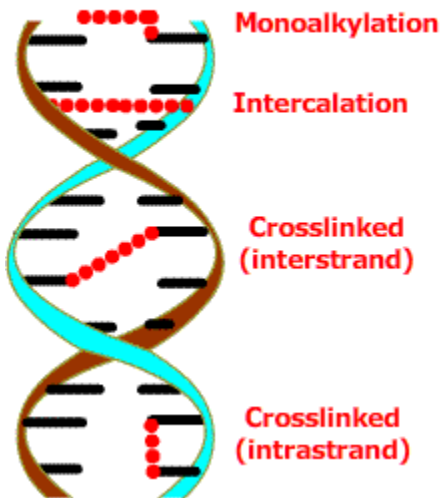
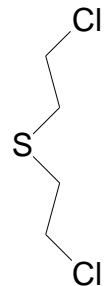


Αλκυλιωτικοί παράγοντες

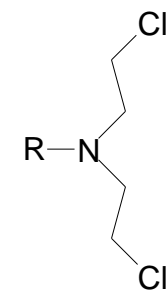
Alkylated DNA

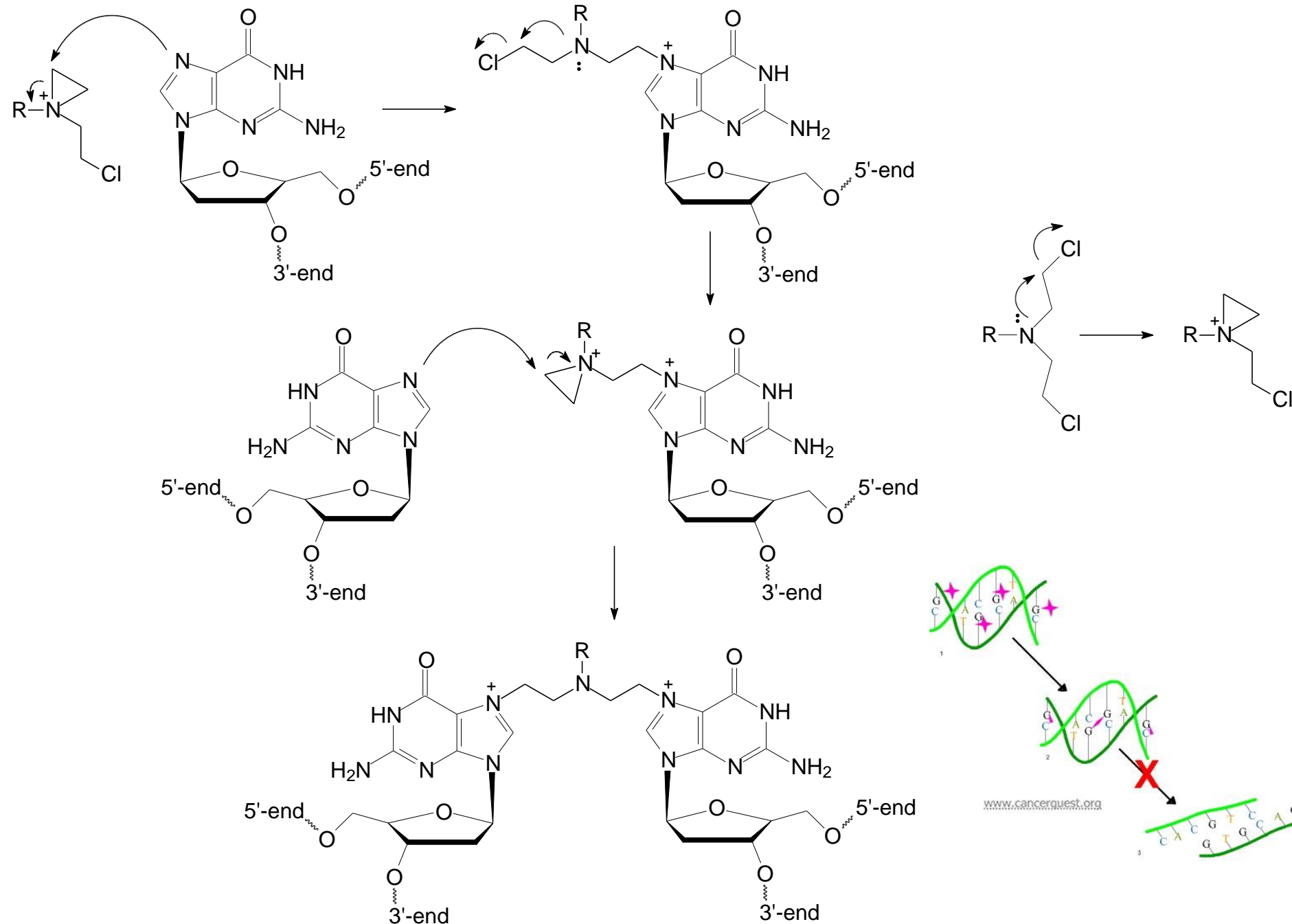


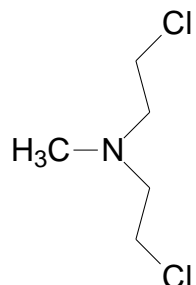
Sulfur mustards



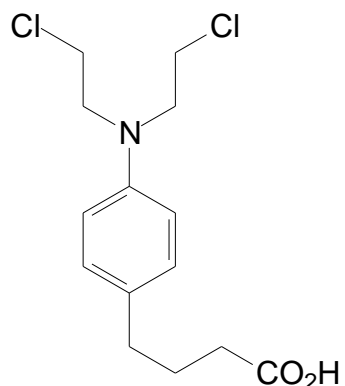
Nitrogen mustards



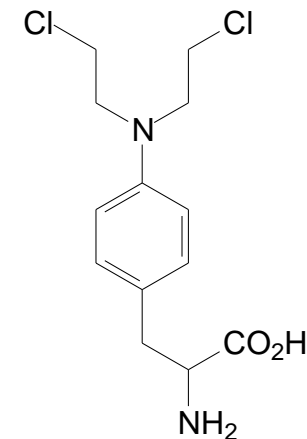




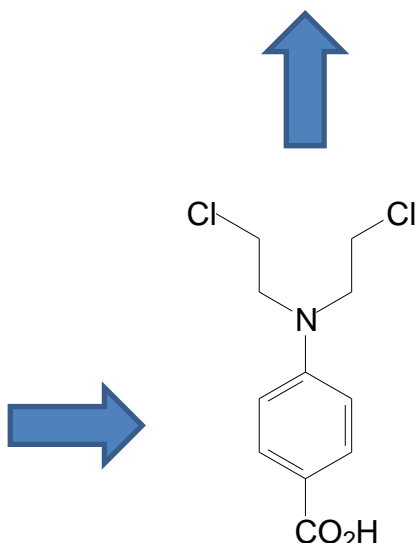
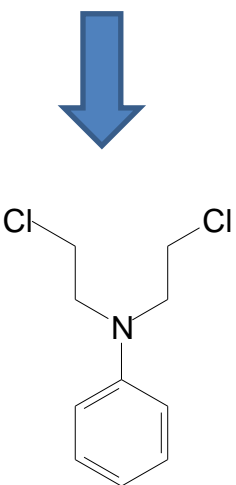
mechlorethamine



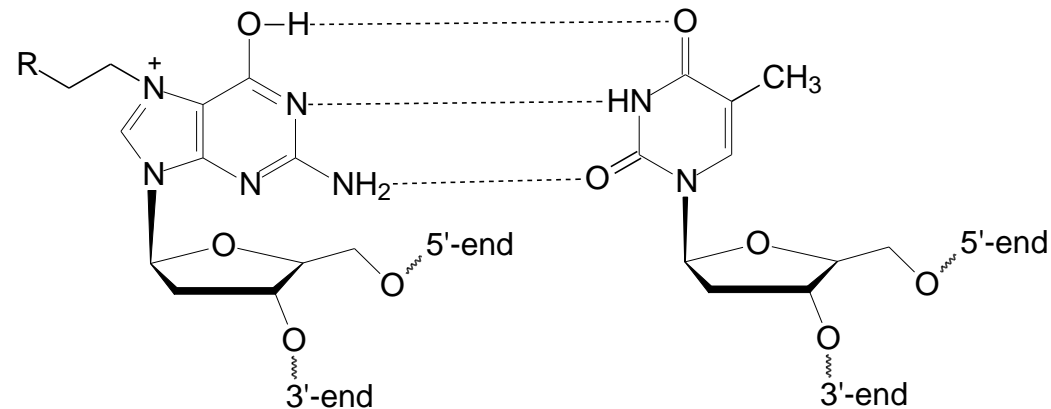
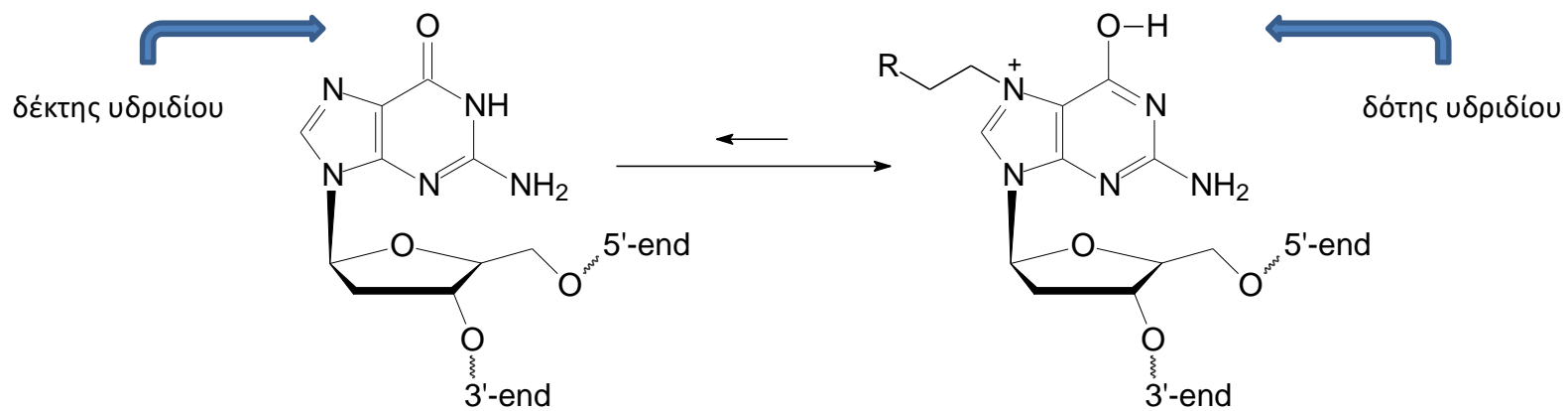
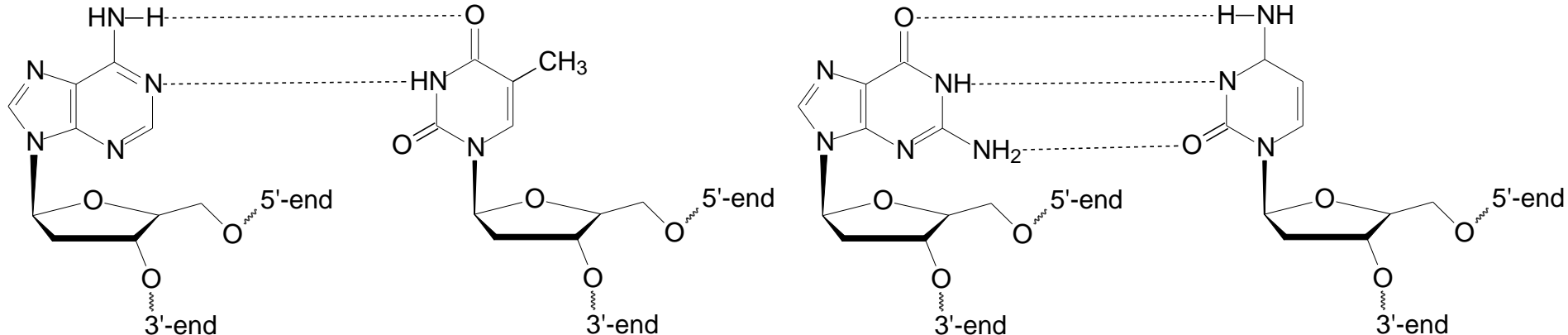
chlorambucil

