INTRODUCTION:

Throughout this Poster I will be showing the researchand knowledge I have gathered not only on the importance of this topic but on how Intercalating agents could possibly be used in the future as a possible very successfulcancer treatment. Discussed will be the mechanism and concepts of the process that these drugsperform inside the body.

Toxicity of DNA Intercalating Agents in the Human Body By: Jacob Snell

Chemical Mechanisms of Intercalation occurring

Agents, Mechanisms and Toxicity:

- There are multiple intercalating agents under research, the three main ones being, Acridine, Actinomycin, Ethidium Bromide.
- These agents bind to DNA which in turn inhibits a key replication enzymecalled TOPO some rasel
- Due to this inhibition and space between base pairing, mutations canarise in DNA replication
- Not only is replication inhibited but all DNA repair and translation becomes effected leading to cell death



Affinity:

- successates in DNAbinding
- complexmechanismswith.

Statistical Measurements of successful binding affinity to DNA

Possibilities for the Future:

cancer being uncontrollable cell With replication this research and technology with extreme studies performed could lead to a breakthrough in successful ancerresearch If a way to attach a signal ligand to tumor cells and avoid healthy cells was discovered, this would lead to an extreme increase in cell selectivity and binding affinity. It is impossible for treatments such as taiaigmrf2007703.014. Epub2007 Apr

Acridine and Measurement of Binding

Due to the property of fluorescenceAcridine has it makes it very easyto measureand study in the body

Acridine has a very high binding affinity leading to high

The permanent inhibition this agent has on DNA replication makes it very useful to study intercellular