

# X-tremeGENE HP DNA Transfection Reagent

For transient and stable transfection of eukaryotic cells

 Cat. No. 06 365 752 001
 Trial-pack

 Cat. No. 06 366 244 001
 0.4 ml

 Cat. No. 06 366 236 001
 1 ml

 Cat. No. 06 366 546 001
 5 × 1 ml

Version 1.0
Content version: October 2010

Store at -15 to -25°C

#### 1. What this Product Does

#### **Number of Tests**

Using the standard procedure, 1 ml of X-tremeGENE HP DNA Transfection Reagent can be used to perform up to 10,000 transfections in 96-well plates.

#### **Formulation**

X-tremeGENE HP DNA Transfection Reagent is a proprietary blend of lipids and other components supplied in 80% ethanol, filtered through 0.2  $\mu$ M pore size membrane, and packaged in glass vials. It does not contain any ingredients of human or animal origin.

#### Storage and Stability

Store X-tremeGENE HP DNA Transfection Reagent at -15 to  $-25^{\circ}$ C, with the lid tightly closed. The reagent is stable until the expiration date printed on the label when stored under these conditions.

X-tremeGENE HP DNA Transfection Reagent remains fully functional even after repeated opening of the vial (at least five times over a two-month period), as long as the vial is tightly recapped and stored at -15 to -25°C.

#### **Special Handling**

- Always bring the vial to +15 to +25°C and mix X-tremeGENE HP DNA Transfection Reagent prior to removing the amount required vortexing for one second.
- Do not aliquot X-tremeGENE HP DNA Transfection Reagent; store in the original glass vials.
- Minimize the contact of undiluted X-tremeGENE HP DNA Transfection Reagent with plastic surfaces.
- For use, the minimum amount of X-tremeGENE HP DNA Transfection Reagent: DNA complex is 100 μl. Complex formation at lower volumes can significantly decrease transfection efficiency.
- ⚠ Do not use tubes or microplates made of polystyrene for X-tremeGENE HP Transfection Reagent: DNA complex preparation. When not able to avoid polystyrene materials, make certain to pipet the transfection reagent directly into the serum-free medium (e.g., Opti-Mem).
- ⚠ Do not use siliconized pipette tips or tubes.

#### **Additional Equipment and Reagents Required**

Additional reagents and equipment required to perform transfection assays using X-tremeGENE HP DNA Transfection Reagent include:

# Standard Laboratory Equipment.

- Standard cell culture equipment (e.g., biohazard hoods, incubators)
- Standard pipettes and micropipettes
- Vortex mixer

#### For Plasmid Preparation

- Purified plasmid stock (0.1 µg/µl 2.0 µg/µl) in sterile TE (10 mM Tris, 1mM EDTA, pH 8.0) buffer or sterile water
- Genopure Plasmid Midi Kit\* or Genopure Plasmid Maxi Kit\* to prepare plasmid

#### For Verification of Vector Function

- Assay appropriate for transfected gene
- G-418 Solution\* or Hygromycin B\* (optional for stable transfection experiments)

#### For Transfection-Complex Formation

- Opti-MEM I Reduced Serum Medium or serum-free medium
- Sterile polypropylene tubes or round-bottom 96-well plates

#### Growing Cells

- Select subconfluent cultures in log phase for preparation of cell cultures
- Quantify cell number to reproducibly plate the same number of cells

#### **Application**

X-tremeGENE HP DNA Transfection Reagent is a high performance transfection reagent, free of animal-derived components. Benefits of X-tremeGENE HP DNA Transfection Reagent include:

- Designed to transfect a broad range of eukaryotic cells, including insect cells, many cell lines not transfected well by other reagents, and hard-to-transfect cell lines (e.g., HT-1080, K-562, HepG2).
- Can be successfully used in a variety of applications, such as gene
  expression analysis and protein production using transiently transfected cells, generation of stable cell lines, expression of shRNA for
  gene knockdown studies, drug discovery programs, and target
  evaluation. Samples and detailed transfection protocols are available at <a href="http://www.powerful-transfection.com">http://www.powerful-transfection.com</a>.
- Produces minimal cytotoxicity or changes in morphology when adequate numbers of cells are transfected, eliminating the requirement to change media after adding the transfection complex.
- · Suitable for transient and stable transfection.
- Functions very well in the presence or absence of serum.

