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- (本文编辑: 俞燕)

· 消息 ·

本刊关于实验动物样本数的规定

为了提高杂志的学术质量,《中国当代儿科杂志》决定从2016年8月起,对凡是在本刊刊出的动物实验研究论文中实验动物样本数做出如下规定,请广大作者注意并严格执行。对于未达到以下要求的动物实验性文章投稿,本刊则一律不再接收。

(1) 活体动物实验每组动物数不得少于8只;

(2) 对于细胞或分子水平检测方法,可适当放宽对样本数的要求,如:PCR法检测mRNA水平,每个单元组的样本数不得少于6例;Western blot法检测蛋白水平,每个单元组的样本数不得少于3例;

(3) 组织切片,要求每个样本组织的切片数不得少于3张,每张切片随机观察视野不得少于5个。

《中国当代儿科杂志》编辑部
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