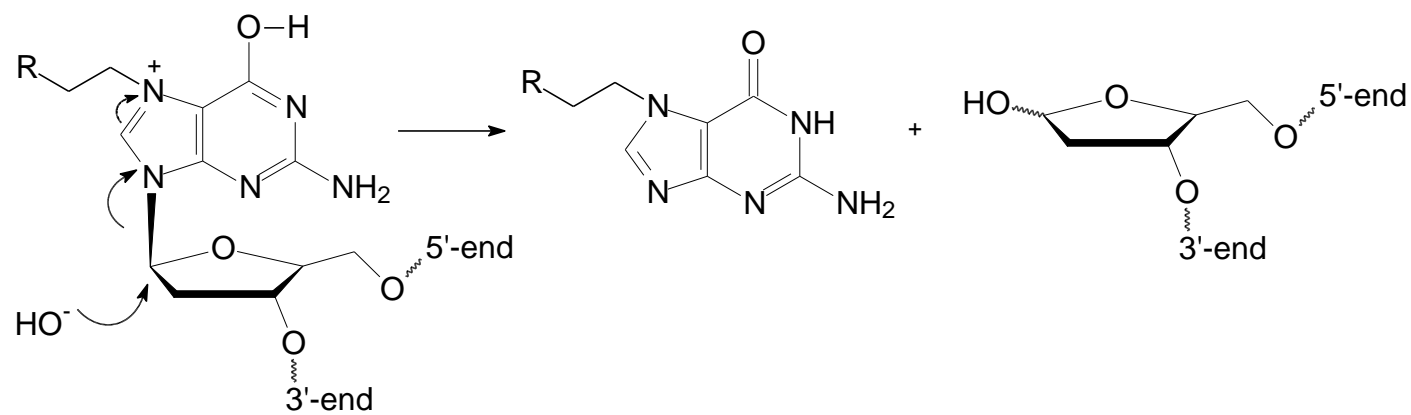


melphalan

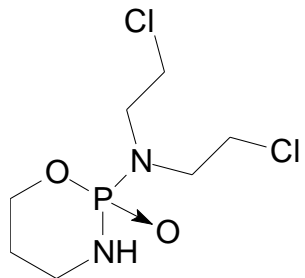


bendamustine

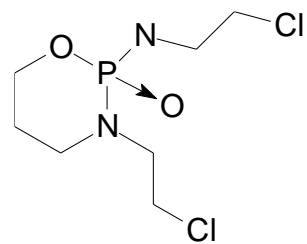




Oxazaphosphorines

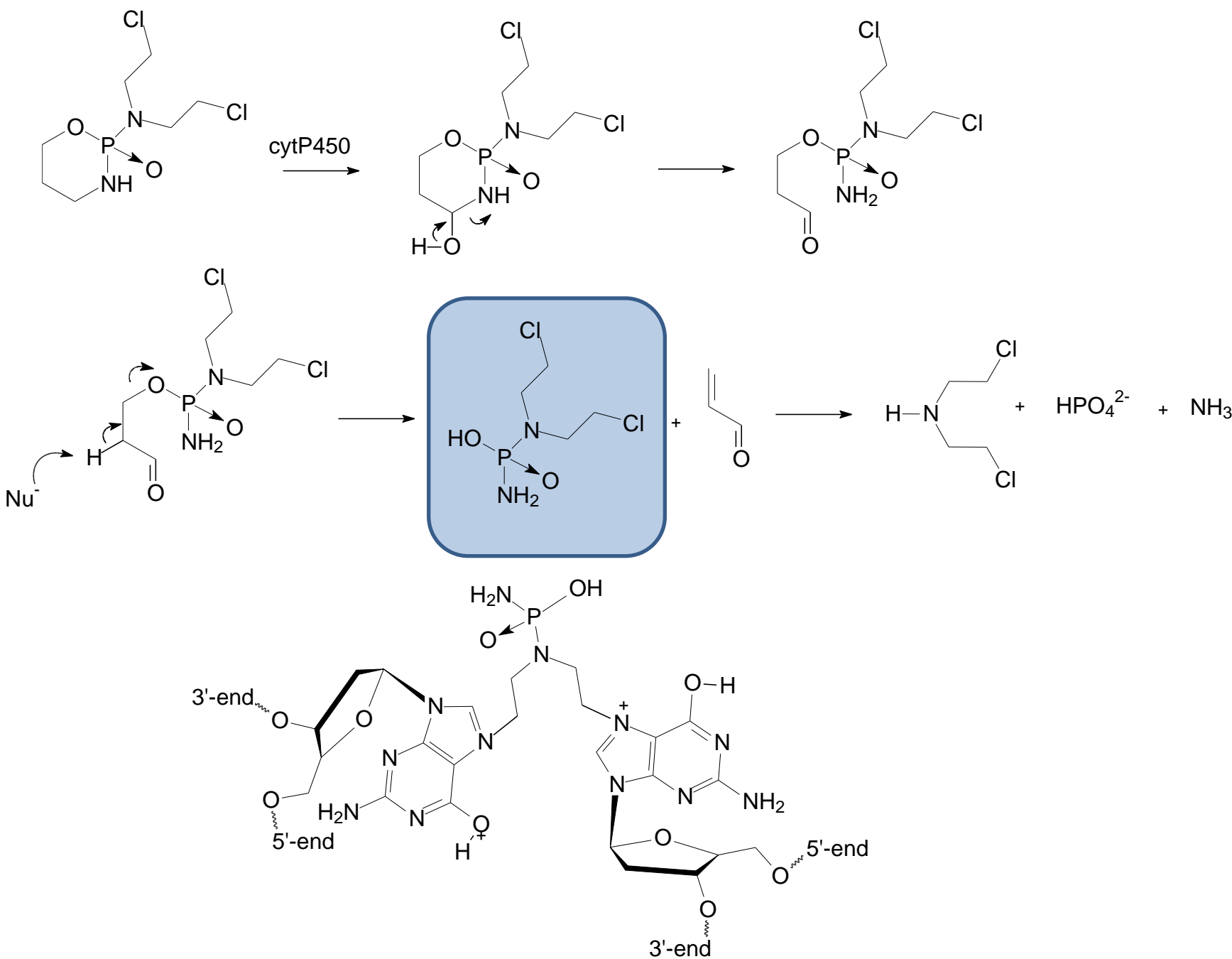


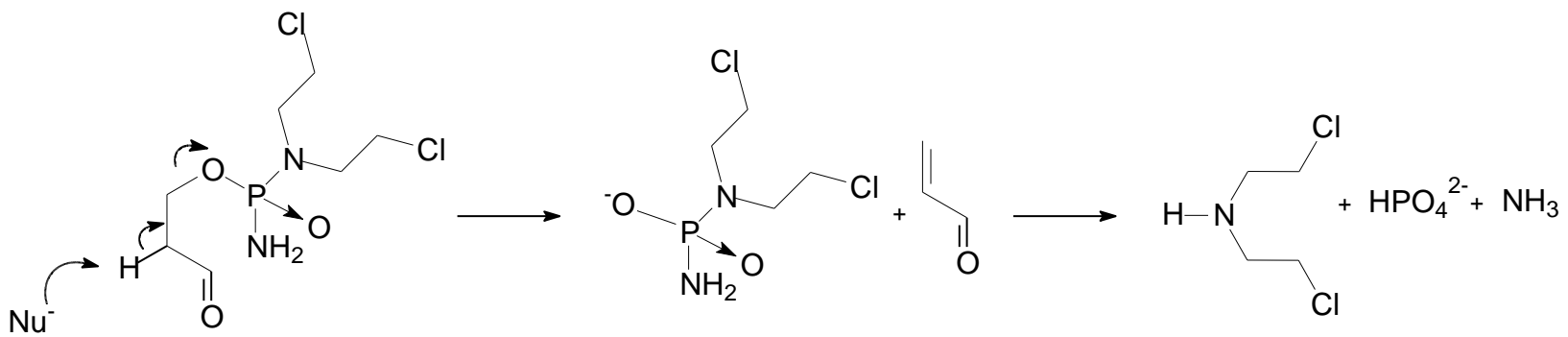
cyclophosphamide



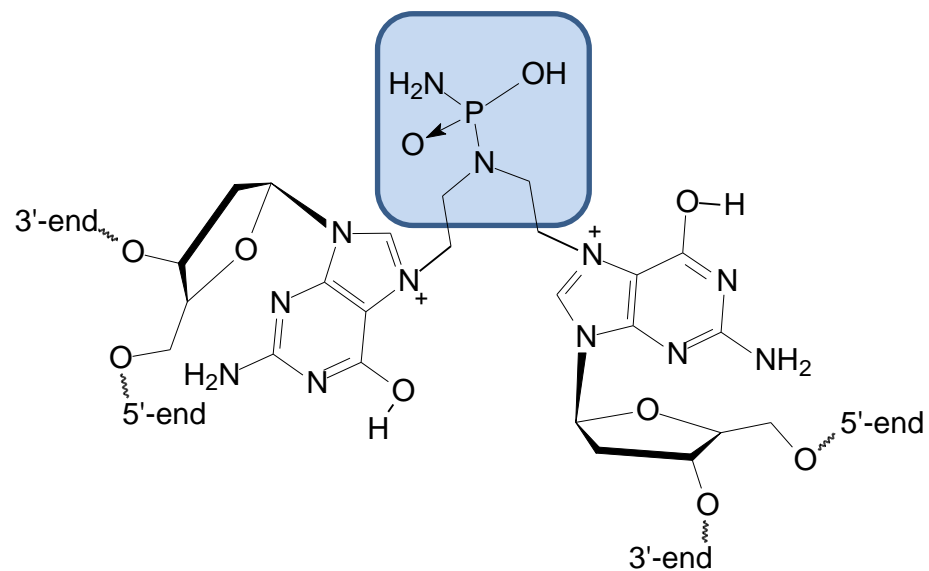
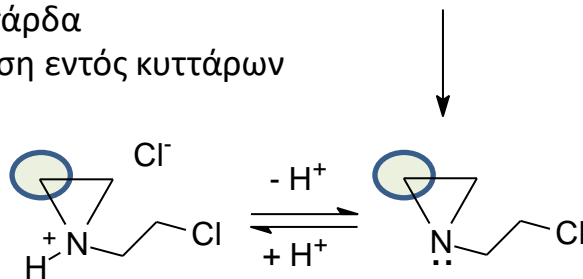
ifosfamide

Στα καρκινικά κύτταρα, σε αντίθεση με τα φυσιολογικά, υπερεκφράζονται οι φωσφοραμιδάσες.



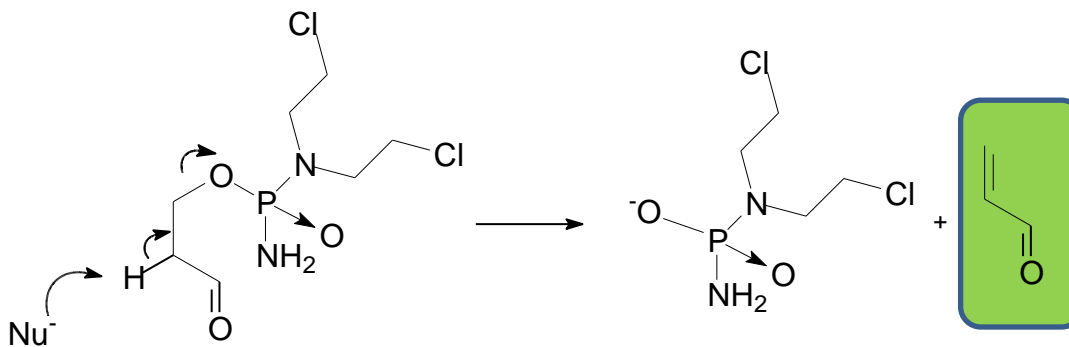


Φωσφοραμιδο-μουστάρδα
