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Insert + Vector DNA Ligation

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Protocol status: Working

We use this protocol and it's working

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Abstract

Pipette mix the following components:

	Double Digested DNA	ng/uL	size (kb)	bp ratio	1	3	5	I:V Ratio
	AXXX Vector, linearized + dephos			#VALUE!		#VALUE!		ng insert
	AXXX Gen, digested					#VALUE!		uL insert
	Volume (uL)	Ratio	Insert	Vector	T4 Ligase	Water	T4 Buffer	Total
	Ligation 1 Vector only	vector	0		1	17	2	20
	Ligation 2 Vector + Gene ligation	3 to 1	#VALUE!		1	#VALUE!	2	20

Nur die grün markierten felder müssen ausgefüllt werden. Die Formeln berechnen das optimale 3:1 verhältnis von Vektor zu gen. Check wie viel DNA insgesamt eingesetzt wird. Vektor ist bei mir immer so um 25-30 ng und totale DNA menge überschreitet nicht 100 ng dann sollten gute ergebnisse erzielt werden.

Incubate 60 min at room temperature, and/or overnight at 16°C (*higher efficiency*).

Optional: inactivate ligase by incubating for 5 min in 70°C heat block.

Transform into E. coli

