Regioselective Reactions, Spectroscopic Characterization, and Cytotoxic Evaluation of Spiro-pyrrolidine Thiophene

Salwa Abdelwahhab ,Sameh A. Rizk,a* Salwa S. Abdelwahab,b and Hanan A. Sallama

Abstract

Spiro-oxo-indole/pyrrolidine-thiophene base possessed significant pharmacological activity. The [3+2]

cycloaddition reactions of thia-methine ylide respected through multi-component reaction affording regioselective

and stereoselective spiroindoline-3,20-tetrahydrothiophene derivative 3. Reaction of such compound

with different electrophilic and nucleophilic reagents afforded bioactive heterocyclic compounds 4ó16. Biological

evaluation showed that these synthesized spiro-pyrrolidine exhibited moderate to good cytotoxic activity.

Among them, compounds 7 and 14 displayed the best cytotoxic activity against MCF-7 and WI-38

cells with the IC50 values of $7.02"\tilde{0}"0.6$ and $8.97"\tilde{0}"0.9"$ m (very strong), respectively. Compounds 4, 5,

and 12 exhibited strong cytotoxicityøs with IC50 16.28"Õ"1.7, 11.16"Õ"1.1, and 19.14 "Õ"1.7" m, respectively,

against MCF-7 mammary gland cell line. All compound structures were supported by spectroscopic data and elemental analysis.

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