 $pK_a = 4.75$, συσσώρευση εντός κυττάρων



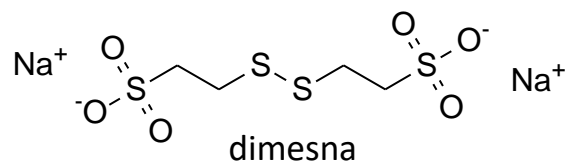
Πρωτεύων μηχανισμός (λόγω αζιριδινικού κατιόντος)

Δευτερεύων μηχανισμός, λόγω μειωμένου ηλεκτρονιόφιλου χαρακτήρα του αζιριδινικού C



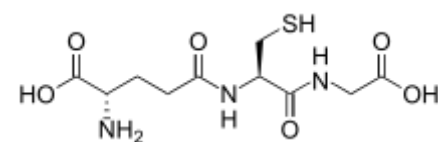
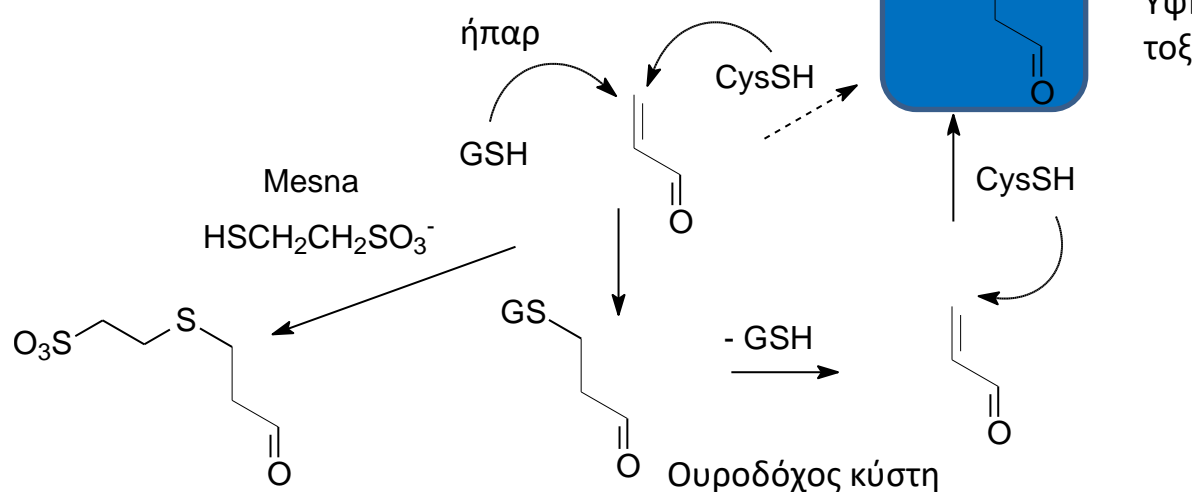
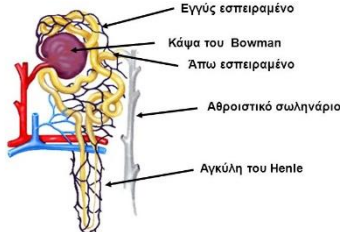
Η ηπατική μεταβολική ενεργοποίηση, συνεπάγεται χαμηλότερη τοξικότητα στο γαστρεντερικό και σχετικά πιο εκλεκτική κυτταροτοξικότητα.

Παρόλα αυτά το φάρμακο δεν είναι απαλλαγμένο παρενεργειών...



