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Produktinformation



Forschungsprodukte & Biochemikalien



Zellkultur & Verbrauchsmaterial



Diagnostik & molekulare Diagnostik



Laborgeräte & Service

Weitere Information auf den folgenden Seiten!
See the following pages for more information!



Lieferung & Zahlungsart

siehe unsere [Liefer- und Versandbedingungen](#)

Zuschläge

- Mindermengenzuschlag
- Trockeneiszuschlag
- Gefahrgutzuschlag
- Expressversand

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hTR(Texas Red)/CEN3q(FITC) FISH Probe

Catalog # : FA0507

規格 : [200 uL]

List All

Specification

Product Description:	Made to order FISH probes for identification of gene amplification using Fluorescent In Situ Hybridization Technique. (Technology)
Supplied Product:	DAPI Counterstain (1500 ng/mL) 250 uL
Storage Instruction:	Store at 4°C in the dark.
Origin:	Human
Source:	Genomic DNA
Notice:	We strongly recommend the customer to use FFPE FISH PreTreatment Kit 1 (Catalog #: KA2375 or KA2691) for the pretreatment of Formalin-Fixed Paraffin-Embedded (FFPE) tissue sections.
Regulation Status:	For research use only (RUO)

Application Image

Fluorescent In Situ Hybridization (Cell)

Applications

Fluorescent In Situ Hybridization (Cell)

 [Protocol Download](#)

Gene Information

Entrez GeneID: [7012](#)

Gene Name: TERC

Gene Alias: SCARNA19,TR,TRC3,hTR

Gene Description: telomerase RNA component

Omim ID: [127550](#), [602322](#), [609135](#)

Gene Ontology: [Hyperlink](#)

Gene Summary: Telomerase is a ribonucleoprotein polymerase that maintains telomere ends by addition of the telomere repeat TTAGGG. The enzyme consists of a protein component with reverse transcriptase activity, and an RNA component, encoded by this gene, that serves as a template for the telomere repeat. Telomerase expression plays a role in cellular senescence, as it is normally repressed in postnatal somatic cells resulting in progressive shortening of telomeres. Deregulation of telomerase expression in somatic cells may be involved in oncogenesis. Studies in mouse suggest that telomerase also participates in chromosomal repair, since de novo synthesis of telomere repeats may occur at double-stranded breaks. Mutations in this gene cause autosomal dominant dyskeratosis congenita, and may also be

associated with some cases of aplastic anemia. [provided by RefSeq]

Other -
Designations:

Related Disease

[Anemia, Aplastic Bone Marrow Diseases](#) [Breast Neoplasms](#) [Dyskeratosis Congenita](#)
[Fanconi Anemia](#) [Genetic Predisposition to Disease](#) [Heart Defects, Congenital](#)
[Leukemia, Myeloid, Acute](#) [Neural Tube Defects](#) [Pulmonary Fibrosis](#) [Scleroderma, Systemic](#)
[Syndrome](#)

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