Euro Chemistry 2020 Euro Toxicology 2020 Advanced Energy Materials 2020

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July 15-16, 2020

WEBINAR

Hong-Ngoc PHAM, Organic Chem Curr Res 2020, Volume 09

Obtainment of 3,4-Dihydro-2H-1,4-Benzoxazine Enantiopure Analogues For Potential Biological Applications

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2,4-dihydro-2H-1,4-benzoxazine and its derivatives were described as promising candidates for many choices ${\mathfrak I}$ of therapeutic applications. This fascinating class of heterocyclic compounds can exhibit various biological properties such as antibacterial1, cardiovascular protector2, anti-hypertensive3, anti-angiogenesis4, anti-thrombin5 or neuroprotective activities. Due to those potential effects, 3,4-dihydro-2H-1,4- benzoxazine became an attractive target to develop. However, because of receptor selectivity, chirality now becomes a top subject in chemistry. Within one molecule such as levofloxacin or thalidomide, different chirality may exhibit different pharmacological activities. As a result, we raised an awareness about the obtainment of 3,4- dihydro-2H-1,4-benzoxazine enantiopure analogues but there were not many reports in the literature brought up an effortless procedure to straightforward way to directly acquire the chiral ethyl 3,4-dihydro-2H- 1,4-benzoxazine-2-carboxylate. As our targeted compound could be easily prepared by allowing appropriate aminophenol reacts with ethyl 2,3-dibromopropionate, this reaction always lead to racemic 1,4- benzoxazine as the main product in high yield. Our desire is to attempt chiral (R)- or (S)-ethyl-2,3dibromopropionate which had never been described in the literature in order to replace the role of racemic one and (R)- or (S)-ethyl-3,4-dihydro-2H-1,4-benzoxazine-2-carboxylate by two strategies: enantioselective synthesis and preparative enantioseparation of racemate using different chiral stationary phases (CSPs). Once ethyl-3,4-dihydro-2H-1,4-benzoxazine-2-carboxylate enantiopure is obtained, it could be easily to introduce as new β -amino acid analogs in peptidic chains by coupling O- and N-terminal extremities with different α -amino acids. Finally, studies of the conformational behavior of obtained compounds will be led and their potential biological activities will be examined.

Biography

Hong-Ngoc PHAM received her master degree in Drug Development in 2016 from Vietnam-France University (Vietnam). She joined Laboratoire de Chimie Physique Macromoleculaire (LCPM), Université de Lorraine (France) in 2017 as PhD student in Organic Chemistry. Her research focused on the synthesis of heterocycle-based peptidomimetics for different biological applications. She expects to work as a researcher and teaching assisant in university after gaining the PhD diploma.