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Synthesis of ¹⁵N-labeled vicinal diamines through N-activated chiral aziridines: tools for the NMR study of platinum-based anticancer compounds

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ABSTRACT

A new method for the synthesis of 15 N-labeled chiral β -diamines from a common precursor, either optically pure amino acids or *anti*-β-amino alcohols, is reported. The two diastereomeric series of vicinal diamines are produced through the nucleophilic ring opening of activated chiral aziridines. ¹⁵N was introduced by means of [¹⁵N]-benzylamine, prepared from ¹⁵NH₄Cl. The final compounds are highly valuable because [¹H-¹⁵N] NMR is considered a powerful tool for studying the chemical properties of platinum-based complexes.

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The β -diamine (or 1,2-diamine) moiety is of great importance in medicinal chemistry, especially in its chiral form. This structure is particularly well represented in anticancer platinum(II) complexes.¹ Since the discovery of *cis*-diamminedichloroplatinum(II) (Cisplatin) by Rosenberg et al. in the late 1960s,² the continuous search for new compounds is driven, on the one hand, by the limitations of current platinum-containing drugs (acute toxicity, resistance, limited activity spectrum, and poor pharmacokinetics)³ and, on the other hand, by the high efficacy of these drugs against various cancers.⁴ The biological activity and the chemical properties of Pt compounds (i.e., hydrolysis kinetics, dissociation constants, reactivity with bionucleophiles, proteins, nucleic acids) are closely related. Therefore, tailoring these structure-dependent chemical properties is the basis for the rational design of platinum drugs and is essential to provide new leads with better profiles. Moreover, the importance of chirality to the biological properties of these complexes has largely been demonstrated.⁵

Nuclear Magnetic Resonance (NMR) methods, especially ¹⁹⁵Pt and ¹⁵N spectroscopy, have proven to be highly useful for the characterization of platinum compounds.⁶ The introduction of [¹H,¹⁵N] NMR techniques, in which the sensitivity is enhanced through polarization transfer,⁷ allowed fine studies on the behavior of platinum anticancer derivatives in aqueous solution.^{8,5f} However, sensitivity remains a major issue at natural abundance of ¹⁵N (0.365%). The possibility of synthesizing ¹⁵N-enriched optically pure vicinal diamines is therefore crucial for the understanding of the structure-activity relationships of platinum coordination complexes. Only quite simple complexes were studied until now by ¹⁵N NMR and few publications actually mention the preparation of ¹⁵N-enriched diamines. Indeed, reported achiral ligands are: alkyl amine derivatives,⁹ polynuclear complexes,¹⁰ diethylenetriamine compounds,¹¹ dipyridine coordinates,¹² diimines,¹³ and nitroimidazole compounds.¹⁴ A platinum(IV) chiral compound, using propane-1,2-diamine as the ligand, was reported by Drahoňovský et al.15a

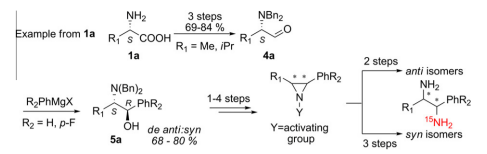
Based on their biological and structural properties, seven previously synthesized platinum compounds^{5b-d} were selected for ¹⁵N labeling prior to a comprehensive NMR study of their chemical properties. A new synthetic method was developed for the introduction of ¹⁵N in the target compounds.

Commercially available, optically pure α -amino acids, were used as starting compounds. β-amino alcohols **5** are first produced as described by Reetz et al.¹⁶ and then converted into vicinal diamines with the introduction of ¹⁵N (Scheme 1). Alternative synthetic methods, as the organometallic addition to imines derived from $\mathbf{4}^{17}$ and the diastereoselective reduction of 1,2-diimines¹⁸ were assayed but delivered unsatisfactory results. Because the choice of reagents is considerably limited by the need for isotopically enriched compounds, various pathways were considered for the conversion of amino alcohols 5 into diamines. The few publications about ¹⁵N-enriched chiral diamines recommended CH₃-¹⁵NH₂ to open a chiral aziridinium ion obtained from ephedrine or ¹⁵N-enriched amino acids, which are very expensive.¹⁵ Asymmetric synthesis through aziridines was demonstrated to be



