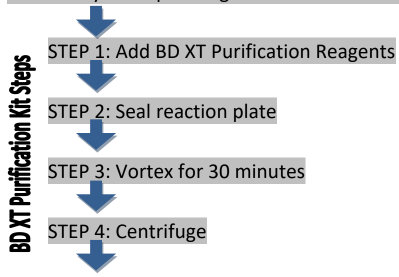


Quick Reference Guide

ADS BD XT Purification Kit

Sequencing workflow

Perform cycle sequencing with BD XT Purification Kit



Run purified samples on your DNA Analyzer

Overview

The BD XT Purification Kit sequesters cycle-sequencing reaction components such as salt lons, unincorporated dye terminators, and dNTPs to prevent their coinjection with dye-labled extension products into a CE DNA analyzer. The BD XT Purification reagents can be pipette separately and sequentially into reaction plate, or premixed together before being pipette into a reaction plate.

Ordering Information

Refer to the BD XT Purification Kit Protocol for recommended vortexers and required accessories.

	Approximate	Volume of Each		
Kit Size	Number of 20-μL Reactions	Resin	Solution	Part Number
2-mL	100	2	9	
20-mL	1000	20	90	
50-mL	2500	50	225	
800-mL	40000	800	3600	

Important Tips

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- When you pipette directly from the Solution bottle:
 - Before pipetting, mix the Solution until homogeneous,
 - Use wide-bore pipette tips,
 - Avoid pipetting near the surface of the liquid,
 - When you seal the reaction plate, verify that each well is sealed.
- To achieve optimum performance, use a recommended vortexer and follow the protocol when you vortex the reaction plate.
- When you load plates into the CE instrument:
 - Do not heat-denature or use Formamide with samples containing BD XT Purification reagents.
 - Use the ABI run modules specified for your instrument and plate type.

Procedure for Sequential Pipetting

STEP		ACTI	ON					
4	Centrifuge	Follow the cycle-sequencing	protocol. When the reaction is complete,					
T	the	centrifuge the reaction plate for 1 minute to spin down plate contents.						
	sequencing	IMPORTANT! You may need to decrease the amount of DNA template in the						
	reaction	sequencing reactions to compe	nsate for increased signal strength. See "DNA					
	plates.	Quantity Guidelines" on page 6.						
_	Add the	To each well of the reaction pl	ate, add the volume of the Solution specified					
	Solution to	below, using a conventional pipe	tte tip.					
	the reaction	Make sure there are no parti	culates in the Solution before pipetting. If					
	plates	particulates are present, heat th	e Solution to 37°C and mix to redissolve. Cool					
		to room temperature before using	ng.					
		Plate Type and Reaction	Volume of the Solution/Well (μL)					
		Volume/Well	volume of the 30idtion, wen (με)					
		384-well, 5 μL 22.5						
		96-well, 10 μL 45.0						
		96-well, 20 μL 90.0						
		IMPORTANT! For 384-well reactions with reaction volumes less than 5 μ L, add water to bring the volumes to 5 μ L before adding the Solution. For 96-well reactions with reaction volumes less than 10 μ L, add water to bring the volume to 10 μ L before adding the solution.						
	Add the Resin	Add the Resin:						
3	to the	a. Vortex the resin at maximum speed for at least 10 seconds, until it is						
	reaction	homogeneous						
	plates using a	b. Using a wide-bore pipette tip, add to the reaction plate the volume of						
	wide-bore	the Solution specified be	low.					
	pipette tips	Plate Type and	Volume of Resin/Well (μL)					



STEP	ACTION							
			Reaction Volume/Well					
			384-well, 5 μL	5.0				
			96-well, 10 μL	10.0				
			96-well, 20 μL	20.0				
Л	Seal, vortex,	Follow t	Follow the instructions in "After Pipetting Is Complete" on page 4.					
4	4 load and run							
	the plates							

Procedure for Premix Pipetting

Note: The premix is stable only for 5 days. Make only the volume of premix that you will use in 5 days.

