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Molecular Modeling and Docking Techniques for Drug Discovery and Design Bhat, Ajmal Rashid, Ahmed, Sumeer, Kawsar, S. M. Abe, 2025-02-05 In the realm of pharmaceutical research the challenge of efficiently discovering and designing new drugs to combat diseases is ever present Traditional approaches to drug discovery often rely on time consuming and costly experimental methods leading to lengthy development timelines and high failure rates This problem is exacerbated by the complexity of molecular interactions and the vast chemical space to explore As a result there is a pressing need for innovative solutions that can streamline the drug discovery process and improve its success rate Molecular Modeling and Docking Techniques for Drug Discovery and Design addresses this critical challenge by offering a comprehensive guide to advanced computational methods in pharmaceutical research Edited by leading experts in the field the book provides insights into molecular modeling docking and other computational approaches that can significantly accelerate the drug discovery process By leveraging computational tools and software researchers can simulate molecular interactions predict drug efficacy and optimize chemical structures with greater speed and accuracy than traditional experimental methods

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