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
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
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
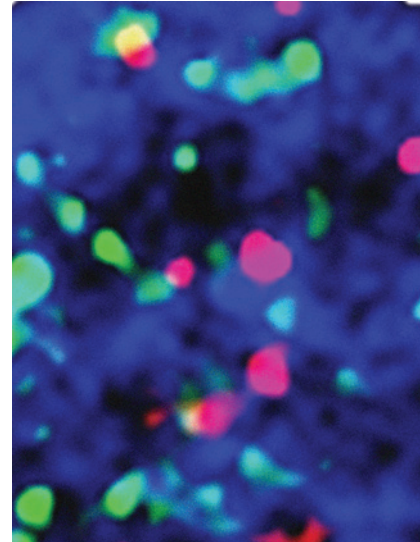
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6-thio-dG induces persistent telomere damage in conjunction with extensive genomic DNA damage in telomerase-positive cells several days after drug withdrawal. This is detail of an image of HFF + hTERT cells stained by FISH-immunofluorescence using a telomere specific PNA probe (red) and γ H2AX staining (green). Yellow fluorescent spots indicate co-localization of DNA damage (γ H2AX) and telomere signal (TelC).



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