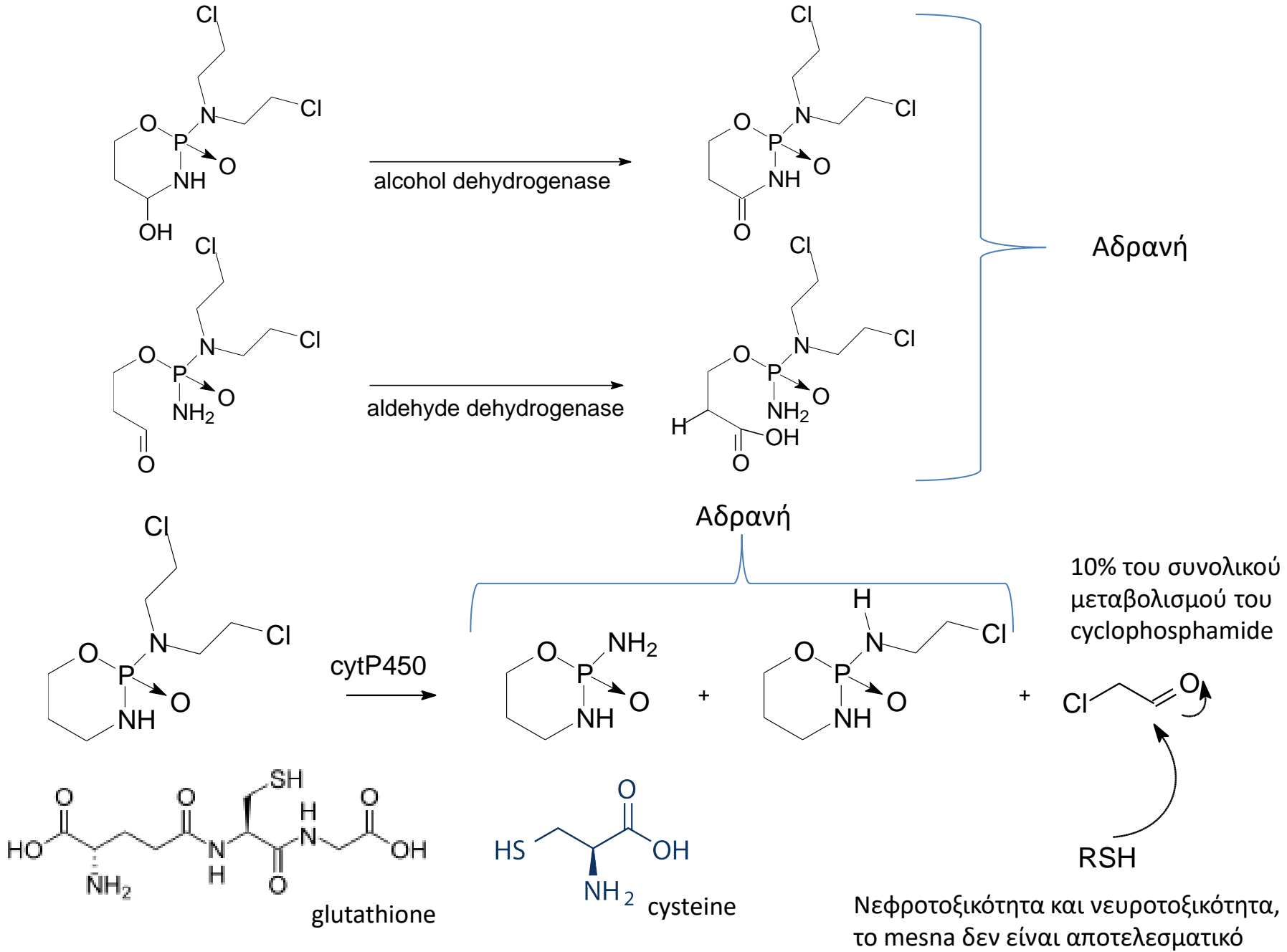
Εκλεκτική αναγωγή στα εγγύς σωληνάρια

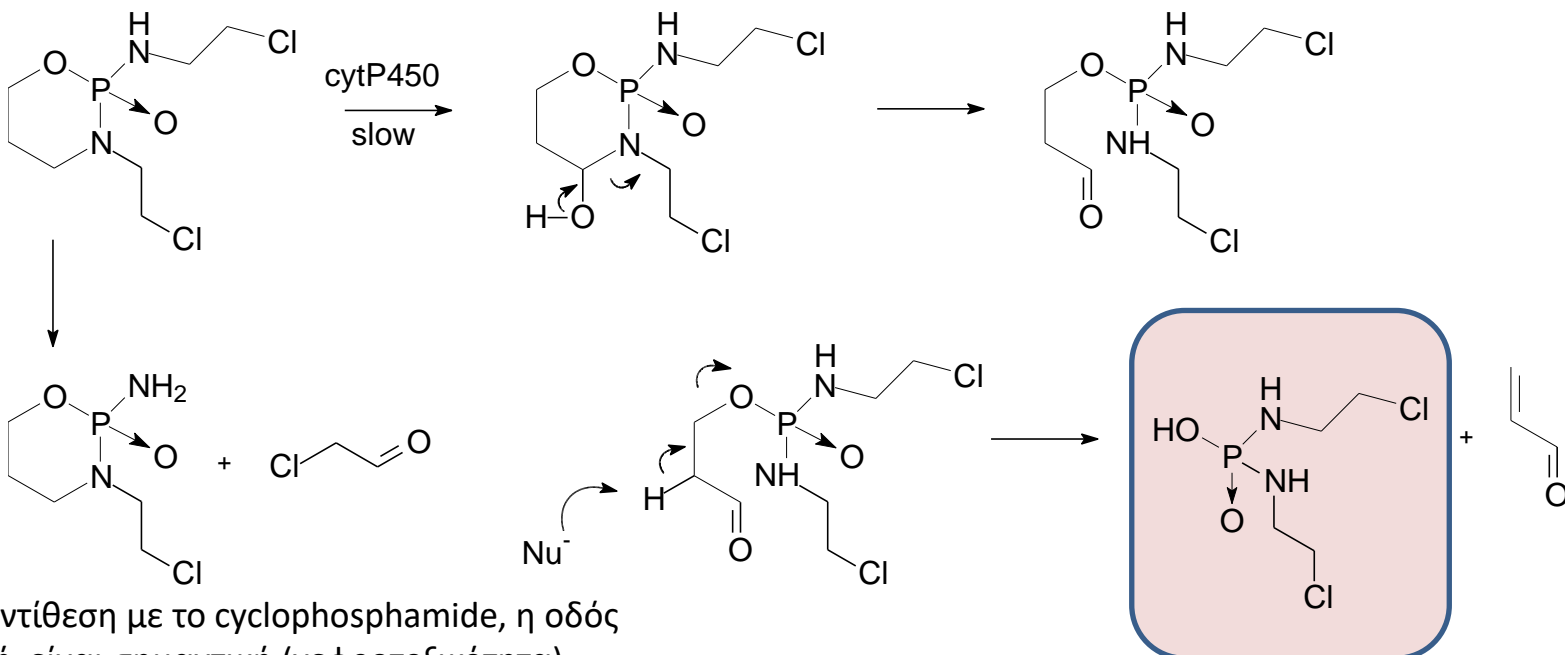
ΝΕΦΡΙΚΑ ΣΩΛΗΝΑΡΙΑ



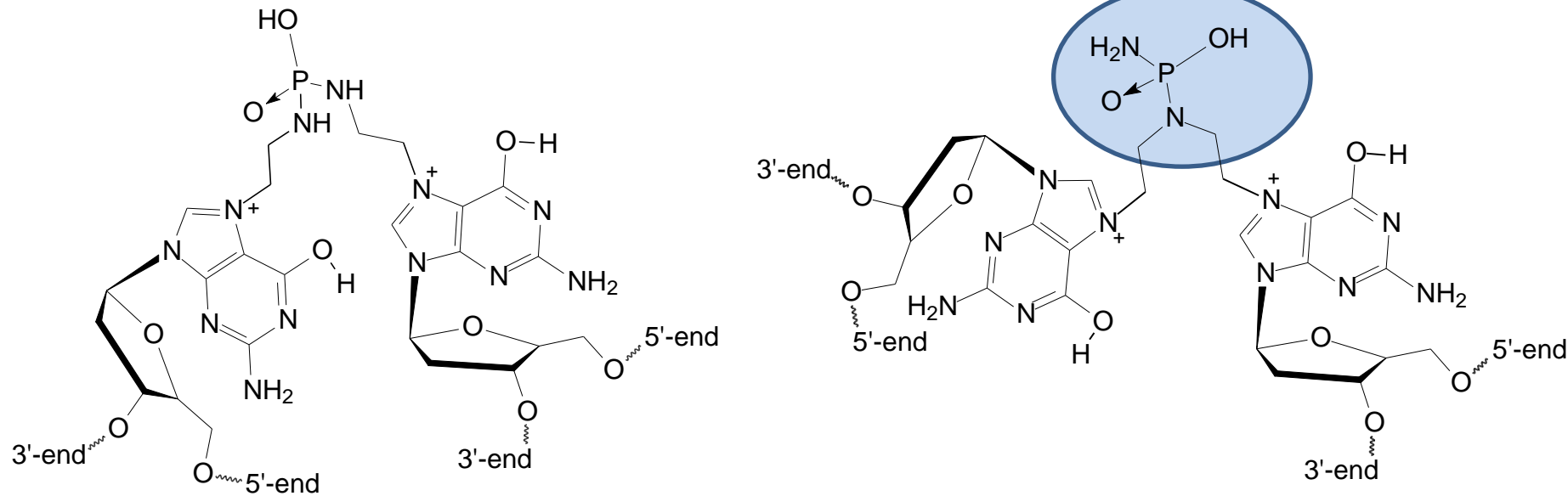
Υψηλή τοξικότητα

Οι σημαντικότερες συγκεντρώσεις mesna είναι στην ουροδόχο κύστη

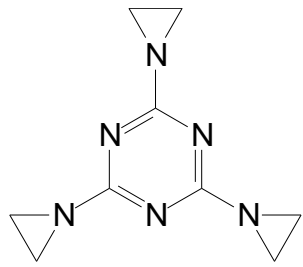




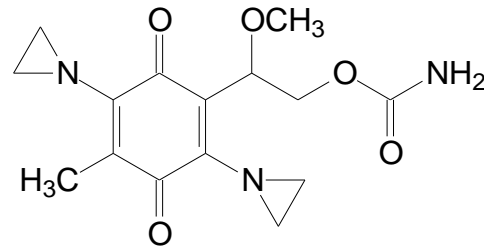
Σε αντίθεση με το cyclophosphamide, η οδός
αυτή είναι σημαντική (νεφροτοξικότητα)



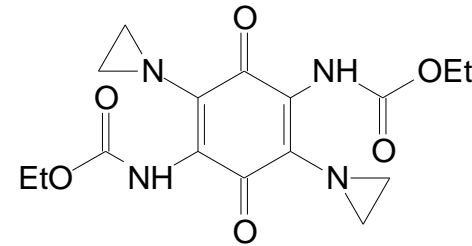
ethylenimines



triethylenemelamine

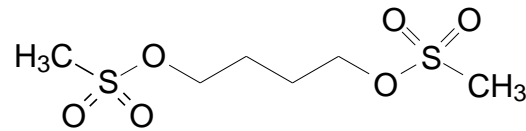


carboquone

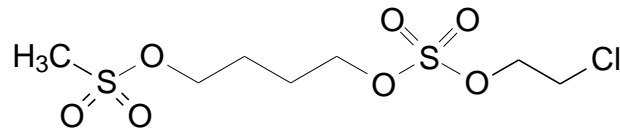


dazoquone

methanesulfonates



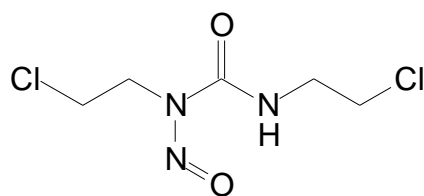
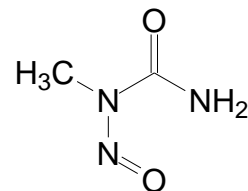
busulfan



