



PLCG2 (NM_002661.5) - cDNA + Protein - 2024-05-19

AGAAGTAGCG AGCGCCGGCG GCGGAGGGCG TGAGCGGGCG TGAGTGACCC -132
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nLysGluAsp CysCysPheT hrIleLeuTy rGlyThrGln PheValLeuS 107

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***PLCG2* (NM_002661.5) - cDNA + Protein - 2024-05-19**