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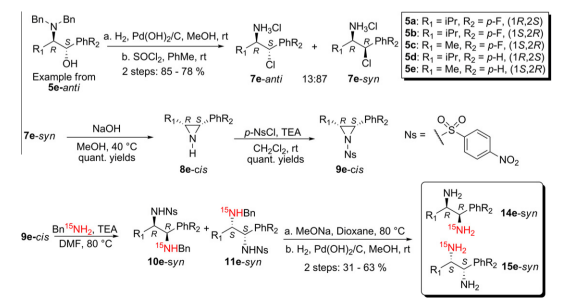
Scheme 1. Overall scheme for the production of 15 N-labeled diamines from α -amino acids.

highly efficient for the preparation of nitrogen-containing molecules and, among these, the vicinal diamine moiety is readily obtained.¹⁹ The stereo- and regio-controlled opening of chiral aziridines was thus finally selected and two distinct pathways were used to produce *syn*- and *anti*-configured diamines.

Syn-diamines were obtained from the nosyl-activated aziridines 9. which were produced from amino alcohols 5 in a four step synthesis (Scheme 2). Nosyl groups allow the very efficient activation of aziridines, comparable to that of aziridiniums with nucleophiles.²⁰ Other activation methods, especially the use of Lewis acids,²¹ were not successful. Simple protonation of the aziridine was not possible, as aziridines are far less basic than benzylamine.²² Amino alcohols 5 were first debenzylated by treatment with $Pd(OH)_2/C$ under H_2 atmosphere (1 atm), leading to unprotected β-amino alcohols. In the second step, their chlorination using thionyl chloride afforded the amino chlorides 7 (as the hydrochloride salts) with a diastereomeric excess of the syn compound; this reagent is indeed known to produce an inversion of configuration in the case of *anti*-configured β-amino alcohols.²³ This diastereomeric mixture was either purified by recrystallization or engaged in the next step.²⁴ Chlorinated compounds **7** were then cyclized to aziridines 8 under basic conditions, by deprotonation of the hydrochloride salt. After reacting with nosyl chloride to produce nosyl-azirdines 9, these were easily converted into ¹⁵N-labeled syn-diamines through nucleophilic ring-opening by reaction with [¹⁵N]-benzylamine (prepared from ¹⁵NH₄Cl).²⁵ This procedure stereospecifically afforded the syn isomers as the reactions proceeded exclusively through a S_N2 mechanism, but

produced both regioisomers in the case of Me substituted aziridines. Indeed, if benzvl substituted aziridines are known to be regioselectively opened at the benzylic position,²⁶ in the case of nosyl-activated aziridines **9c** and **9e** (Me substituted aziridines). the preferred position for nucleophilic opening was shifted to the alkyl-substituted carbon C₃ affording both the regioisomers **10** (C₂ opening) and **11** (C₃ opening) in a 35:65 ratio. However, compounds 10 were the only regioisomers obtained upon opening of the bulkier *i*-Pr substituted aziridines **9a**. **9b**. and **9d**. possibly indicating a kinetically unfavorable attack on C₃, due to steric hindrance. All regioisomeric products were separated and identified by the mean of [¹H ¹³C] 2D-NMR experiments.²⁷ Sulfonamide cleavage upon MeONa treatment²⁸ and catalytic hydrogenation resulted in the final labeled β-diamines. Thiolated nucleophiles (PhSH, thioglycolic acid, mercaptoethanol), better known for their nosyl cleavage ability,^{20a,29} gave unsatisfactory results (i.e., poor vields and/or sluggish reaction rates).

anti-Diamines were synthesized from **5** through a chiral dibenzylaziridinium intermediate (**16**), ^{5a,19b} which was regiospecifically opened at the benzylic position by [¹⁵N]-benzylamine to give **18** after hydrogenation (Scheme 3). Indeed, aziridinium **16** showed a clear preference for nucleophilic attack at the benzylic position, and diamines **17** were the only stereo- and regio-isomeric observed products. Anyway, the presence of a by-product resulting from the opening of dibenzylaziridinium **16** by chloride ions needed products **17** to be purified by flash chromatography prior to catalytic hydrogenation. For pharmacological reasons (i.e., much less active platinum(II) complexes), only two compounds of the



Scheme 2. Conversion of **5** to ¹⁵N-labeled syn-β-diamines.

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