#### 2. How to Use this Product

# 2.1 Before You Begin

# Required Amount of X-tremeGENE HP DNA Transfection Reagent

To optimize, first transfect a monolayer of cells that is 70 - 90% confluent, using 1:1, 2:1, 3:1 and 4:1 ratios of microliter ( $\mu$ I) X-tremeGENE HP DNA Transfection Reagent to microgram ( $\mu$ g) DNA. A ratio of 3:1 of microliter ( $\mu$ I) X-tremeGENE HP DNA Transfection Reagent to microgram ( $\mu$ g) DNA has been shown to be optimal for many cell types.

Q Lower confluencies have also been tested successfully.

The recommended starting concentration is a 3:1. For most cell types, these X-tremeGENE HP DNA Transfection Reagent to DNA ratios provide excellent transfection efficiency.

Surther optimization may increase transfection efficiency in your particular application. In addition to varying the ratio, other parameters may also be evaluated, such as the amount of transfection complex added. For additional optimization guidelines, see Section 3, Troubleshooting and visit <a href="http://www.powerful-transfection.com">http://www.powerful-transfection.com</a>.

#### **Plasmid DNA**

- For best results, accurately determine the plasmid DNA concentration using 260-nm absorption; estimates of DNA by measuring gel band density are not recommended. Determine DNA purity using a 260 nm/280 nm ratio (the optimal ratio is 1.8).
- Prepare the plasmid DNA solution using sterile TE (Tris/EDTA) buffer or sterile water at a concentration of 0.1 to 2.0 μg/μl.
- Use high quality DNA preparation kits to obtain endotoxin-free DNA.

#### **Cell Culture Conditions**

- Minimize intra- and inter-experimental variance in transfection efficiency using cells that are regularly passaged, proliferating well in a log-growth phase, and plated at a consistent density.
- For best results, accurately quantify cell concentration using a hematocytometer or automated system.
- · Cells must be healthy and free of Mycoplasma.
- Cells should have a low passage number to achieve best results.

#### **Other Media Additives**

In some cell types, antimicrobial agents (e.g., antibiotics and fungicides) commonly included in cell-culture media may adversely affect the transfection efficiency of X-tremeGENE HP DNA Transfection Reagent. If possible, exclude additives in initial experiments. Once high-efficiency conditions have been established, these components can be added back while monitoring transfection results. Cell growth and/or transfection efficiency may be affected by variations in serum quality and medium formulations.

#### **Verification of Vector Function**

Optimize transfection conditions using a known positive-control reporter gene construct before transfecting cells with a new vector construct:

- Determine transfection efficiency using a reporter gene assay, such as β-Gal\*, Luciferase\*, or SEAP\*.
- Sequence flanking vector insert regions to verify the integrity of your new construct.

#### 2.2 Transfection Procedure

**Adherent Cells**: Plate cells approximately 24 hours before transfection making sure cells are at the optimal concentration in the appropriate cell culture vessel.

Suspension Cells: Plate freshly passaged cells at optimal concentra-

#### 2.3 Transfection Procedure

- Allow X-tremeGENE HP DNA Transfection Reagent, DNA and diluent to equilibrate to +15 to +25°C. Briefly vortex the X-tremeGENE HP DNA Transfection Reagent vial.
- 2 Dilute DNA with appropriate diluent (*e.g.*, serum-free medium) to a final concentration of 1 μg plasmid DNA /100 μl medium (0.01 μg/μl). Mix gently.
- 3 Place 100 μl of diluent, containing 1 μg DNA into each of four sterile tubes labeled 1:1, 2:1, 3:1, and 4:1.
  - Δ Use a minimum of 100 µl of diluent. Lower volumes may significantly decrease transfection efficiency.
  - Use sterile tubes or tissue culture treated round-bottom, 96-well plates to produce the complex.
- Pipet the X-tremeGENE HP DNA Transfection Reagent (1, 2, 3, or 4  $\mu$ I) directly into the medium containing the diluted DNA without coming into contact with the walls of the plastic tubes. Mix gently.
  - To avoid adversely affecting transfection efficiency, do not allow undiluted X-tremeGENE HP DNA Transfection Reagent to come into contact with plastic surfaces. Do not use siliconized pipette tips or tubes.
- Incubate the transfection reagent:DNA complex for 15 minutes at +15 to +25°C.
  - Some ratios and cell types may required longer incubation (up to 30 min). Determine this for your particular cell line and the ratio used.
- Remove the culture vessel from the incubator. Removal of growth medium is not necessary. Add the transfection complex to the cells in a dropwise manner.
  - See Table 1 to determine component amounts corresponding to the surface area of the cell culture vessel used

Gently shake or swirl the wells or flasks to ensure even distribution over the entire plate surface. If available, use a rotating platform shaker for 30 seconds at low speed for mixing 96-well plates.

