

Design, Optimization and Synthesis of Certain Heterocyclic Compounds with Potential Targeted Anticancer Activity.

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بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ

قالوا

سبحانك لا علم لنا
إلا ما علمتنا إنك أنت
العليم العظيم

صدق الله العظيم

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تصميم وتعظيم وتشبيد بعض المركبات غير متجانسة الحلقة التي لها فاعلية محتملة ضد السرطان.

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