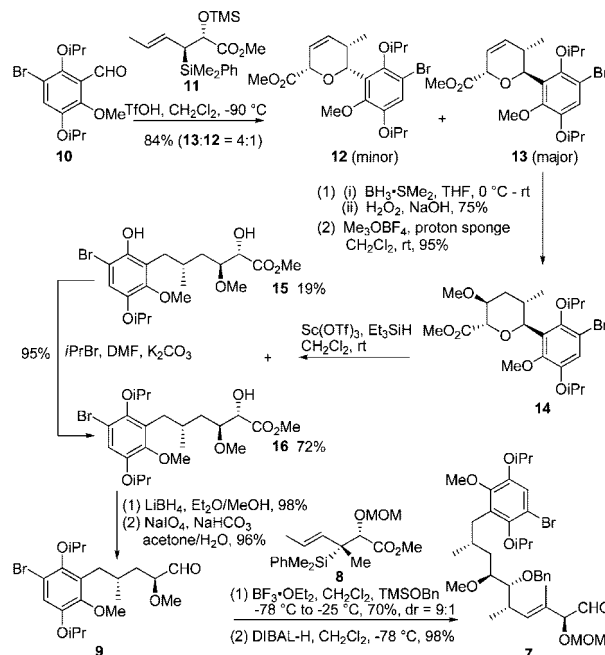
would be formed from the acyclic amide **5** through an intramolecular aryl amidation reaction similar to that described in our synthesis of reblastatin.^{7,9} The (*E,Z*)-diene could be installed by reduction of the enyne from alkylation of precursors **6** and **7** which could be generated from easily accessible aldehyde **9** and chiral silane reagent **8**. Organosilane **8** has unique features as it establishes the C10–C11 syn stereo-

chemistry while simultaneously creating the C8–C9 (*E*)-trisubstituted olefin.

Recently, we have reported the selective hydridic opening of aryl *C*-glycosides using a Sc(OTf)₃/Et₃SiH-mediated reduction of the benzylic C–O σ -bond resulting in the formation of a stereochemically well-defined acyclic system.¹⁰

The synthesis of the C6–C21 fragment **7** (Scheme 2) started from functionalized aromatic aldehyde **10**, which was easily

Scheme 2. Synthesis of Aromatic C6–C21 Fragment **7**



obtained in three steps from commercially available 2-methoxyhydroquinone in 55% overall yield. [4 + 2]-Annulation of this aldehyde with silane **11** provided a mixture of dihydropyrans **12** and **13** in 84% yield (*trans/cis*: 4:1).¹¹ Regio- and stereoselective hydroboration of the pyran double bond provided the secondary alcohol, which was subsequently methylated with Meerwein's reagent to provide tetrahydropyran **14** in good yield and excellent selectivity (dr > 20:1). Reductive opening of pyran **14** with Sc(OTf)₃/Et₃SiH gave a mixture of **15** (which was easily converted to **16** in 95% yield) and **16** in excellent yield, essentially deoxygenating C15 (cf. macbecin). Reduction of ester **16** with LiBH₄ afforded the 1,2-diol, which was subjected to oxidative cleavage using sodium periodate to furnish the α -methoxyaldehyde **9**.

At this point, we turned toward the generation of the C7, C10, and C11 stereocenters and the installation of the trisubstituted alkene. Gratifyingly, after a variety of conditions were explored, a double-stereodifferentiating crotylation¹² of silane **8** and aldehyde **9** promoted by BF₃·OEt₂ provided benzyloxy