Once the transfection reagent: DNA complex has been added to the cells, there is no need to replace with fresh medium (as may be necessary with other transfection reagents)

Following transfection, incubate cells for 18 – 72 hours before measuring protein expression. The duration of incubation will depend on many factors, including the transfected vector construct, the cell type being transfected, the cell medium, cell density, and the type of protein being expressed. After the incubation period, measure protein expression using an assay appropriate for your system.

#### **Notes:**

- As with any experiment, include appropriate controls. Prepare culture wells with cells that remain untransfected, cells with transfection reagent alone, and cells with DNA alone.
- Solution or Hygromycin B).
  For stable transfection experiments, the complex-containing medium should be left unchanged until the cells are passaged. At that time, include appropriate selection antibiotics (e.g., G 418 Solution or Hygromycin B).
- To prepare transfection complexes for different-sized containers or parallel experiments, adjust component amounts corresponding to the surface area of the cell culture vessel used (see Table 1)
- Solution For ease-of-use when transfecting small volumes into 96-well plates containing 0.1 ml culture medium per well, prepare 100 μl of transfection complex, and then add 10 μl to each well (depending on cell type)
- The optimal ratio of transfection reagent to DNA, and the optimal total amount of complex, will depend on the cell line, cell density, day of assay, and gene expressed
- After performing the optimization experiment in which several different ratios are tested, select a ratio in the middle of the plateau optimum for future experiments

**Tab. 1:** Guidelines for Preparing X-tremeGENE HP DNA Transfection Reagent: DNA Complex for Various Culture Vessel Sizes

Culture vessel	Surface Area (cm²)	Total volume of medium (ml)	Suggested amount of 100 µl transfection complex to add to each well (µl)	DNA (µg) using 1:1 or 4:1 Ratio	Final amount of X-tremeGENE HP DNA Transfection Reagent (µl) using 1:1 Ratio	Final amount of X-tremeGENE HP DNA Transfection Reagent (µl) using 4:1 Ratio
96-well plate (1 well)	0.3	0.1	10	0.1	0.1	0.4
48-well plate (1 well)	1.0	0.3	30	0.3	0.3	1.2
24-well plate (1 well)	1.9	0.5	50	0.5	0.5	2
12-well plate (1 well)	3.8	1	100	1	1	4
35-mm dish	8	2	200	2	2	8
6-well plate (1 well)	9.4	2	200	2	2	8
60-mm dish	21	5	500	5	5	20
10-cm dish	55	10	1000	10	10	40
T-25 flask	25	6	600	6	6	24
T-75 flask	75	20	2000	20	20	80

