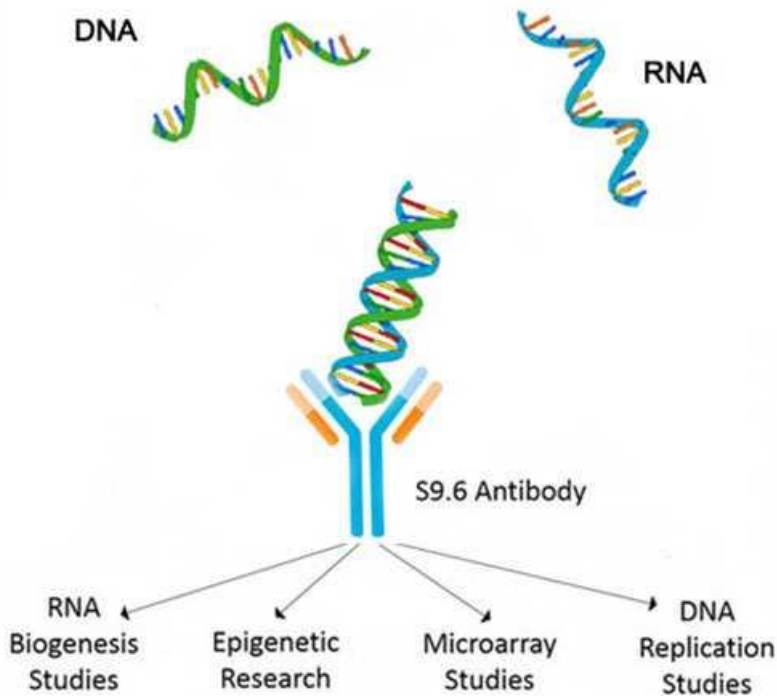


抗DNA-RNA杂种[S9.6]抗体,1mg(10x100ug)

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产品英文名称

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货号/SKU

ENH002

货号/规格

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库存与交货期

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产品基础信息

From the laboratory of Stephen H. Leppla, PhD, National Institute of Allergy and Infectious Diseases/NIH.

产品描述信息

Product Type:

Antibody

Name:	Anti-DNA-RNA Hybrid [S9.6]
Antigen:	S9.6 ϕ X174 bacteriophage-derived synthetic DNA-RNA antigen
Isotype:	Rabbit IgG
Fusion Tag(s):	Mouse Fab version contains His-tag
Clone Name:	S9.6
Reactivity:	High specificity and affinity for DNA/RNA hybrids and other A-form nucleic acid hybrids
Immunogen:	ϕ X174 bacteriophage-derived synthetic DNA/RNA
Purification Method:	Protein A/G
Buffer:	ENHOO1: PBS, 0.05% (w/v) Sodium Azide Ab01137- : PBS with 0.02% Proclin 300

Dot Blot Analysis: 0.2 μ g/mL.

Affinity Binding Assay: Clone S9.6 bound the DNA-RNA heteropolymer and poly(I)-poly(dC) equally, but 100-fold higher levels of poly(A)-poly(dT) were required to achieve a similar degree of binding. Single-stranded DNA, double-stranded DNA and RNA, and ribosomal RNA were not bound by clone S9.6 (Boguslawski, S.J., et al. (1986). J. Immunol Methods. 89(1):123-130).

Chromatin Immunoprecipitation (ChIP) Analysis: A representative lot detected increased DNA RNA hybrids at four actively transcribed genes upon shRNA-mediated knockdown of BRCA1 or BRCA2, but not PCID2 or RAD51 in HeLa cells (Bhatia, V., et al. (2014). Nature. 511(7509):362-365).

Chromatin Immunoprecipitation (ChIP) Analysis: A representative lot detected R-loops formed over beta-actin gene using HeLa chromatin preparation. RNase H treatment of the chromatin preparation prevented clone S9.6 from immunoprecipitating target chromatin fragments (Skourtis-Stathaki, K., et al. (2011). Mol. Cell. 42(6):794-805).

Chromatin Immunoprecipitation-sequencing (ChIP-seq) Analysis: A representative lot detected genome-wide distribution of DNA-RNA hybrids in budding yeast by ChIP-seq analysis (El Hage, A., et al. (2014). PLoS Genet. 10(10):e1004716).

Immunocytochemistry Analysis: Representative lots immunolocalized nuclear R loops by fluorescent immunocytochemistry staining of methanol-fixed H1 human embryonic stem cells (hESCs) and formaldehyde-fixed HeLa cells (Bhatia, V., et al. (2014). Nature. 511(7509):362-365; Ginno, P.A., et al. (2012). Mol. Cell. 45(6):814-825).

