Application	Amount	Volume	Conc.	Quality	Size	Comment
		(μL)	(ng/μL)		(bp)	

Illumina Short-Read Sequencing

DNA Sequencing	General rema	rks and requir	ements for DN	IA samples:							
. 5	- Buffer	: TRIS-HCL 10	mM or lowTE								
	- 260/2	80 ratio 1.8-2.	0 (or according	g to quality column)							
	- In case of genome sequencing: high molecular for best results										
	All samples of an order must be adjusted to a uniform concentration within the specifications.										
	Samples that do not meet the requirements listed here cannot be processed and will be rejected!										
Whole Genome - Low input (incl. PCR)	150 – 300ng	>=15	>10	see general remarks	-	FFPE possible with constraints in					
The continue Lew Impact (men's eny				above		output quality					
Whole Genome - PCR Free	>300 ng	>=15	>20	see general remarks	-						
				above							
Bacterial genome sequencing	15 ng	>=15	1	see general remarks							
				above							
Whole Exome Sequencing	>75 ng	>=15	5	see general remarks	-	FFPE possible if amount based on					
				above		qPCR measurement is sufficient					
Gene Panel	600 ng	>=15	40	260/280 ratio 1.7-2.2	-	-					
				260/230 ratio 1.2							
Amplicons	>75 ng	>=15	5	260/280 ratio >1.5	-	less material possible on request					
, , , , , , , , , , , , , , , , , , ,											
Microbiome 16S	150 ng	>=15	10	260/280 ratio >1.5	-	less material possible on request					
Microbiome Shotgun Metagenomics	75 ng	>=15	5	260/280 ratio >1.5	-	less material possible on request					

Application	Amount	Volume	Conc.	Quality	Size	Comment
		(μL)	(ng/μL)		(bp)	

Illumina Short-Read Sequencing

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	Seque	9

General remarks and **requirements** for totalRNA samples:

- Buffer: nuclease free water
- DNAse treated and cleaned up
- 260/280 ratio >2.0
- RQN>=8; RQN Δ between samples <1
- ultra low input (< 1 ng total amount) possible, but must be planned beforehand

All samples of an order must be adjusted to a uniform concentration within the specifications. Samples that do not meet the requirements listed here cannot be processed and will be rejected!

Whole Transcriptome – standard input	>375 ng	>=15	25 - 80	see general remarks above	-	-
Whole Transcriptome - low input	>15 ng	>=15	1 - 5	see general remarks above	-	very low input amount may lead to constraints in output quality
Expression Profiling (mRNA) – standard input	>375 ng	>=15	25 - 80	see general remarks above	-	-
Expression Profiling (mRNA) – low input	>15 ng	>=15	1 - 5	see general remarks above	-	very low input amount may lead to constraints in output quality
3'Prime RNA Seq -standard input	>375 ng	>=15	25 - 80	see general remarks above		-
3'Prime RNA Seq -low input	>15 ng	>=15	1 - 5	see general remarks above		very low input amount may lead to constraints in output quality
Small RNA Profiling (miRNA, Inc-RNA etc.)	50 ng	>=15	50	see general remarks above	-	Low input possible on request

Application	Amount	Volume (μL)	Conc. (ng/μL)	Quality	Size (bp)	Comment		
Sequencing of prepared NGS libraries	50 ng	>=15	5	No primer dimer residuals	; -	Buffer: TRIS-HCL 10 mM		
10X Single-Cell RNA-Sequencing								
Single-Cell Transcriptomics	1.000 – 30.000 cells /sample	50 μΙ	500-1.000 cells/μl	Vitality >70 %	-	-		
Spatial Transcriptomics	Tissue sections placed on Visium slides, RNA extracted from tissue section RIN >7							
Bionano Optical Mapping								
Bionano Saphyr Chip	1 μg	-	>36	260/280 = 1.8 260/230 = 2.0-2.2	Megabase range	· -		

A	Application	Amount	Volume	Conc.	Quality	Size	Comment
			(μL)	(ng/μL)		(bp)	

