PROTOCOL



Dharmacon[™] Edit-R[™] Cas9 nuclease protein and synthetic guide RNA transfection protocol

The following is a protocol for transfecting <u>Edit-R™ Cas9 Nuclease protein</u> <u>NLS</u>, (Cat #CAS11XXX) with synthetic guide RNA into cultured mammalian cells using DharmaFECT™ transfection reagents (Cat #T-20XX-xx). Synthetic guide RNA can be either Edit-R synthetic tracrRNA (Cat #U-002005-xx) complexed with crRNA (predesigned or custom), or Edit-R synthetic sgRNA (custom). For a more detailed protocol please see the <u>Technical Manual</u>.

The protocol is written for transfection into 96-well tissue culture plates (100 μL final volume).



96-well protocol			
Day 1			
Cell plating	Seed cells at a density that gives 70-90% confluency on the next day		
Day 2			
Prepare working solutions of reagents for transfection	Synthetic guide RNA	Dilute and mix crRNA and tracrRNA to a working concentration of 2 μ M in 10 mM Tris-HCl pH 7.4 or Dilute synthetic sgRNA to a working concentration of 2 μ M in 10 mM Tris-HCl pH 7.4	
	Cas9 nuclease protein	Dilute Cas9 nuclease protein to a working concentration of 2.5 μ M in serum-free medium	
Combine working solutions for transfection mixture		For one well	For mulitple wells
	Tube 1		
	Synthetic guide RNA	2.5 μL	_ μL
	Cas9 nuclease protein	1 μL	_μL
	Serum-free medium	To 50 μL	_μL
Prepare working solution of DharmaFECT for transfection	Tube 2		
	DharmaFECT transfection reagent	0.1–0.8 µL	_ µL
	Serum-free medium	To 50 μL	_ µL
	Incubate at room temperature for 5 minutes before next step		
Combine transfection mixture	Combine Tube 1 and Tube 2 and carefully mix by pipetting		
	Incubate at room temperature for 20 minutes before next step		
	Total	100 µL	_μL
Transfect cells	Replace growth medium on cells with 100 μL of transfection mixture		
Return to full media	After 14-18 hours; replace transfection mixture on the cells with typical cellular growth medium		

If you have any questions, contact

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