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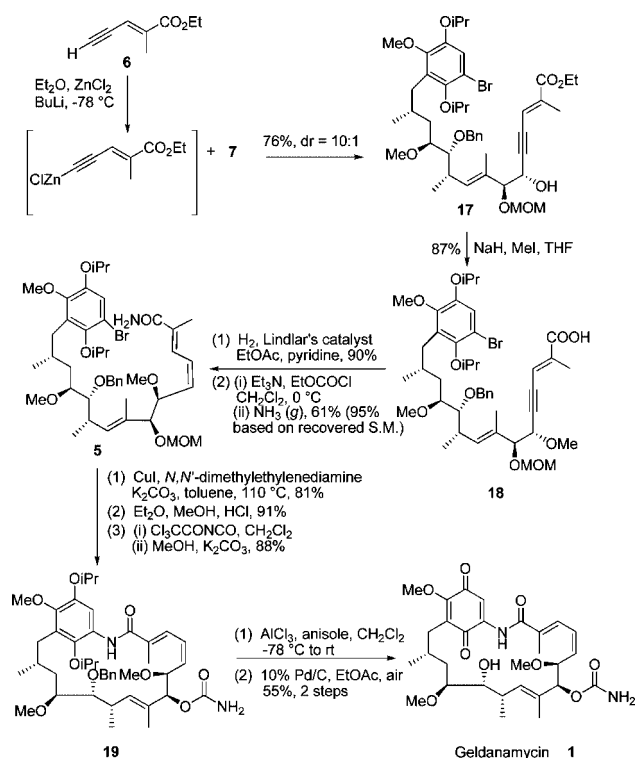
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homoallylic ether bearing not only the three desired stereocenters and trisubstituted alkene, but also a benzylic ether protecting group at C11. The reaction proceeded in 70% yield with high selectivity (dr = 9:1). Reduction of the product methyl ester with DIBAL-H yielded aldehyde 7.

To access (*E,Z*)-unsaturated macrocycle **19** through the intermediacy of bromo ester **17**, aldehyde **7** was homologated through diastereoselective addition¹³ of a metallated acetylide (Scheme 3). Accordingly, fragment **17** was obtained through

Scheme 3. Completion of the Total Synthesis of Geldanamycin



chelation-controlled coupling of acetylene **6** and aldehyde **7** in 76% yield (dr = 10:1).^{14,15}

To complete the synthesis, methylation of propargylic alcohol **17** using NaH and MeI occurred with simultaneous dealkylation of the ester to the corresponding acid¹⁶ and gave the enyne-acid **18** in 87% yield, which was subjected to Lindlar reduction.¹⁷ The resulting acid was subsequently converted to the (*E,Z*)-unsaturated amide **5**. The amide **5** was then subjected to an

intramolecular copper(I)-mediated aryl amidation reaction to provide the desired macrocyclic lactam in 81% yield. Deprotection of the MOM ether was achieved utilizing anhydrous $\text{HCl}/\text{Et}_2\text{O}/\text{MeOH}$ to obtain the secondary alcohol in nearly quantitative yield without affecting other protecting groups. The secondary alcohol was then converted to the desired carbamate **19** in 88% yield.^{7,18} Finally, deprotection of both the diisopropyl and benzyl ethers was accomplished with AlCl_3 in the presence of anisole¹⁹ to generate dihydrogeldanamycin, which was immediately treated with catalytic palladium on carbon²⁰ (10%) under air atmosphere to give geldanamycin in 55% yield over two steps. The spectroscopic data obtained for synthetic material were in agreement with those for authentic geldanamycin (optical rotation, ^1H and ^{13}C NMR, IR, and HRMS).²¹

In summary, we achieved the total synthesis of geldanamycin in 20 linear steps and 2.0% overall yield from commercially available 2-methoxyhydroquinone. Notable features of our synthetic route include the following: a concise synthesis of the C11–C21 fragment through reductive pyran-opening approach; an efficient and selective crotylation with silane **8** that simultaneously set two stereocenters (C10, C11), created an (*E*)-trisubstituted olefin, and put the stereocenter C7 in place; an asymmetric alkynylation to install stereocenter C6; and the first example of synthesis of the *E,Z*-diene of ansamycins through Lindlar reduction of the enyne precursor.²² An intramolecular copper(I)-mediated amidation reaction was used to close the 19-membered macrolactam.

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Supporting Information Available: Experimental details and selected spectral data for all new compounds. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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