Note:	The premix is stable only for 5 days. Make only the volume of premix that you will use in 5 days.						
STEP			ACTI	ON			
4	Calculate the	Based on	your plate and	reaction size,	calculate the	volume of the	
1	required volume of	Solution an	d Resin needed.				
	the Purification	Note: All volumes below include an additional 10% to account for dead					
	reagents.	volume in the reagent trough.					
		For 384-well plate, 5-μL reactions:					
		Reagent	Volume/Well	Volume/Plate	Number of	Final Volume	
		Reagent	(μL)	(μL)	Reactions	Needed	
		Solution	24.75	9504			
		Resin	5.5	2112			
		For 96-well	plate, 10-μL rea	ections:			
		Danasat	Volume/Well	Volume/Plate	Number of	Final Volume	
		Reagent	(μL)	(μL)	Reactions	Needed	
		Solution	49.5	4752			
		Resin	11	1056			
		For 96-well	plate, 20-μL rea	ections:			
		Decemb	Volume/Well	Volume/Plate	Number of	Final Volume	
		Reagent	(μL)	(μL)	Reactions	Needed	
		Solution	99	9504			
		Resin	22	2112			
	Combine the	Cambinath	ne Solution and I	Docine			
2					mum speed fo	or the least 10	
	reagents to create		onds, until it is h		num speed n	or the least 10	
	the premix		•	ŭ	a graduated o	dinder add the	
			_	e of Resin to a cle	_	ylinder, add the	
			•			he liquid	
		IMPORTANT! Avoid pipetting near the surface of the liquid.c. Using a conventional pipette tip or a graduated cylinder, add the					
			-		-	itainer with the	
			sin.	ie or the solution	on to the con	italliei with the	
		Ne:)III.				



STEP		ACTION						
		Make sure there are no particulates in the Solution before						
		pipetting. If particulates are present, heat the Solution to 37°C						
		and mix to redissolve. Cool to room temperature before using.						
		d. Mix the reagents until homogeneous.						
		Note: The premix can be stored in a clean,	capped container at 4°C for up					
		to 5 days.						
2	Centrifuge the	Following the cycle-sequencing protocol. \	When the reaction is complete,					
3	sequencing	centrifuge the reaction plate for 1 minute t	o spin down plate contents.					
	reaction plates.	IMPORTANT! You may need to decrease t	he amount of DNA template in					
		the sequencing reactions to compensate	for increased signal strength.					
		See "DNA Quantity Guidelines" on page 6.						
Л	Add the premix to	Using a conventional pipette tip, add to each well of the reaction plate the						
4	the reaction plates.	volume of the thoroughly mixed premix specified below.						
		IMPORTANT! For 384-well reactions with reaction volumes less than 5 μL,						
		add water to bring the volumes to 5 µL before adding the premix. For 96-						
		well reactions with reaction volume less t	han 10 μL, add water to bring					
		the volume to 10 µL before adding the prer	nix.					
		Plate Type and Reaction Volume/Well	Volume of Premix/Well (μL)					
		384-well, 5 μL	27.5					
		96-well, 10 μL	55.0					
		96-well, 20 μL	110.0					
		IMPORTANT! Mix the premix as needed	to maintain a homogeneous					
		solution. Dispense the premix within 1 minutes of aspiration to avoid						
		separation of the reagents in the pipette tip.						
	Seal, vortex, load,	Follow the instructions in "After Pipetting Is	s Complete" on page 4.					
5	and run the plates							

After Pipetting Is Complete

STEP	ACTION						
4	Seal the reaction	Seal the plate, using:					
T	plates.	 A heat seal at 160oC for 2 second 	onds				
		or					
		 MicroAmp Clear Adhesive Film 	ns or any other go	ood adhesive films			
		Verify that each well is sealed.					
		IMPORTANT! If you are using an ABI 3730 DNA Analyzer and plan to use					
		direct injection, only ABI Heat Seal Film for Sequencing and Fragment					
		Analysis Sample Plates is supported					
	Vortex the reaction	Vortex the reaction plate for 30 minutes using the following conditions:					
2	plates.	Vortexer	Plate Type	Speed			
		Digital vortex-Genie 2	96-well	1800 rpm			