Pacific Biosciences Long-Read Sequencing

Revio System

- ·							
Transcriptome	Iso-Seq	>2-3 μg	>15	>200	RIN 8-10 260/280 = 2.0 260/230 = 2.2	-	DNase digested, buffer: RNase- free water
Whole Conome	Coguanaina						
Whole Genome .			. 420	70	250/200 4.0	. 501 1	La constant and the state of th
HiFi Rea	ds – Standard Protocol	≥8 µg	>120	~70	260/280 = 1.8 260/230 = 2.0-2.2	>50kb	homogenous HMW DNA, RNase digested, buffer: EB, DNA size distribution: 80% ≥ 50Kb
	HiFi Reads – Low Input	>3 μg	>50	60	260/280 = 1.8 260/230 = 2.0-2.2	>50kb	homogenous HMW DNA, RNase digested, buffer: EB, DNA size distribution: 80% ≥ 50Kb
HiFi R	eads – Ultra-Low Input	>50 ng	>15	>3	260/280 = 1.8 260/230 = 2.0-2.2	>50kb	homogenous HMW DNA, RNase digested, buffer: EB DNA size distribution: 80% ≥ 50Kb
Trageted Sequer	ncing						
	Amplicon Sequencing	2-4 μg (depends on size)	50	-	-	-	clean, target-specific, buffer: EB
Metagenomics							
	Multiplexed Microbial	>2-3 µg per sample	20-100	-	260/280 = 1.8 260/230 = 2.0-2.2	>30kb	Homogenous HMW DNA, RNase digested, buffer: EB DNA size distribution: 80% ≥ 30Kb

Application	Amount	Volume (μL)	Conc. (ng/μL)	Quality	Size (bp)	Comment
Kinnex workflows						
Full-lenght RNA	2-3 μg (depends on # samples to concatenate)	50	> 200	RIN 8-10 260/280 = 2.0 260/230 = 2.2	-	DNase digested, buffer: RNase- free water up to 8-fold concatenation
16S rRNA Amplicons	>500 ng - 3 µg (depends on size)	50	-	-	~ 1500 bp	Purified amplicons up to 12-fold concatenation (~ 19 Kb) Applicant must order Kinnex 16S Forward and reverse primers. From any oligo vendor, HPCL purification is recommended.
Single-cell RNA	50 ng of 10x Chromium 3` or 5` single cell DNA.	25	-		500-1500 bp	Purified up to 16-fold concatenation Single-cell cDNA generated using the 10x Chromium Next GEM Single Cell 3' kit v3.1/5' kit v2 standard throughput. cDNA in the respective buffer.

Application	Amount	Volume (μL)	Conc. (ng/μL)	Quality	Size (bp)	Comment
Sequel II/IIe Systems						
Transcriptome (Iso-Seq)	>2 μg	>15	>130	RIN 8-10 260/280 = 2.0 260/230 = 2.2	-	DNase digested, buffer: RNase free water
Whole Genome Sequencing						
HiFi Reads – Standard Protocol	>6 µg	>120	50	260/280 = 1.8 260/230 = 2.0-2.2	>50kb	homogenous HMW DNA, RNase digested, buffer: EB DNA size distribution: 80% ≥ 50Kb
HiFi Reads – Low Input	>1 μg	>50	20	260/280 = 1.8 260/230 = 2.0-2.2	>50kb	homogenous HMW DNA, RNase digested, buffer: EB DNA size distribution: 80% ≥ 50Kb
HiFi Reads — Ultra-Low Input	>30 ng	>15	2	-	>50kb	homogenous HMW DNA, RNase digested, buffer: EB DNA size distribution: 80% ≥ 50Kb
Continous Long Reads	>6 μg	>120	50	260/280 = 1.8 260/230 = 2.0-2.2	>50kb	homogenous HMW DNA, RNase digested, buffer: EB
Amplicon Sequencing	500 ng – 3 μg (depends on size)	50	-	-	-	clean, target-specific, buffer: EB
Multiplexed Microbial	>1 µg per sample	20-100	-	260/280 = 1.8 260/230 = 2.0-2.2	>30kb	-