Immunoprecipitation Analysis: A representative lot immunoprecipitated in vitro transcribed R-loop substrate (DNA-RNA hybrid), but not double-stranded DNA (dsDNA) (Ginno, P.A., et al. (2012). Mol. Cell. 45(6):814-825).

See also: S9.6 Publications by Application

产品安全信息

Anti-DNA-RNA Hybrid [S9.6] Antibody - Publications by Application »Phillips DD, Garboczi DN, Singh K, Hu Z, Leppala SH, Leysath CE. The sub-nanomolar binding of DNA-RNA hybrids by the single-chain Fv fragment of antibody S9.6. J Mol Recognit. 2013 Aug;26(8):376-81.Boguslawski SJ, Smith DE, Michalak MA, Mickelson KE, Yehle CO, Patterson WL, Carrico RJ. Characterization of monoclonal antibody to DNA:RNA and its application to immunodetection of hybrids. J Immunol Methods. 1986 May 1;89(1):123-30.Yehle CO, Patterson WL, Boguslawski SJ, Albarella JP, Yip KF, Carrico RJ. A solution hybridization assay for ribosomal RNA from bacteria using biotinylated DNA probes and enzyme-labeled antibody to DNA:RNA. Mol Cell Probes. 1987 Jun;1(2):177-93.Miller CA, Patterson WL, Johnson PK, Swartzell CT, Wogoman F, Albarella JP, Carrico RJ. Detection of bacteria by hybridization of rRNA with DNA-latex and immunodetection of hybrids. J Clin Microbiol. 1988 Jul;26(7):1271-6.Casebolt DB, Stephensen CB. Monoclonal antibody solution hybridization assay for detection of mouse hepatitis virus infection. J Clin Microbiol. 1992 Mar;30(3):608-12.Hu Z, Zhang A, Storz G, Gottesman S, Leppala SH. An antibody-based microarray assay for small RNA detection. Nucleic Acids Res. 2006 Apr 13;34(7):e52.Székely L, Rákosi Z, Bálint BL, Kókai E, Imre L, Vereb G, Bacsó Z, Goda K, Varga S, Balázs M, Dombrádi V, Nagy L, Szabó G. Ribonucleoprotein-masked nicks at 50-kbp intervals in the eukaryotic genomic DNA. Proc Natl

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主要内容

该小鼠单克隆抗体抗 ϕ 174噬菌体衍生的合成DNA-RNA抗原产生，并识别各种长度的RNA-DNA杂交物。高灯：在检测R-Loopshigh特异性和对DNA-RNA杂交物的亲和力不交叉反应的情况下有用已经观察到单链DNA或双链DNA或双链DNA的交叉反应（~5倍），用于富含Au的双链RNA.high亲和力结合，用于LightRecomant的8,10,15和23个碱基对的杂交种来自我们姐妹公司的版本，绝对抗体：使用来自杂交瘤S9.6DNA-RNA杂种的可变区（即特异性）的绝对抗体的重组平台制造是真核细胞内的自然发生，这些杂种在位点增加高转录活性，例如在转录起始，抑制和伸长期。因为RNA-DNA杂交种影响基因组不稳定性，所以S9.6抗体是有用的试剂，以帮助研究在DNA复制或其他细胞过程中通过这些杂种形成的R圈和病变的后果。此外，S9.6抗体可有效识别微阵列研究的RNA-DNA杂交。通过斯蒂芬H.Leppla，博士学位，国家过敏和传染病研究所的实验室。

厂牌介绍

关于Kerafast Inc.

Kerafast 是一家位于波士顿的试剂公司，其主要使命是为QuanQiu科学界提供易于使用的独特实验室研究工具。我们的产品组合包括细胞系、抗体、小分子、染料等，其中许多在其他地方无法获得。自 2011 年成立以来，来自全球 190 多个机构的研究人员通过我们的在线平台提供了他们的创新试剂，无需通过传统的材料转让协议流程即可快速获取材料。

我们处理提供实验室的所有销售和运输物流，并从每次销售中返还丰厚的特许权使用费。因此，我们帮助提供实验室节省时间和资源，同时为进一步研究提供额外资金。采购科学家可以更轻松地发现和获取其他地方通常无法获得的独特试剂，同时还可以资助其他研究人员的工作。这创建了一个QuanQiu科学家社区，他们贡献和获取Reagent for the Greater Good，以加速他们自己的研究以及整体科学进步。

2018 年，Kerafast 与 [Absolute Antibody](#) 合并，后者是一家总部位于英国的公司，其愿景是为所有研究人员提供重组抗体技术。[此次合并](#)将两家公司聚集在一起，共同致力于改善科学界可用的研究工具的选择。

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