



2-Quinoxalinol salen ligands incorporated into functionalized resins for selective solid-phase extraction of copper(II)

Xianghong Wu, Anne E. V. Gorden *

Department of Chemistry and Biochemistry, College of Science and Mathematics, 179 Chemistry Building, Auburn University, Auburn, AL 36849-5319, USA

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ABSTRACT

In exploring selective extraction systems for use in environmental remediation or in metal scavenging agents for use in combinatorial chemistry, a novel reagent for the selective extraction of copper(II) has been developed. 2-Quinoxalinol salen ligands supported on an aminomethyl-polystyrene resin has been shown to efficiently and selectively extract copper(II) ions from organic solvents within 30 min under a variety of experimental conditions. Mild reducing conditions allow for metal ion recovery.

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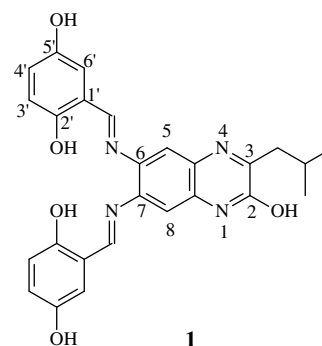
Solid-phase extraction (SPE) technologies are being used in a wide number of areas including environmental chemistry, medicinal chemistry, combinatorial chemistry, agriculture, and food science. This technology has been used in environmental applications to monitor or extract toxic heavy metal cations such as Hg^{2+} , Pb^{2+} , and As^{3+} .^{1–7} Others have applied solid-phase micro-extraction (SPME) techniques to analyze biological fluids in diagnostic medicine.⁸ In food science, SPE technologies are used extensively in food analysis and quality control.^{9–12} Currently, the most common application of SPE technology in use is in the identification of drug components.^{13,14} One promising area of SPE technology not as widely investigated is as a scavenging agent to remove excess catalysts or metal ions used in synthetic methods or combinatorial chemistry.^{15–17}

Copper salts are often applied as catalysts in a variety of organic synthetic procedures. An overabundance of copper in the environment is of concern due to possible harmful effects to agriculture, fish, or wildlife.^{18–22} SPE technology for the selective recovery of copper has not previously been described in the literature. Such technology has considerable potential and would allow many catalytic or coupling reactions commonly used in the synthetic laboratory to be more accessible for use in combinatorial processes or in industrial applications, where excess or trace metal ions are likely to complicate subsequent reactions. In addition, because of the high cost of copper reagents, this would create a more efficient and more readily recyclable system thereby limiting the need for costly copper reagents.

In our previous research, both symmetric and unsymmetric 2-quinoxalinol salen ligands have been prepared (see structure **1** as an example).^{23,24} It has been found that these ligands can coordinate +2 metal cations. This made us consider the possibility of incorporating these 2-quinoxalinol salen ligands into solid-phase

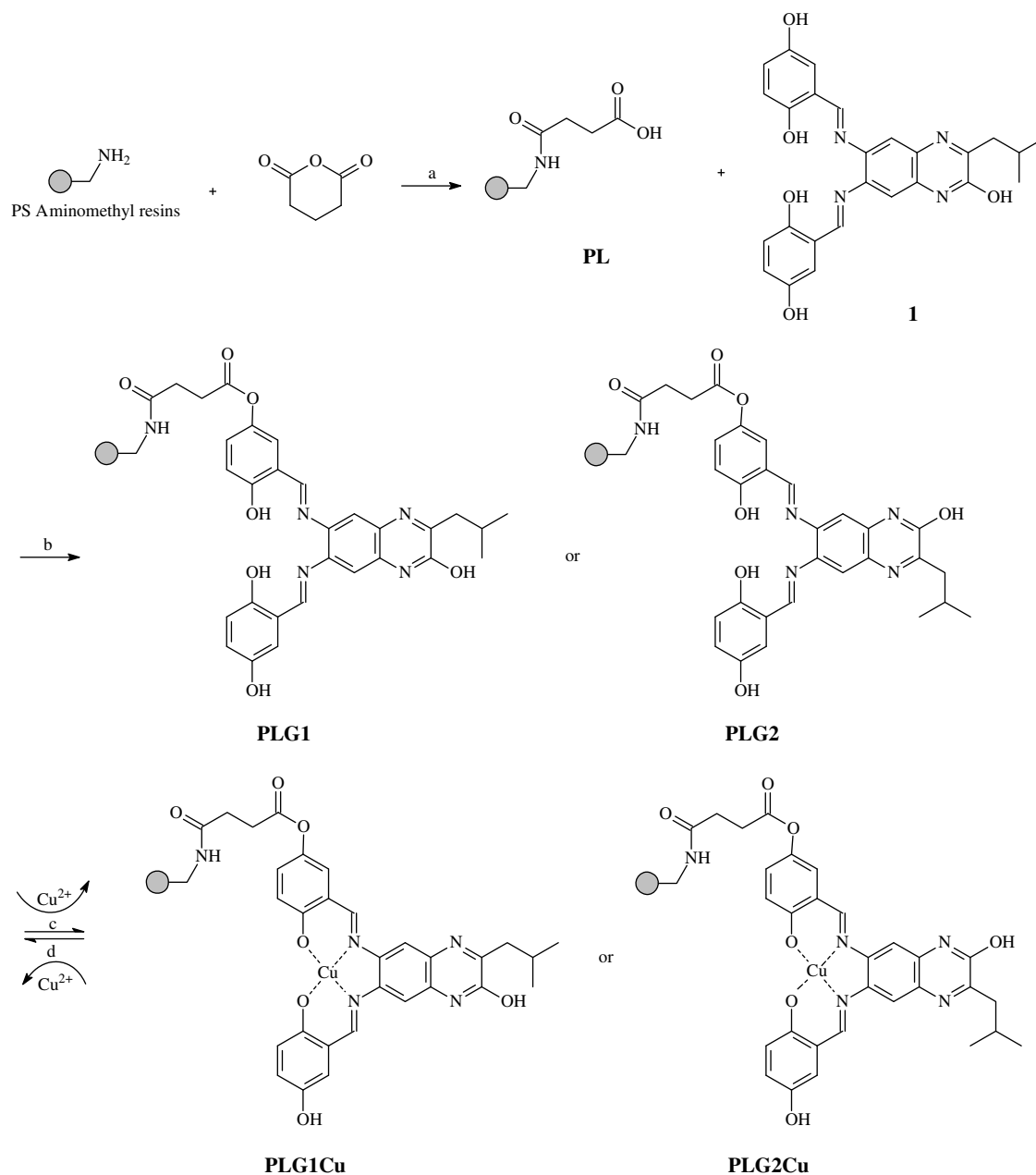
resins. Here, we will detail the methods of preparing solid-phase reagents and their application in the selective extraction and recovery of copper cations.