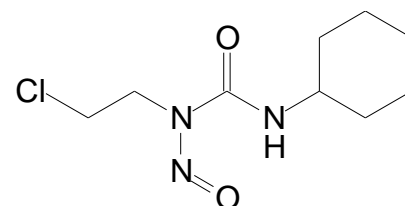
clomesone

nitrosoureas

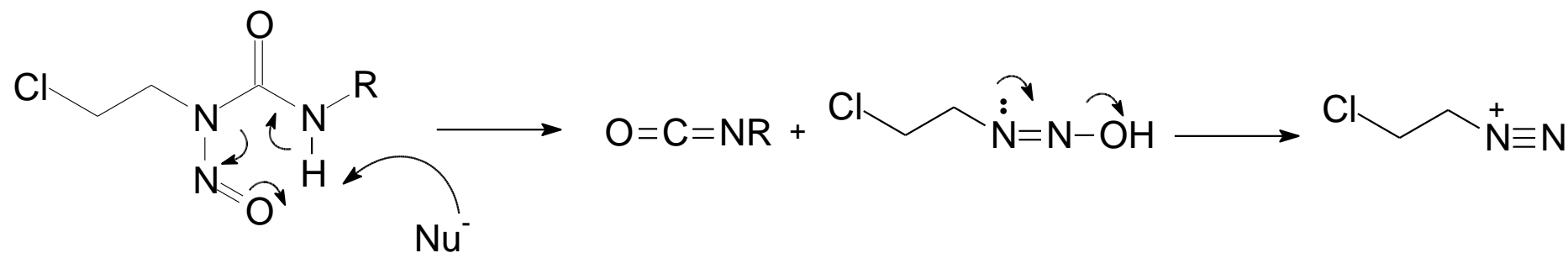
Ένωση-οδηγός,
Χαμηλής δραστικότητας

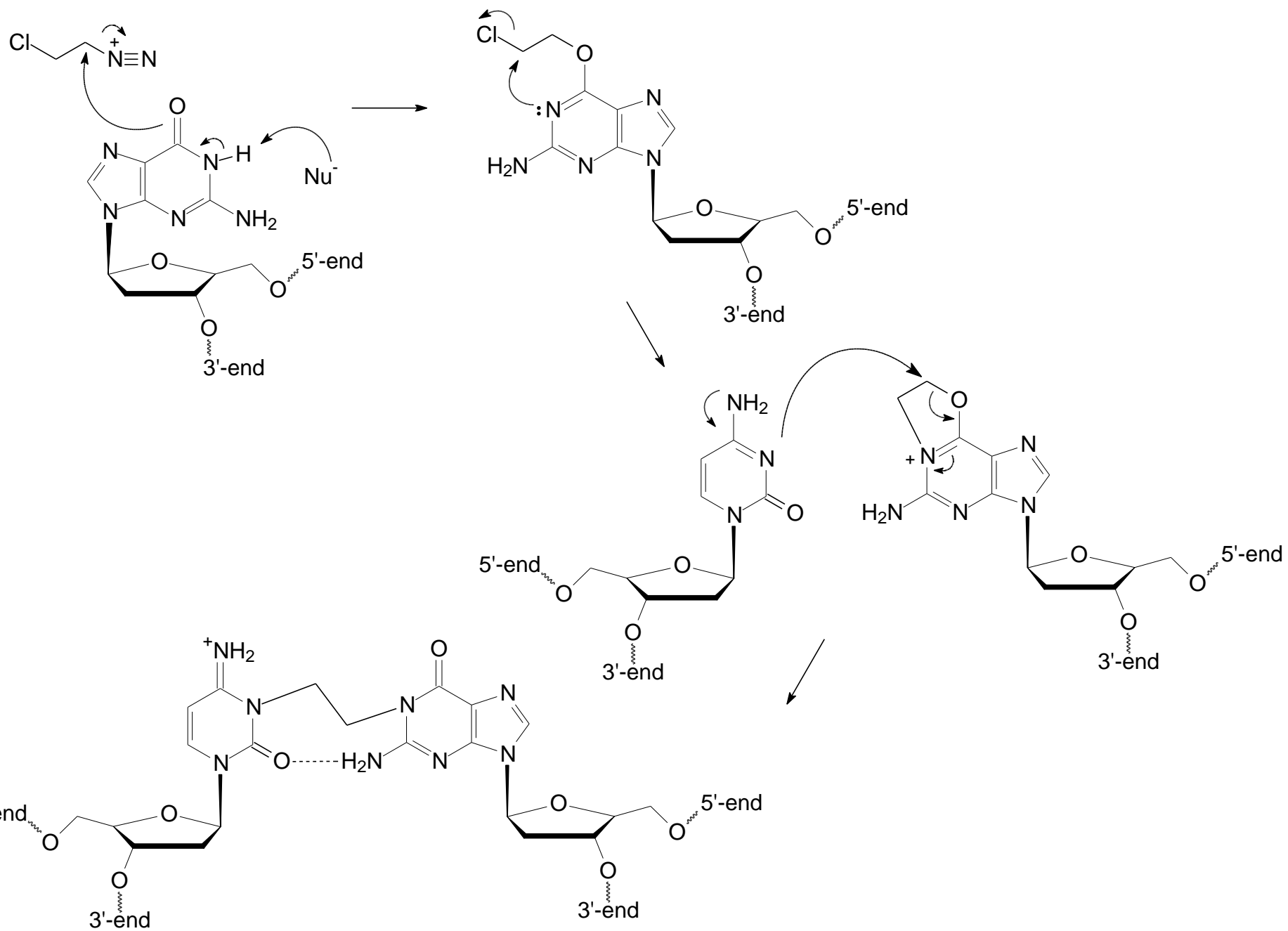


carmustine

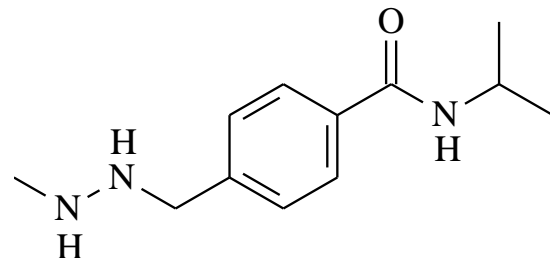


lomustine

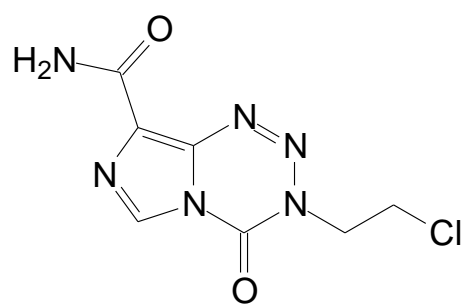




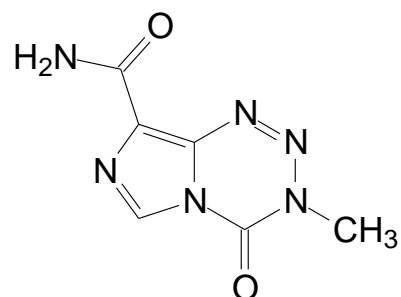
procarbazine



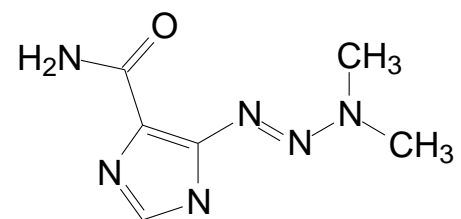
triazenes



mitozolomide

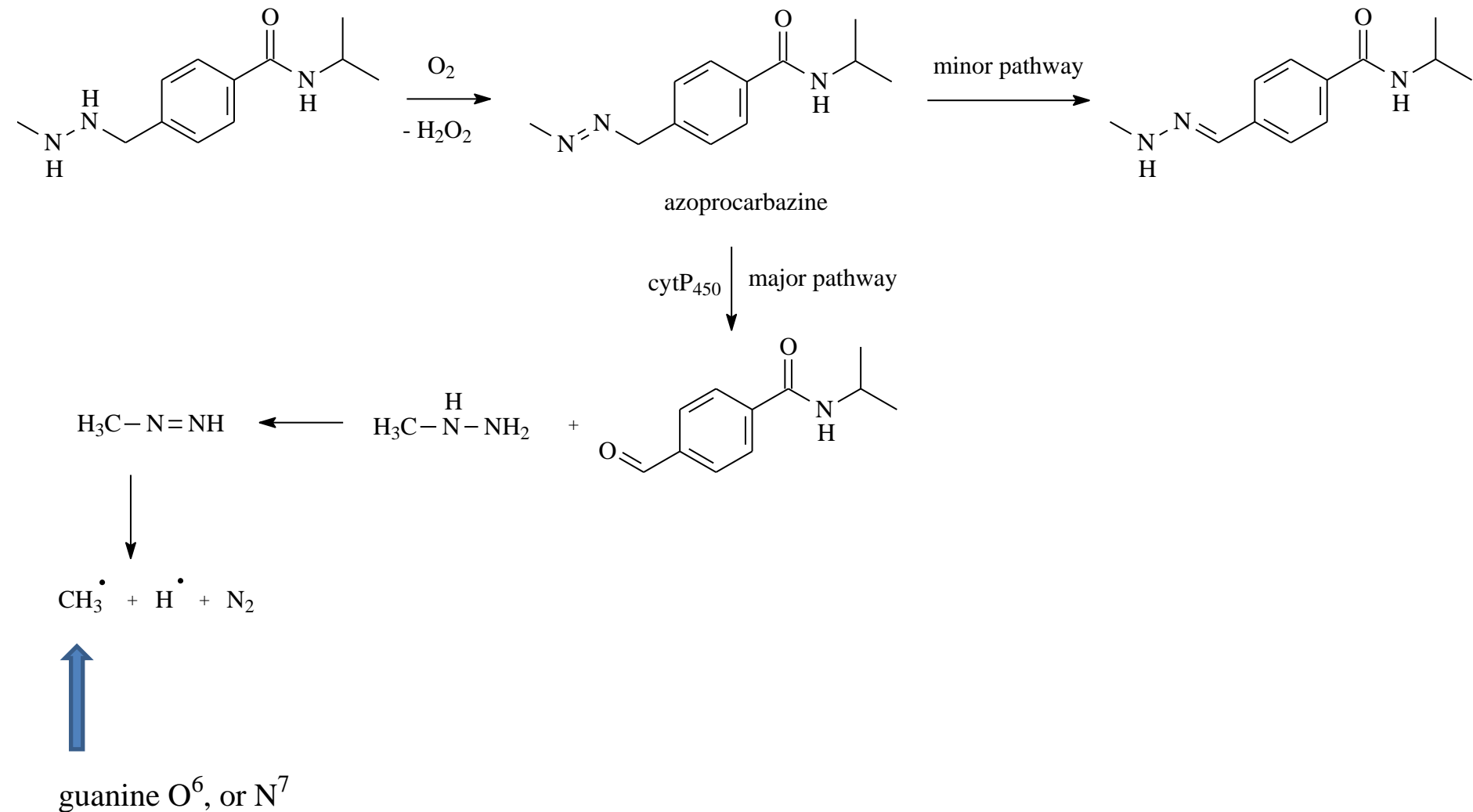


temozolomide (TMZ)