AD.	Tour heliable Sariger Sequencing heagent Farther					
STEP			AC	TION		
					384-well	2000 rpm
		Eppendorf MixMate			384-well	2600 rpm
		IKA MS3 Digital			Either	2000 rpm
		IKA Vortex	3		Either	Setting 5
			roMixer E-36		Either	Maximum
		L	ntific Vertical Sh		Either	Setting 100
					ise vortexing afte	er 1 minute to verify
			ntents are we			
2	Centrifuge the	In a swingir	ng-bucket cent	trifuge, spin t	he plate at 1000	x g for 2 minutes.
3	reaction plates					
	Prepare the plates	Place the re	eaction plate i	n The CF inst	rument. (To stor	re and run the plate
4	for the instrument	later, see s	•	02		o and ran the plate
_	run.	Plate				
		Туре	Instrument	Seal	Instr	ructions
		384-well	3730 /	Heat seal	Place directly in	n the instrument.
			3730xl	MicroAmp		e clear adhesive
				Clear		e with a heat seal,
				Adhesive	and then place in the	
				Film	instrument.	
		Transfer 10 μL of				
				it to a clean plate,		
				-	a septa mat, place	
		in instrument				
			3100/	Either		of supernatant to
			3100, 3100Avant,	Little	•	over with a septa
			3130/		mat, then place	•
			3130xl, or		instrument.	. III CIIC
			310		moer america	
			Genetic			
			Analyzer			
		96-well	3730 /	Heat seal	Place directly in	n the instrument
			3730xl	MicroAmp		al, replace with a
				Clear	septa mat, plac	•
				Adhesive	instrument.	in the
				Film		
			3100/	Either	Remove the sea	al, replace with a
			3100/ 3100Avant	2101101	septa mat, plac	•
			or 3130/		instrument.	
			3130xl			
	1	I [313071	1	1	



STEP	ACTION						
		310)	Either	Transfer 10 μL of supernatant to		
		Ger	netic		a clean plate, cover with a septa		
		Ana	ılyzer		mat, then place in the		
					instrument.		
	Calaattlaa	Calast the are		DiaDara VA			
5	Select the	1			erminator run module for your		
	appropriate run	instrument and	plate type	2.			
	module	Note: Use standard run modules if you transferred the supernatant to a					
		clean plate after centrifuging.					
	Run the reaction	Run the plate.					
6	plates	If the reaction plates are not run immediately, you can store them under					
		the following conditions:					
		Room temperature – Plates sealed with heat seal film, adhesive film,					
		or septa for up to 48 hours at room temperature (20 to 25°C).					
		Refrigerated storage – Plates sealed with heat seal film or adhesive					
		film for up to 10 days at 4°C (recommended).					
		Frozen storage – Plates sealed with heat seal film or adhesive film for					
		up to 10 day	•				

DNA Quantity Guidelines

DNA sequencing reactions purified with the BD XT Purification Kit result in high signal strength when analyzed on a DNA sequencer. Therefore, when you prepare sequencing samples for purification with the BD XT Purification reagents, you may need to decrease the amount of DNA template in the sequencing reactions to keep the fluorescence signals on scale during analysis. Use the following table as a guide to the amount of template DNA for the initial cycle sequencing.

IMPORTANT! If you decrease the template concentration, also decrease the amount of any template controls proportionately. For example, if you run a pGEM control, dilute if 1:2 or 1:4 and add only 1 to 2 μ L.

Template Type	DNA Quantity/Reaction (ng)	Template Type	DNA Quantity/Reaction (ng)
PCR products		Other types of template	
100 to 200 bp	0.5 to 3	Single-stranded DNA	10 to 50
200 to 500 bp	1 to 10	Double-stranded DNA	50 to 300
500 to 1000 bp	2 to 20	Cosmid or BAC DNA	200 to 1,000
1000 to 5000 bp	5 to 40	Bacterial genomic DNA	1,000 to 3,000
>2000 bp	10 to 50		