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Damaged DNA

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***E. coli* RNase H–substrate interaction**

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Hydration structure

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Interfacial tension

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isoDNA·RNA duplex

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CDK4; Protein kinase (Tian, G. (95) 79)

Molecular dynamics

Nucleic acids evolution; *E. coli* RNase H-substrate interaction; *E. coli* RNase H inhibitor interaction; *isoDNA*·RNA duplex; *isoDNA* duplexes and triplexes (Premraj, B.J. (95) 253)

Molecular dynamics

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Molecular dynamics

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Sugar pucker; DNA dodecamer; X-Ray structure; Damaged DNA (Tofazzal Hossain, M. (95) 69)

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Nucleic acids evolution

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Phosphatidylcholine monolayer

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Protein C deficiency

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Release and penetration of flunitrazepam

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Tethered-ligand

Parathyroid hormone (PTH); PTH receptor; G-protein coupled receptors; Coupling of G-proteins to PTH receptor; Constitutively active receptors (Monticelli, L. (95) 165)

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