## Synthesis of pyrrole derivatives of serinol for functionalization of carbon allotropes



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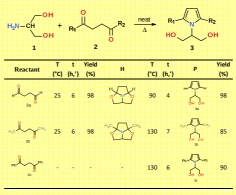
**Summary:** N-Pyrrole-based heterocycles are present in many natural products,[1] medicinal agents,[2] and functional materials,[3,4] therefore substantial attention has been paid to develop efficient methods for pyrroles synthesis.[5,6] Moreover, they are precursors for the synthesis of poly N-alkyl pyrroles which have wide ranging applications in electronics and sensors due to their tunable optoelectronic properties.

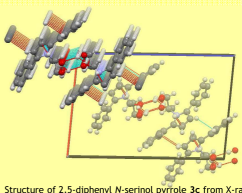
We present here one operationally simple, practical and economical Paal-Knorr pyrrole condensation of serinol (2-amino-propan-1,3-diol, 1) with dicarbonyl compounds 2 (and related precursors acetal/ketals or enolesters), under neat conditions in the absence of any catalysts, which allows the synthesis of *N*-serinolpyrrole derivatives 3 in good to excellent yield.

N-serinolpyrroles 3a,b were obtained in two steps from dicarbonyl compounds 2a and 2b through oxazolidine intermediates.

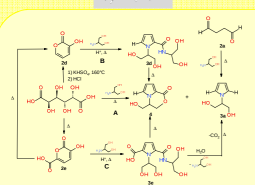
The first step occurs at room temperature while in the second step oxazolidines are converted into pyrroles by heating.

Pyrrole 3c was obtained in one step by heating compound 2c. In all cases solvents and catalysts were not used and yields were all good to excellent.





Structure of 2,5-diphenyl *N*-serinol pyrrole **3c** from X-ray. Highlighted is the hydrogen bond of 4 oxygen network which divides the high polar and unpolar framework.



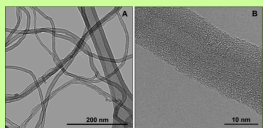
Scheme: Pathways to serinolpyrrol derivatives from mucic acid and its di-insaturated lactones.

Mucic acid (as all aldaric acid) has an interesting thermal reactivity: on heating this compound loses water forming the corresponding unsaturated lactone **2e** which on further warming can be converted by decarboxylation into the 3-hydroxy-2-pyrone (**2d**). Pyrones **2e** and **2d** can undergo a nucleophilic attack by amines on lactone carbonyl group opening the ring to the corresponding substituted 1,4-diketons.

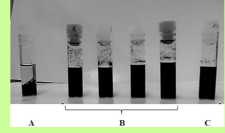
This reactivity was used to synthesize pyrroles **3a**, **3d**, **and 3e** by a mild and green process (50-70°C) using serinol (1) as nucleophilic agent (pathways **B** and **C**).

The pyrroles 4 and 3a can be obtained conveniently also heating a mixture of mucic acid and serinol at 160°C for 3 hours (pathway A).

Polyurethanes (PU) were synthesized from 2-(2,5-dimethyl-1*H*-pyrrol-1-yl)-1,3-propandiol (**3b**-SP) and hexamethylen-diisocyanate (HDI). Stable supramolecular interaction with multiwalled carbon nanotubes (MWCNT) was established.



HRTEM micrographs of: (A) individual tubes, (B) the MWCNT-PU adduct



Dispersions of MWCNT in acetone: (A) without additive; (B) with PU (four different polymer grade) after 12 months storage, (C) with PU after centrifugation (third vials from left to right of B).

## References

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