Application Amo	unt Vo (μL	lume)	Conc. (ng/μL)	Quality	Size (bp)				
oxford Nanopore Long-Read Sequencing									
Transcriptome									
Direct mRNA Sequencing	>500 ng polyA+ or	>12	>40	260/280 = 2.0 260/230 = 2.0-2.2	-	DNase digested, buffer: RNase free water			
	1.5 μg Total- RNA	>12	>125						
cDNA Sequencing	>200 ng polyA+ or	>10	>20	260/280 = 2.0 260/230 = 2.0-2.2	-	200 ng cDNA can be used as input DNase digested, buffer: RNase free			
	>1 µg Total- RNA	>10	>100			water			
cDNA PCR Sequencing	>20 ng polyA+ or	>15	>1.3	260/280 = 2.0 260/230 = 2.0-2.2	-	DNase digested, buffer: RNase free water			
	>600 ng Total- RNA	>15	>40						
Whole Genome Sequencing									
Ligation Sequencing	>2 µg	>50	>40	260/280 = 1.8 260/230 = 2.0-2.2	>30 kb	homogenous HMW DNA, RNase digested, buffer: EB			
Ligation Sequencing with Size-Selection	>10 µg	>50	>200	260/280 = 1.8 260/230 = 2.0-2.2	>30 kb	homogenous HMW DNA, RNase digested, buffer: EB			
Ligation Sequencing Whole-Genome Amplification	>1 ng	>10	>0.1	260/280 = 1.8 260/230 = 2.0-2.2	>30 kb	homogenous HMW DNA, RNase digested, buffer: EB			
Native Barcoding	>1 µg per sample	>50	>20	260/280 = 1.8 260/230 = 2.0-2.2	>30 kb	homogenous HMW DNA, RNase digested, buffer: EB			

>750

>10

>70

>30

260/280 = 1.8

260/280 = 1.8

260/230 = 2.0-2.2

260/230 = 2.0-2.2

>100

kb

>30

kb

homogenous UHMW DNA,

inquire about EEB buffer)

homogenous HMW DNA,

RNase digested, buffer: EB

RNase digested, buffer: EEB (please

Ultra Long Reads >50 µg

>300 ng

Rapid - gDNA

Application	Amount	Volume	Conc.	Quality	Size	Comment
		(μL)	(ng/μL)		(bp)	

Oxford Nanopore Long-Read Sequencing

Amplicon Sequencing						
Amplicons by Ligation	>300 ng per sample	>20	>15	260/280 = 1.8 260/230 = 2.0-2.2	-	clean, target-specific, buffer: EB
Rapid - 16S	>20 ng	>20	>1	260/280 = 1.8 260/230 = 2.0-2.2	-	clean, target-specific, buffer: EB

Sanger Sequencing

Full Service						
Plasmid	>500 ng	-	100 – 250	260/280 = 1.8 260/230 = 2.0-2.2	<5 kb	Buffer: 10 mM Tris/HCl or water Primer concentration 10μM
	>800 ng	-	150 - 600	260/280 = 1.8 260/230 = 2.0-2.2	5 kb – 15 kb	Buffer: 10 mM Tris/HCl or water Primer concentration 10μM
	>1,5 μg	-	>600	260/280 = 1.8 260/230 = 2.0-2.2	>15 kb	Buffer: 10 mM Tris/HCl or water Primer concentration 10μM
Full Service / Xpress Service						
PCR-product	max. 10 ng	-	<2,5	260/280 = 1.8 260/230 = 2.0-2.2	<100 bp	purified PCR products Primer concentration 10μM
	50 ng	-	< 5ng	260/280 = 1.8 260/230 = 2.0-2.2	100 bp – 1 kb	purified PCR products Primer concentration 10μM

Application	Amount	Volume (μL)	Conc. (ng/μL)	Quality	Size (bp)	Comment
	400 ng	-	20 - 50	260/280 = 1.8 260/230 = 2.0-2.2	>1 kb	purified PCR products Primer concentration 10μM
Sanger Sequencing						
Xpert Service	-	-	-	-	-	Sample requirements as for Full Service
Pre-mixed Service						
Plasmid	300 – 600 ng	-	-	-	<10 kb	5 pmol Primer/reaction 7,5 µL total volume
	>700 ng	-	-	-	>10 kb	5 pmol Primer/reaction 7,5 μL total volume
PCR-products	<100 ng	-	-	-	100 bp − 1 kb	5 pmol Primer/reaction 7,5 µL total volume
	>100 ng	-	-	-	>1 kb	5 pmol Primer/reaction 7,5 μL total volume
Ready-to-load Service						
Plasmid, PCR-products	-	20	-	-	-	Purified sequencing reaction
Fragment Analysis						
Ready-to-rur	7 -	20	-	-	<600 bp	Fully prepared samples dissolved in formamide
Pre-pared	d -	10	-	-	<600 bp	samples dissolved in formamide without length standard

	Application Amo	nount	Volume (μL)	Conc. (ng/μL)	Quality	Size (bp)	Comment
STR analysis	50 r	ng	-	5	260/280 = 1.8 260/230 = 2.0-2.2	-	Buffer: 10 mM Tris/HCl, low TE or water