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3, Respectively. The PCR Conditions, With TaKaRa LA Taq Polymerase And GC Buffer

I, Were Essentially The Same As Those Described Previ-ously (10). The Nucleotide Sequence Of The Amplified DNA Fragment (507 Bp From The 5-end Of Pld1) Was 12th, 2024

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And Cloned Into The Same Sites Of PcDNA3.1/Hygro (Invitrogen, Carls-bad, CA), Resulting In PcDNA3.1-C4ST-1 (the Name Of C4ST-1 Was Given After The Determination Of Acceptor Specificity). The Second Gene Was Initially Identified In AA233362 And AA777237 Derived F 9th, 2024

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Proc. Natl. Acad. Sci. USA92 (1995) Withphenol, DNAwasprecipitatedwithethanol Andspooled Onto A Glass Rod. Construction And Screening Ofa AExpression Library. S. Solfataricus Genomic DNAwaspartially Digested With Sau3A And Fragments From 0.5 To 2.5 Kb Were Ligated To BamHI- Digested And Dephosphorylated AZAPExpress Arms (Strat- Agene); Packagingwasperformed According To The ... 2th, 2024

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And 1 Kb Plus DNA Ladder, Invitrogen (catalogue No. 10787-018) Was Used As Marker To Determine The Size Of DNA Fragment. DNA Recovery The Amplified DNA Fragment Verified With Gel Electrophoresis Was Extracted Using TM Purelink Quick Gel Extraction Kit From Invitrogen (catalogue No. K2100-12). Four Tubes Of PCR Reaction (25 16th, 2024

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Gao Et Al. 3149 Table 1. Primers Used In This Study. Primers Name Sequence(5' To 3') SMART IVTM Oligonucleotide AAGCAGTGGTATCAACGCAGAGTGGCCATTACGGCCGGG CDS III/3 ... 13th, 2024

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4Boyer, R F (1986) Modern Experimental Biochemistry, Addison- Wesley Publishing Co, Reading, MA, USA -5Maniatis, T, Fritsch, E F, And Sambrook, J (1982) Molecular Cloning. A Laboratory Manual, Cold Spring Harbor Laboratory, New York, NY, USA 6Clark, John M, Switzer, R L (1977) Experimental Biochemistry, W HCited By:

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