Shipping Options



Online Ordering



Sequencing Primers



Genomics

Use our **eco-friendly paper boxes** for secure shipping of your sample plates.



Drop it in your nearest **DropBox** for **free pick-up**.



Alternatively use our Sample Bag (padded envelope), which you may have ordered online already.



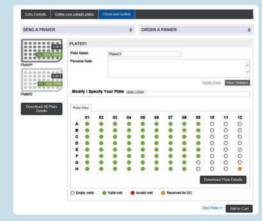
If you have a postal service in your institute or company, address stickers with our sequencing lab adresses are available online, free of charge.



Go to the **PlateSeg Supreme** order page on eurofinsgenomics.eu and select the entry format.



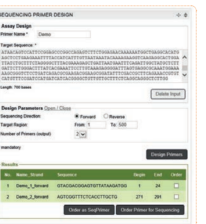
Specify your samples and reaction conditions on the second step. Based on the sample type, the respective PlateSeg Kit and PlateSeg Coupon codes can be selected per plate.



On the last step you have the possibility to modify and finally check your plates. Confirm your sample plates by adding them to your cart.

OPTIMUM PRIMER CONDITIONS

- The optimum primer length is between 16-25 bases
- Primer melting temperature (Tm) should be 50-62 °C
- The GC content of the primer should be 35-60 %
- Ideally one G or C should be located at the 3' primer end
- The number of 3' Gs or Cs should not exceed 2 Gs or Cs
- If possible, avoid >3 identical bases in a row in the sequence
- Primers must not contain phosphorylation or fluorescent dyes



Use our free **Sequencing Primer Design Tool** to design the optimum sequencing primers.

PRIMER CONCENTRATION & VOLUME

- Exactly 10 pmol/μl primer concentration is required per sequencing reaction
- Each primer must have a total volume of 20 μl (double distilled water or 5 mM Tris-HCI)
- 2 µl of primer volume is required for every additional sequencing reaction
- Concentration of **primers with wobble bases** must be calculated according to the following formula: n^x x Conc_{Primer}

n = number of bases within a wobble according to IUPC code; **X** = number of wobbles within the primer sequence. E.g. 1 V (AGC) = 3^1 x 10 pmol/µl; 2 V (AGC) (AGC) = 3^2 x 10 pmol/µl







Premixed samples



- Use our purple PlateSeg Kit Mix or black PlateSeg Kit Mix – NightXpress for your premixed samples
- Alternatively you can use the PCR Plate from our sequencing accessories
- Templates should consist of **15 µl purified DNA** with either of the concentrations given in below table
- DNA concentration must be **normalised** across the plate
- Add 2 µl of primer with a concentration of 10 pmol/µl
- The total volume of your premixed sample must be 17 μl
- Well H12 should be kept free for internal quality control
- Seal your plates using 8-cap strips to prevent material loss
- If you are using your own please use our PlateSeg Labels to label your plate on the plate frame
- Submit samples at **ambient temperature** to our lab

Purified DNA samples



- Use either our blue **PlateSeg Kit DNA** or the PCR Plate from our sequencing accessories
- Plates may contain plasmid DNA and purified PCR products
- Sample concentration must be **normalised** across the plate
- Quantify the DNA concentration via agarose gel or a photometer
- Well H12 should be kept free for internal quality control.
- Seal your plates using 8-cap strips to prevent material loss
- If you are using your own plates, please use our PlateSeq **Labels** to label your plate on the plate frame
- Samples should be sent at ambient temperature

- Concentration must be **normalised** across the plate
- PCR **product size should not vary** by more than a factor of 3
- Well H12 should be kept free for internal quality control
- PCR products should be sent liquid in a total volume of 15 µl
- Seal your plates using 8-cap strips to prevent material loss
- If you are using your own plates, please use our PlateSeg Labels to label your plate on the plate frame
- Ship samples at ambient temperature to us

Sample concentration & volume

Sample type Plasmid DNA	Product length	Sample conc.	Sample vol.
Purified	150-300 bp	1 ng/μl	15 µl
PCR	300-1000 bp	5 ng/μl	15 µl
Products	1000-3000 bp	10 ng/μl	15 µl

Quantify your template concentration via agarose gel or a photometer to ensure accurate results.

Sample concentration & volume

Plasmid DNA	-	50-100 ng/μl	15 µІ
Purified PCR Products	150-300 bp 300-1000 bp 1000-3000 bp	1 ng/μl 5 ng/μl 10 ng/μl	15 µl 15 µl 15 µl

Sample concentration & volume

Unpurified	150-300 bp	4 ng/μl	15 μl
PCR	300-1000 bp	10 ng/μl	15 μl
Products	1000-3000 bp	20 ng/μl	15 μl

Plasmid clones as stab culture



Unpurified PCR products

• Use either our **PlateSeg Kit Clone** or Agar Plate from

our sequencing accessories with appropriate antibiotic

• Use sterile toothpicks to **pick single colonies** from your petri dish and inoculate a single well with one colony

• Incubate plate at **37 °C for 8-12 hours** (overnight)

Labels to label your plate on the plate frame

• Seal the plate with an adhesive plastic foil

PLASMID CLONES AS GLYCEROL CULTURE

• Fill each well with 200 µl of liquid medium

• Use transparent 96well plates with 350 µl/well

• Cover the plate with a lid and loosely wrap with cellophane

• If you are using your own plates, please use our **PlateSeg**

• Ship your stab cultures at **ambient temperature** to us

- Use either our green **PlateSeg Kit PCR** or the PCR Plate from our sequencing accessories
- Quantify the concentration via agarose gel or a photometer

- Include the appropriate antibiotic and add 40 µl glycerol (final glycerol concentration: 10-20 %)
- Use sterile toothpicks to pick single colonies from your petri dish and inoculate a single well with one colony
- Alternatively transfer already arrayed clones from a storage glycerol plate to a freshly prepared 96well plate
- Cover the plate loosely and incubate at 37 °C overnight
- Verify that the **plate surface is dry** before you seal the plate tightly with an adhesive plastic foil
- Use our **PlateSeq Labels** to label your plate
- Freeze the plate at -80 °C
- Ship your glycerol cultures on **sufficient dry ice** to us