

Product Information			
Extracta Plus DNA, 10 rxn			
Part Number	95213-010		
Number of Reactions	10 Reactions		
Storage Temperature	15⁰C to 25ºC		
Lot Number	169036764		

Product Description:

Extracta Plus DNA kits offer rapid purification of total DNA from a variety of samples sources. Genomic, mitochondrial and pathogenic DNA can be collected from fresh/frozen animal cells, tissues, blood, and bacteria. The Extracta Plus Spin column workflow enables simultaneous processing of multiple samples into purified DNA suitable for PCR, Southern blotting, RAPD, AFLP and next generation sequencing.

Purified DNA is eluted in low salt buffer or water. The fragments purified DNA from this kit will predominate in the 30 kb range, with fragments as large as 50 kb and as small as 100 bp being recovered.

Component Part Numbers:

1123997	Proteinase K	1.25 ml
1123999	Buffer EPDTL	1.8 ml
1124002	Buffer EPDL	2 ml
1124005	Buffer EPDW1	5 ml
1124008	Buffer EPDW2	5 ml
1124011	Buffer EPDE	2 ml
1124014	Extracta Plus-Spin Column	10 pc
1124016	Collection Tubes 2ml	20 pc

Product Specifications			
95213-010			
Assay	Functional Assay		
Result	Pass		

Quality Control Analysis and Specifications:

Membrane control

Membrane binding capacity was tested by determining recovery of genomic DNA.

Yield of \ge 50 μg was obtained per Spin Column.

Pass

As tested by PCR, no PCR inhibitors were released from the membrane.

Pass

Limitations of Use

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Buffers

Conductivity and pH of buffers and solutions were tested and were within the ranges below:

	Conductivity (mS/cm)	рН	
EPDL	38.40 – 45.40 (1/10 dilution)	5.90 – 6.10	Pass
EPDTL	20.50 – 28.20	8.10 – 8.50	Pass
EPDW1	50.20 - 59.80 (1/10 dilution)	_	Pass
EPDW2 ^t	28.60 – 34.20	7.40 - 7.60	Pass
EPDE	0.24 - 0.31	8.90 - 9.10	Pass

^t Contains sodium azide as a preservative

Buffer EPDL was tested by preparation of genomic DNA from 200 μ l of whole human blood. Average yield of genomic DNA was \geq 90% of reference yield, with an A_{260}/A_{280} ratio of 1.70 – 2.00.

Functional absence of RNase was checked for Buffer EPDW2[†].

Proteinase K

Activity was determined by incubation of hemoglobin substrate solution with Proteinase K in water, at 30°C for 10 min, followed by the determination of acid-soluble tyrosine at 750 nm.

Pass

Absence of DNase and RNase activity was checked by incubation of DNA or RNA, with Proteinase K in enzyme incubation buffer at 37°C.

pass

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