Polystyrene (PS) aminomethyl resin was selected as solid carrier, because this resin can swell in many organic solvents. Glutaric anhydride was selected as linker between solid carrier and 2-quinoxalinol salen ligand. The isopropyl-2-quinoxalinol salen ligand (**1**) was selected as the coordination ligand, both because the 5'-hydroxyl group on the outer portion of the salen is a convenient site for incorporating the resin, and because its yield is the highest of the symmetric library.²³



Presented in Scheme 1 is the optimized method for the synthesis of solid reagents PLG1 or PLG2. Different molar ratios and various solvents were investigated to optimize the acylation of the amino group onto the (PS) aminomethyl resin. Finally, a molar ratio of glutaric anhydride to (PS) aminomethyl resin 5:1 using dichloromethane as solvent at room temperature for 24 h was found to be the optimum conditions for acylation of the amino group. After acylation was completed, the resin was washed with DCM, DMF, and MeOH, three times each. The loaded resin PL can be further acylated with 1.5 equivalent of ligand **1** by using 2.2 equivalents of 4-dimethyl-aminopyridine (DMAP) and 2.2 equiv

* Corresponding author. Tel.: +1 334 663 3223; fax: +1 334 844 4043.
E-mail address: gordeae@auburn.edu (A. E. V. Gorden).



Scheme 1. Reaction route for solid reagents PLG1 and PLG2. Reagents and conditions: (a) 5 equiv, glutaric anhydride, DCM, rt, 24 h; (b) 2.2 equiv, DMAP, 2.2 equiv, DIC, DMF, rt, 72 h; (c) Cu^{2+} , DCM, MeOH, rt, 30 min; (d) $\text{NaBH}(\text{OAc})_3$, DCM, AcOH, rt, 4 h.

of *N,N'*-diisopropyl-carbodiimide (DIC) in distilled, dry DMF at room temperature for three days. Extending the reaction time does not increase loading. Without DMAP, the loading is very low. If wet DMF was used, the loading capability is decreased, because DIC can be decomposed easily by moisture.

There are three potential binding sites (the 2, 2', and 5' positions) for the carboxylic acid linker to attach to ligand **1**. Positions 2 and 2' cannot react with the carboxylic acid moiety due to reactive inertia of the 2 position and steric hindrance at the 2' position. Hydrogen bonding also limits the reactivity of the hydroxyl group on the 2' position.²⁵ Reaction with the hydroxyl groups in the two 5' positions results in two possible products (i.e., PLG1 and PLG2). They have the same coordination capability to bind +2 valence metal cations, because the salen coordination cavity is not affected by their position. By this procedure, PL was obtained with 100.0% loading by ninhydrin test, and PLG1 and PLG2 were obtained with

60.0–65.0% loading, as determined by mass. Identification of resins PLG1 and PLG2 was made by comparing the major peaks of an IR spectra of ligand **1** (3381.2, 1658.8, 1622.3, 1577.8, 1489.1, 1278.8 cm^{-1}).

With this solid ligand functionalized reagent in hand, extraction studies were run in several different solvent combinations: DMF, DMF/MeOH, MeOH, THF/MeOH, DCM/MeOH, and DCM/EtOH. Finally, we found that DCM/MeOH is the best combination for our extraction experiment, because the DCM can best swell the PS resins while MeOH still readily dissolves the metal salts. Metal salts of Cu^{2+} , Mn^{2+} , and Ni^{2+} were used in extraction studies. Solutions containing these metal ions were prepared using copper acetate, manganese acetate, and nickel nitrate salts in a 50:50 solution of DCM/MeOH. To determine extraction capability, 3 ml of a prepared solution of 2×10^{-4} mol/l Cu^{2+} was mixed with 5 mg, 10 mg, 15 mg, 20 mg prepared PLG1(**2**) (The molar ratio of Cu^{2+} to PLG1(**2**) is

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