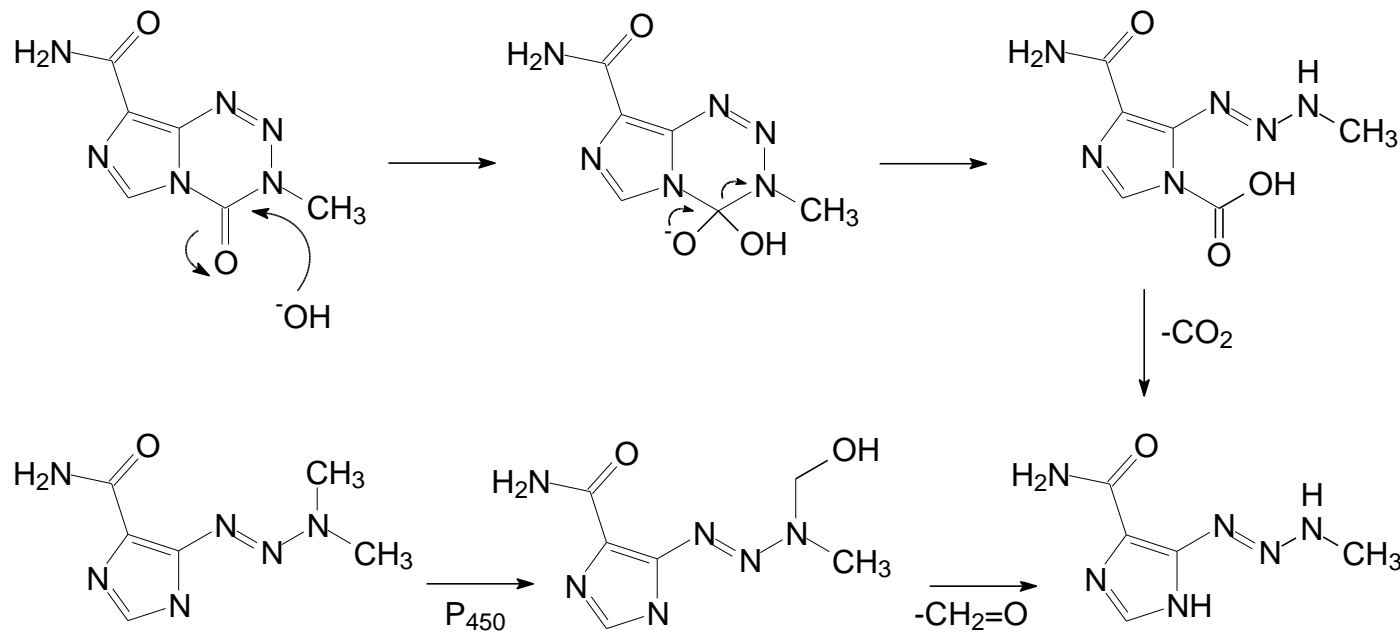


dacarbazine

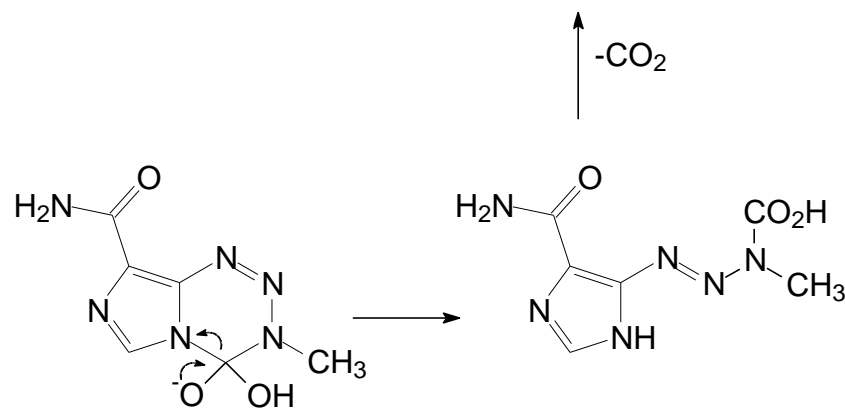
procarbazine

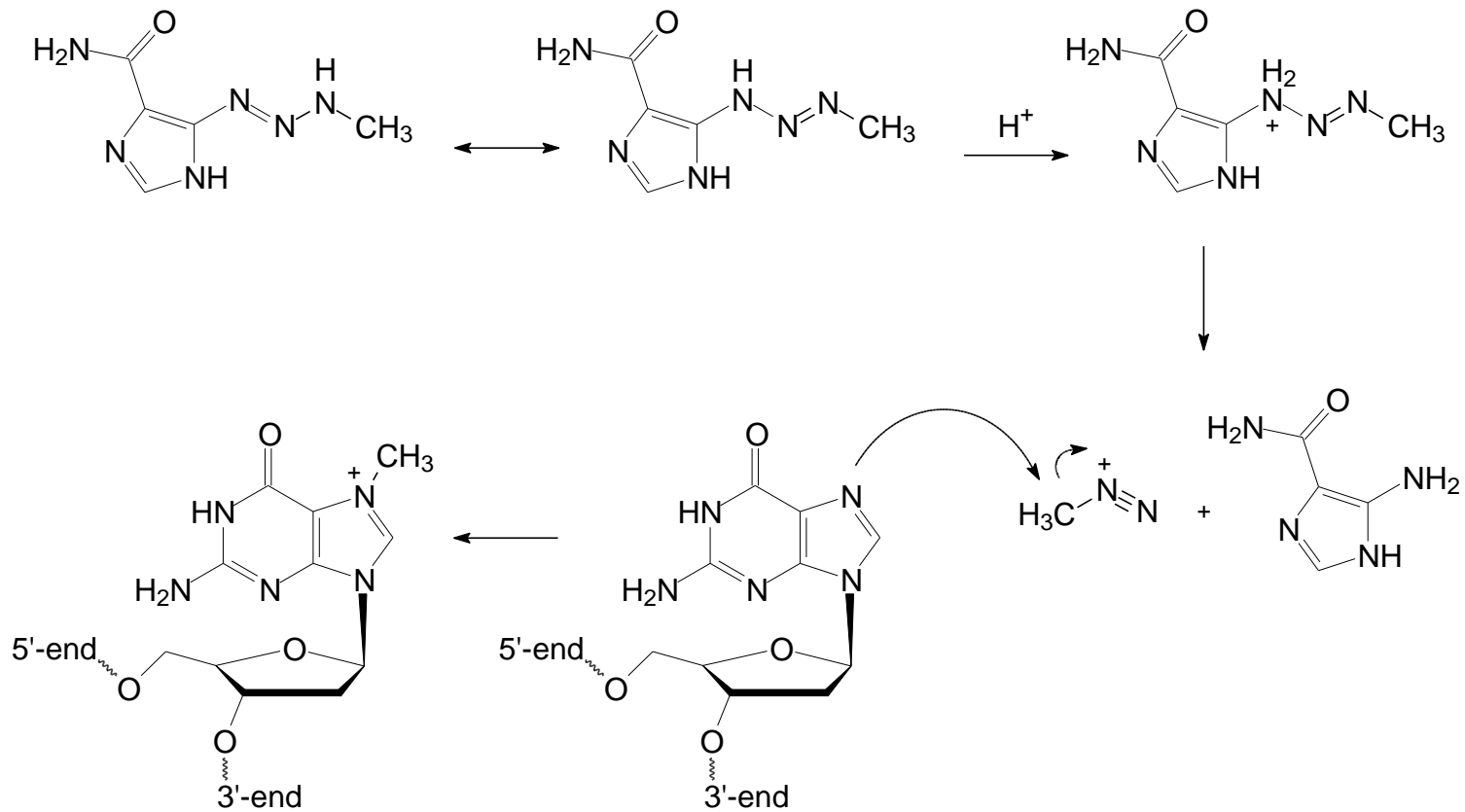


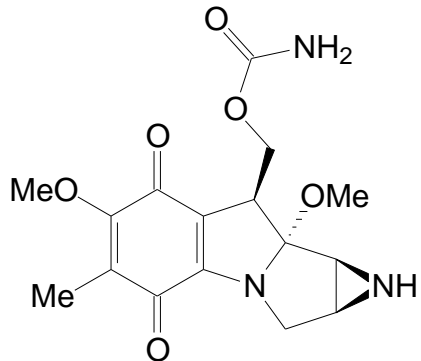
temozolomide (TMZ)