# 3. Troubleshooting

Observation	Possible Cause	Recommendation
Low Transfection Efficiency	Suboptimal X-tremeGENE HP DNA Transfection Reagent : DNA ratio	Titrate optimal X-tremeGENE HP DNA Transfection Reagent : DNA ratio. Refer to the text in Section 2.1 "Before you begin".
	Insufficient number of cells	Determine optimal cell density for each cell type. For most cell types, 70 – 90% confluence at transfection is optimal.
	X-tremeGENE HP DNA Transfection Reagent : DNA complexes did not form well	Prepare complexes in serum-free medium (e.g., Opti-MEM). Do not use siliconized pipet tips or tubes. Do not aliquot the X-tremeGENE HP DNA Transfection Reagent.
	Incubation time of transfection	Determine the optimal incubation time (18 - 72 h). Optimal for most cell types and plasmids is 24 - 48h.
	Inhibition by media components	Some media components (e.g., polyanions) may influence the transfection.
	Low volume of X-tremeGENE HP DNA Transfection Reagent : DNA complex	The minimum amount of X-tremeGENE HP DNA Transfection Reagent to DNA complex is 100µl. Complex formation at lower volumes may significantly decrease the transfection efficiency; refer to the text in Section 1, "Special Handling".
High Cytotoxicity	Cells are cultured in serum-free medium	Transfection using X-tremeGENE HP DNA Transfection Reagent in cells cultured in serum-free medium is possible, however, toxicity may be higher when serum is absent.
	X-tremeGENE HP DNA Transfection Reagent : DNA complexes and cells not mixed well	Add X-tremeGENE HP DNA Transfection Reagent dropwise to the cells. Gently rock the dish/plate back and forth and from side to side to evenly distribute the complexes.
	Plasmid preparation contaminated with endotoxin	Use highly purified, sterile, contaminant-free DNA for transfection.
	Transfected protein is cytotoxic or is produced at high levels	Reduced viability or slow growth rates may be due to high levels of protein expression, with cellular metabolism directed toward production of the heterologous protein. Note that the expressed protein may also be cytotoxic at the expressed levels.
	Too much transfection complex for number of cells	Increase the number of plated cells, and/or decrease the total amount of complex added to the cells.

#### 4. Additional Information on This Product

#### **Quality Control**

Each lot of X-tremeGENE HP DNA Transfection Reagent is tested using established quality control procedures.

#### **Functional Analysis**

Cells are transfected with a reporter gene vector DNA using X-tremeGENE HP DNA Transfection Reagent (ratio 3:1  $\mu l/\mu g$  DNA). Reporter gene activity is monitored by chemiluminescent detection. Using a standard curve analysis method, total amounts of recombinant protein per well are measured to ensure levels that are within specification.

#### 5. Supplementary Information

#### Conventions

In this document, the following symbols are used to highlight important information:

Symbol	Description
<b>©</b>	Information Note: Additional information about the current topic or procedure.
	Important Note: Information critical to the success of the procedure or use of the product.

#### **Text Conventions**

To make information consistent and understandable, the following text conventions are used in this Instruction Manual:

Text Convention	Use
Numbered instructions labeled <b>1</b> , <b>2</b> etc.	Steps in a procedure that must be performed in the order listed.
Asterisk *	Denotes a product available from Roche Applied Science.

## **Changes to Previous Version**

First edition

#### **Ordering Information**

Roche Applied Science provides a large selection of reagents and systems for life science research. For a complete overview of related products and manuals, visit and bookmark our home page, www.roche-applied-science.com, and our Special Interest Site on transfection, www.powerful-transfection.com

Product	Pack Size	Cat. No.		
Apoptosis and Cell Death Products				
Cell Proliferation Reagent WST-1	25 ml (2,500 tests) 8 ml (800 tests)	11 644 807 001 05 015 944 001		
Cytotoxicity Detection Kit <sup>PLUS</sup> (LDH)	1 kit 400 tests in 96 wells 1 kit 2,000 tests in 96 wells	04 744 926 001		
		04 744 934 001		
Gene Knockdown Rea	gent			
X-tremeGENE siRNA Transfection Reagent	1 ml (400 transfections in a 24-well plate)	04 476 093 001		
Mycoplasma Detectio	n Reagents			
Mycoplasma Detection Kit	1 kit (25 tests)	11 296 744 001		
Mycoplasma PCR ELISA	1 kit (96 reactions)	11 663 925 910		
Plasmid Isolation Prod	lucts			
Genopure Plasmid Midi Kit	1 kit (for up to 20 preparations)	03 143 414 001		

