

Product Information**pAd1127-06**

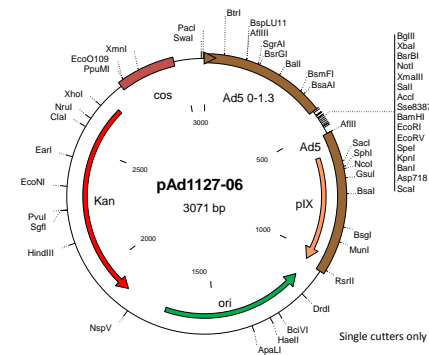
Research use only

Catalog No: QP-24**Lot No:** 1002**Contents:**pAd1127-06 plasmid DNA, 20 µg,
1 µg/µL in TE pH 7.5**Storage:** -20°C**Features and Applications:**pAd1127-06 is a vector designed for inserting expression cassettes in place of the E1 region of the Ad5 genome and to manipulate the pIX promoter and coding region. It contains *PacI* and *SwaI* sites flanking the first 440 base pairs from the Ad5 genome (including

the left ITR and packaging signal), a multiple cloning site, and the pIX coding region. Expression cassettes inserted into the multiple cloning site should contain a promoter, coding sequence and a polyA signal. The sequences encompassing the kanamycin-resistance gene, the λ cos site, the adenovirus 0-1.3 map units, the multiple cloning site and the pIX coding sequence are flanked by two *SfiI* restriction sites. These sites generate non-symmetrical sticky ends suitable for directional cloning with the other AdenoQuick2.0 plasmids (pAd1128, pAd1129, pAd1130, and their derivatives).

pAd1127-06 is a derivative of pAd1127-02, in which the packaging signal has been extended from psn 350 to psn 440 (in the Ad5 genome), to include all 7 packaging "A" repeats (I, II, III, IV, V, VI, and VII). The complete packaging signal region might confer a growth advantage to the virus, according to Youil *et al* (Human Gene Therapy 14: 1017-1034). Because of the size of the

E1 deletion (440-3510), the vectors generated from pAd1127-06 have minimal or no homology with the Ad5 sequences inserted in the chromosome of the helper cells such as PER-C6, thereby minimizing the probability of RCA generation.

Selection:*prokaryotic* – kanamycin 25 µg/mL**Replication:***prokaryotic* – pUC ori

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