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Highly sensitive detection of Epstein-Barr virus-infected cells by EBER flow FISH

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1. 文部科学省科学研究費補助金（萌芽研究，2006-2008） 家族性血球貪食症候群の迅速診断法の確立とその類縁疾患の解析（研究代表者：秦 大資，研究分担者：堀内久徳（京都大学））

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