dacarbazine







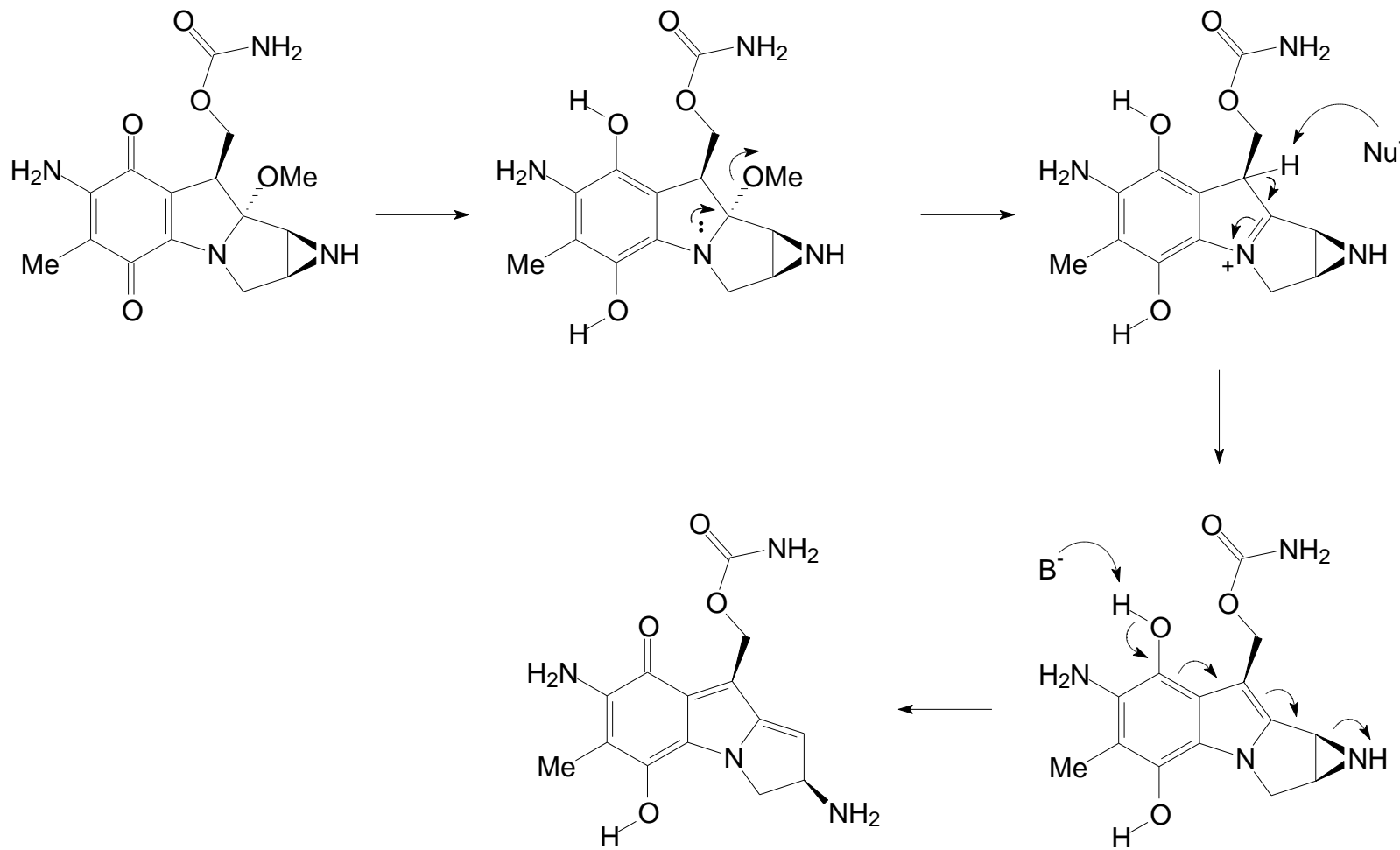
mitomycin A

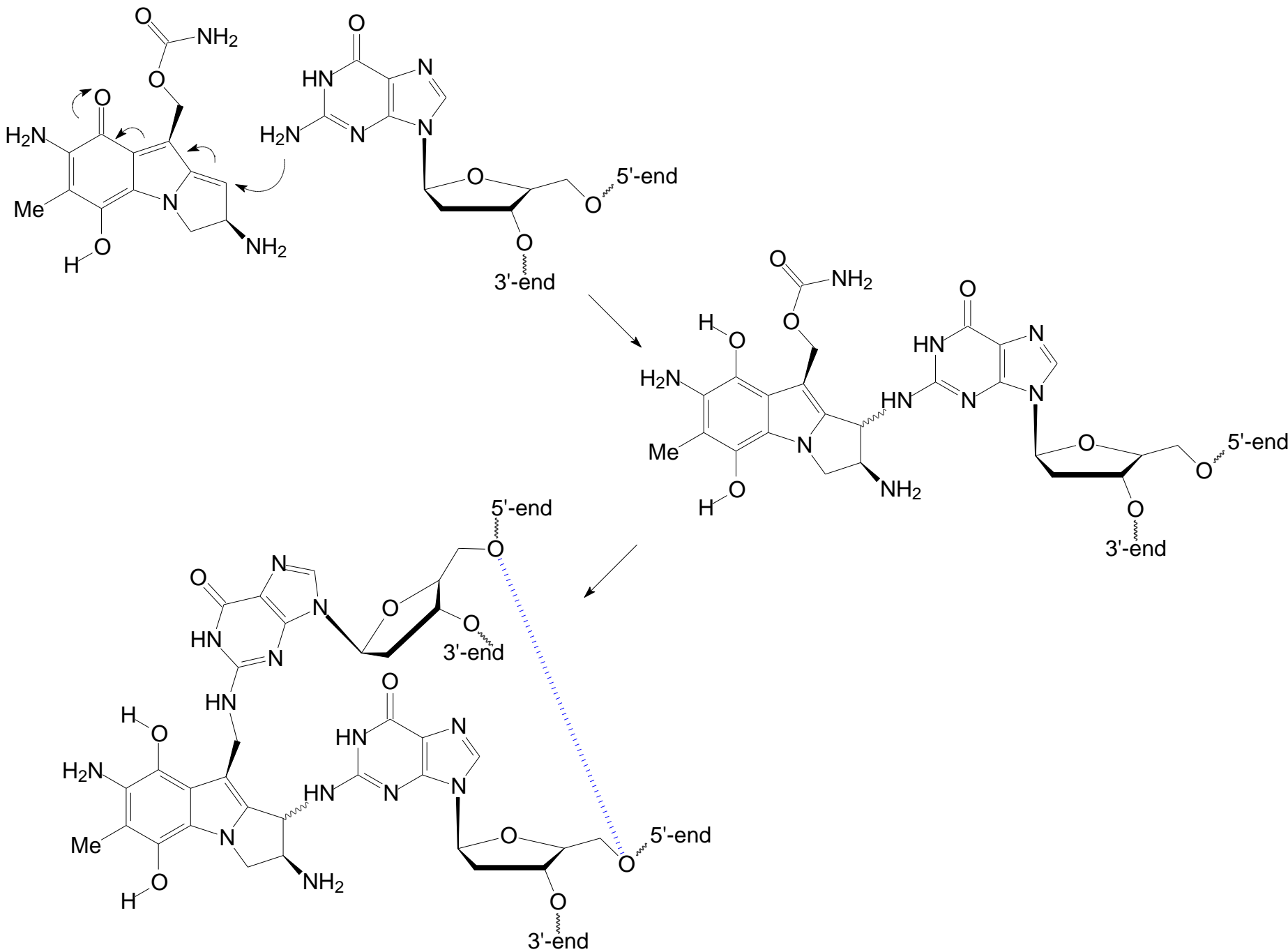
Απομονώθηκε το 1956 από *Streptomyces caespitosus*.
Διευκρινίστηκε η δομή το 1962.

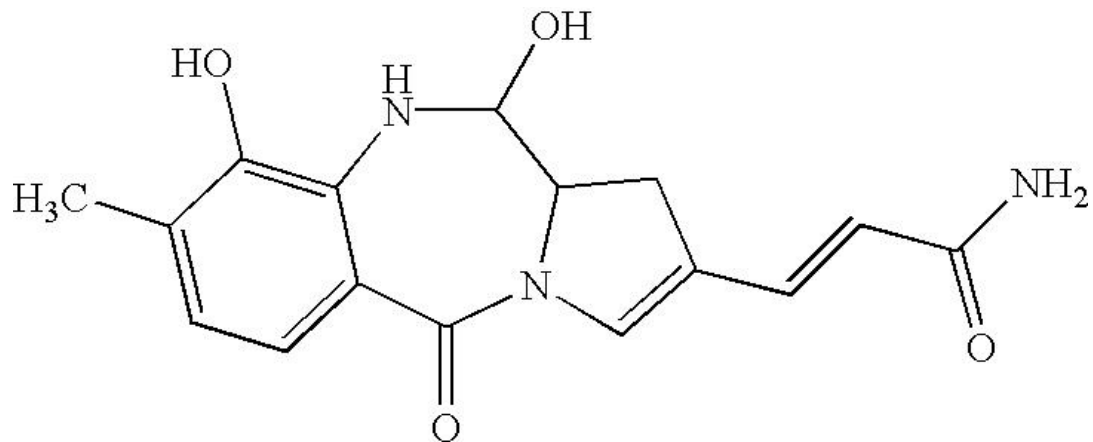
Είναι γνωστά 17 παράγωγα, 16 από τα οποία βιολογικώς δραστικά (αντιβιοτική, αντικαρκινική δράση).

Κυκλοφορεί για χρήση σε καρκίνους οισοφάγου, μαστού, παγκρέατος, ουροδόχου κύστεως.

Απομονώνεται από βακτήρια – δεν υπάρχει εμπορικά διαθέσιμη συνθετική μέθοδος.

