

22825 DAVIS DRIVE STERLING, VIRGINIA 20164

GENERAL PROCEDURE FOR LABELLING OF AMINO-MODIFIED OLIGONUCLEOTIDES

PHONE

703-437-6191

800-327-GLEN

<u>FAX</u>

703-435-9774

INTERNET

WWW.GLENRES.COM

This general procedure can be used to conjugate amino-modified oligonucleotides with active ester or isothiocyanate derivatives of fluorescent dyes which are not suitable for use as cyanoethyl phosphoramidites. At pH 9, conjugation occurs virtually exclusively at the amino group and does not react with the exocyclic amino groups of the nucleosides.

Note that oligos deprotected in AMA may require a desalting step prior to conjugation to prevent methylamine from reacting with the NHS ester.

 Dissolve the product from a 0.2 μmole synthesis of amino- modified oligonucleotide (i.e., approximately 0.1 - 0.2 μmoles of free primary amines) in 0.7mL of sterile distilled water.

- Add 0.1mL of 10X conjugation buffer (1M NaHCO₂/Na₂CO₂, pH9).
- 3. Freshly prepare a 10mg/mL solution of active ester in DMF. Add 0.2mL of the solution to the reaction mixture.
- 4. Vortex and allow the mixture to stand at least 2 hours. (Note: Overnight reaction may be more convenient.)
- Desalt the reaction mixture on a reverse phase cartridge or a column containing Sephadex G-25 or G-50 to remove the excess label. Purify the product using RP HPLC if necessary.