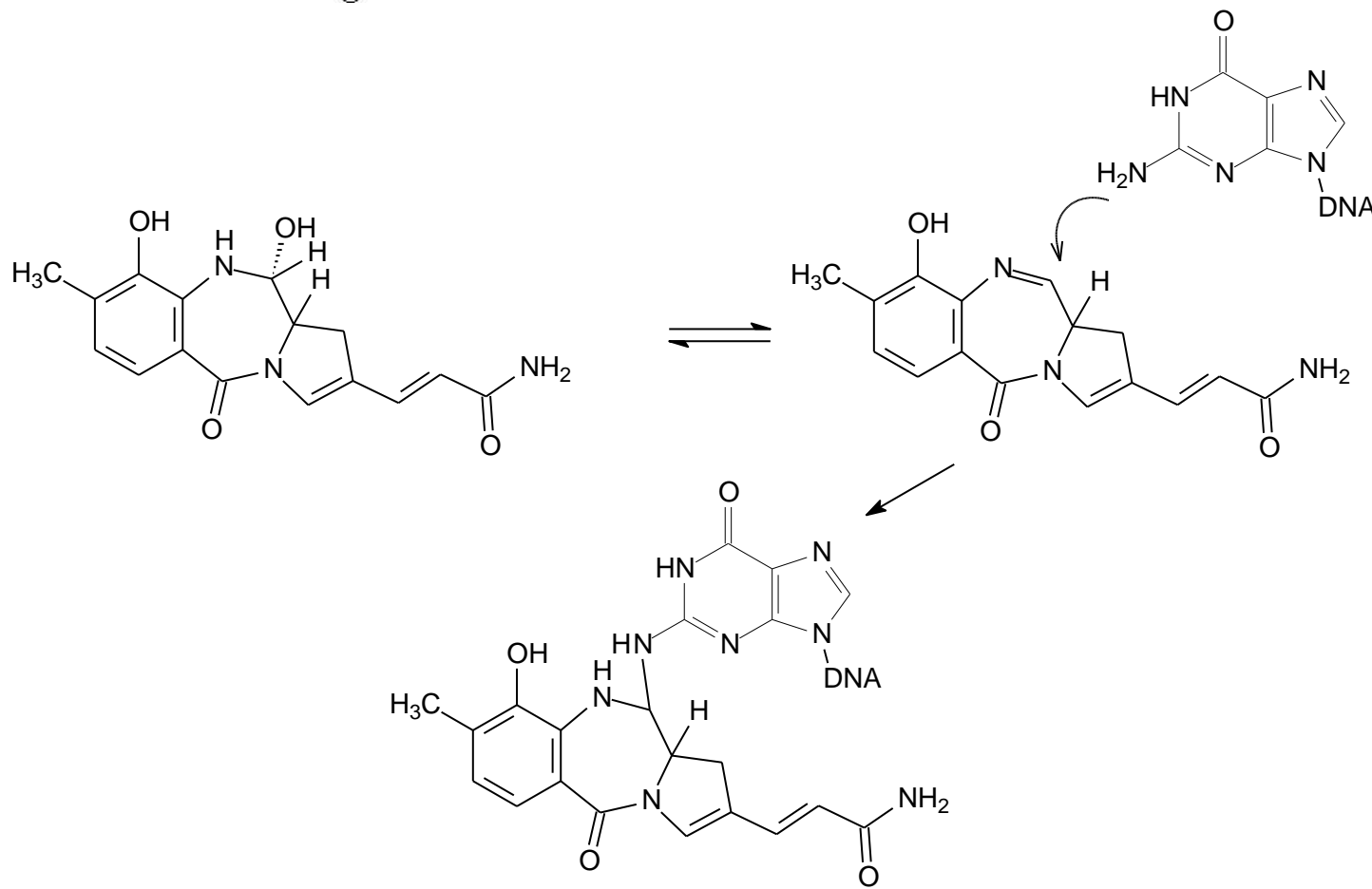


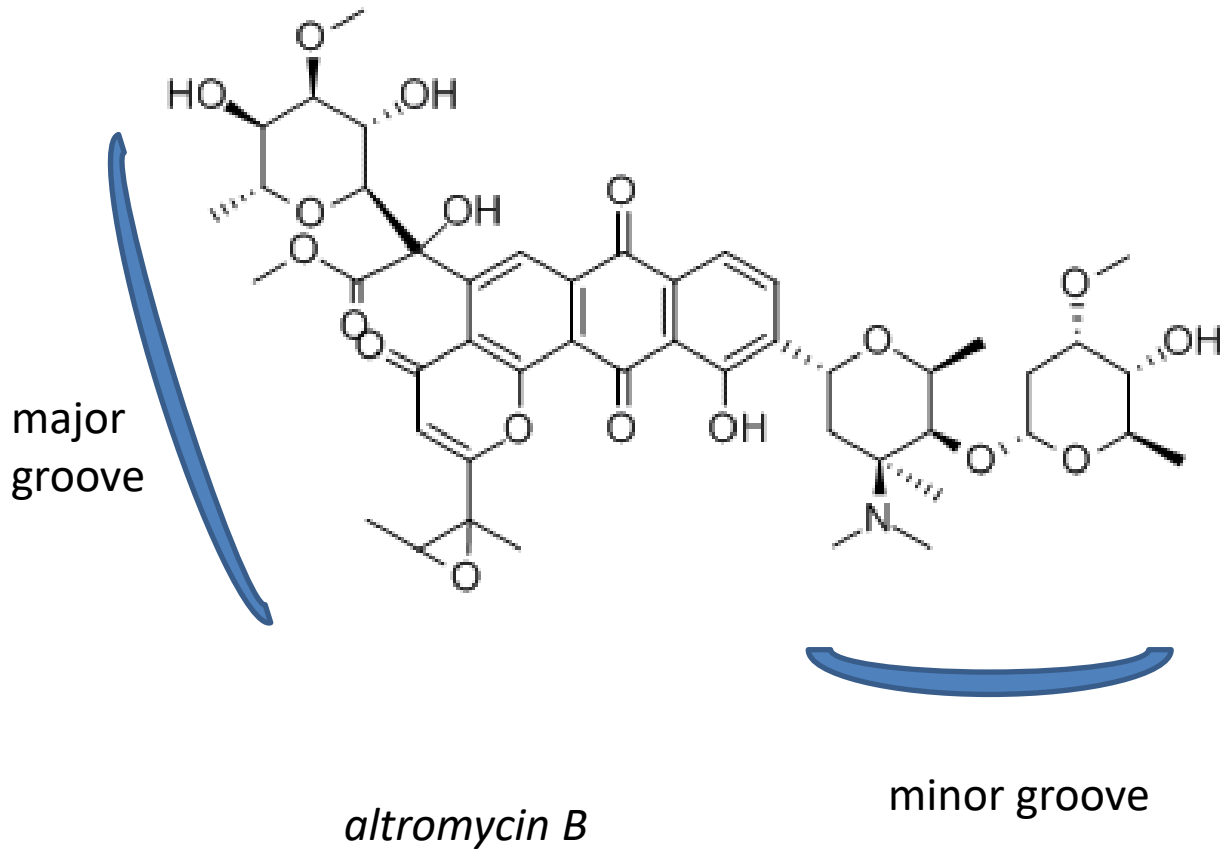




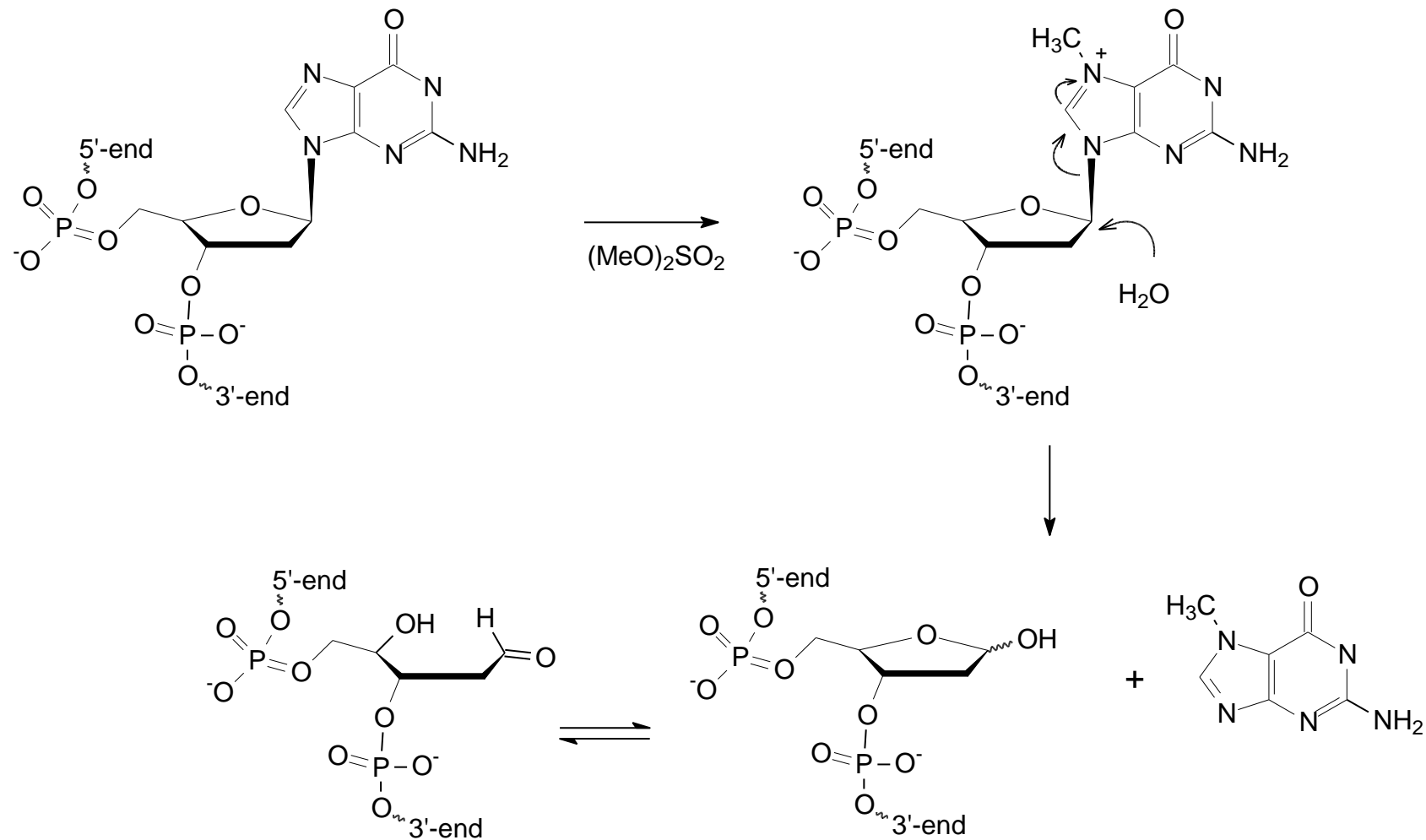
anthramycin

Sequence selectivity: 5'-PuGpu-3'

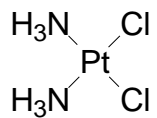




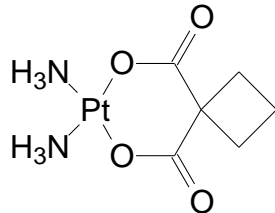
Αλκυλιωτικά αντιδραστήρια!!



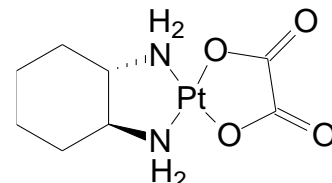
Platinum complexes



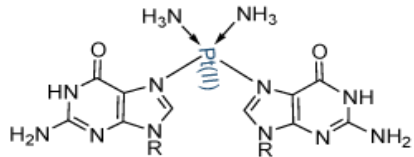
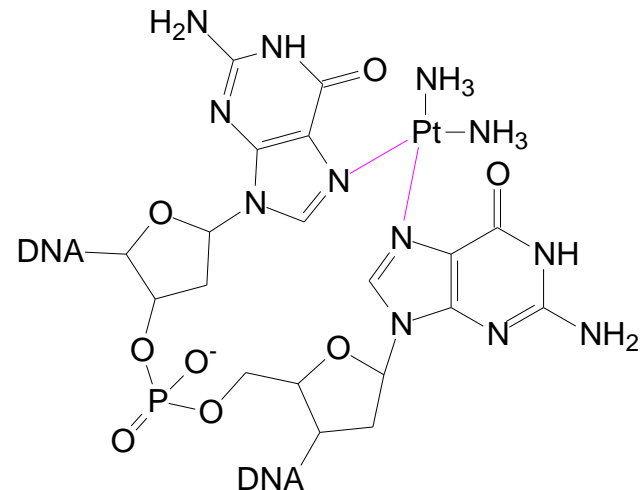
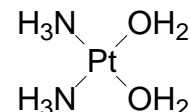
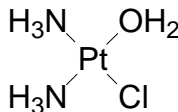
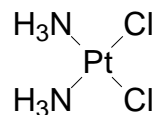
cisplatin



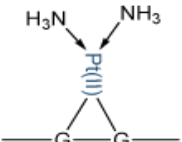
carboplatin



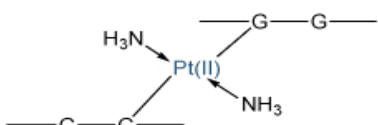
oxaliplatin



cisplatin reacts with *N*(7) of guanine



cisplatin forms **intra-strand** crosslinks:
poorly repaired



trans-platin forms **inter-strand** crosslinks:
repaired more efficiently