Product   Pack Size   Cat. No.			
Protease Inhibitor Tablets and Lysis Reagents			
Complete   20 tablets in glass vials   3 x 20 tablets in glass vials   20 tablets in glass vials   20 tablets in glass vials   20 tablets in EASVpacks   20 tablets in EASV	Kit	tions)	03 143 422 001
3 x 20 tablets in glass vials   11 836 145 001   04 693 116 001   04 693 116 001   04 693 116 001   04 693 116 001   05 056 489 001   3 x 20 tablets in glass vials   11 873 580 001   3 x 20 tablets in glass vials   12 055 056 489 001   05 05	Protease Inhibitor Tab		
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Mammalian cell lysis   And 20 complete Protease Inhibitor Cocktail Tablets	cOmplete, EDTA-free	3 x 20 tablets in glass vials	05 056 489 001
Lysis-M,EDTA-free (for mammalian cell lysis)  Reporter Gene Assays  CAT ELISA 1 kit (192 tests) 11 363 727 001  β-Gal Reporter Gene Assay, chemilluminescent before Core format, 250 assays, tube format, 250 assays, 11 669 893 001  Luciferase Reporter Gene Assay, high sensitivity  SEAP Reporter Gene Assay, high sensitivity  SEAP Reporter Gene Pate format, 250 assays, microplate format, 07 250 assays, tube format)  Selection Antibiotics  G-418 Solution 20 ml 04 727 878 001 04 727 878 001 100 ml 04 727 894 001  Hygromycin B 1 g (20 ml) 10 843 555 001  Transfection Reagents  X-tremeGENE 9 DNA Transfection Reagent 1 ml 06 365 779 001 5 x 1 ml 06 365 809 001 Trial-pack 06 366 511 001  Western Blotting Reagents  Lumi-Light <sup>PLUS</sup> Western Blotting Kit (1,000 cm² membrane)  Western Blotting Reagents  Lumi-Light <sup>PLUS</sup> Western Blotting Membranes 100 ml (1,000 cm² membrane)  Western Blocking Reagent, Solution 100 ml 1 1 921 673 001 (10 blots, 100 cm²)  6 × 100 ml (10 blots, 100 cm²)  6 × 100 ml (60 blots, 100 cm²)  6 × 100 ml (60 blots, 100 cm²)  RTCA Analyzer 05 228 972 001  RTCA SP Station 05 232 368 001  RTCA MP Station 05 232 368 001  6 × 6 Units 05 232 368 001  E-Plate VIEW 96 6 Units 06 472 451 001		and 20 complete Protease	04 719 956 001
CAT ELISA 1 kit (192 tests) 11 363 727 001 β-Gal Reporter Gene Assay, chemiluminescent 1 kit (500 assays, microplate format, 250 assays, tube format) β-Gal ELISA 1 kit (192 tests) 11 539 426 001 hGH ELISA 1 kit (192 tests) 11 585 878 001 Luciferase Reporter Gene Assay, high sensitivity  SEAP Reporter Gene Assay, high sensitivity  SEAP Reporter Gene Assay, chemiluminescent 1 kit (500 assays, microplate format, or 250 assays, chemiluminescent 2 00 ml 100 ml 104 727 878 001 Hygromycin B 1 g (20 ml) 10 843 555 001  Transfection Reagents  X-tremeGENE 9 DNA Transfection Reagent 1 ml 100 ml 106 365 787 001 5 x 1 ml 106 365 787 001 5 x 1 ml 106 365 809 001 Trial-pack 06 366 511 001  Western Blotting Reagents  Lumi-Light <sup>PLUS</sup> Western 100 ml 100 ml 100 ml Blotting Kit (1,000 cm² membrane) Western Blocking Reagents 1 roll (1,000 cm² membrane)  Western Blocking Reagent, Solution 1 100 ml (10 blots, 100 cm²)  Western Blocking Reagent, Solution 1 100 ml (10 blots, 100 cm²)  Cellular Analysis  RTCA Analyzer 05 228 972 001  RTCA SP Station 05 331 625 001  RTCA Control Unit 1.1  E-Plate 96 6 Units 05 232 368 001 E-Plate VIEW 96 6 Units 06 472 451 001	Lysis-M,EDTA-free (for	and 20 complete, EDTA-free Protease Inhibi-	
β-Gal Reporter Gene Assay, chemiluminescent         1 kit (500 assays, microplate format, 250 assays, tube format)         11 758 241 001           β-Gal ELISA         1 kit (192 tests)         11 539 426 001           hGH ELISA         1 kit (192 tests)         11 585 878 001           Luciferase Reporter Gene Assay, high sensitivity         200 assays         11 669 893 001           SEAP Reporter Gene Assay, high sensitivity         1 kit (500 assays, microplate format, or 250 assays, or 250 assa	Reporter Gene Assays		
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Gene Assay, high sensitivity       1,000 assays       11 814 036 001         SEAP Reporter Gene Assay, chemiluminescent       1 kit (500 assays, microplate format, or 250 assays, tube format)       11 779 842 001         Selection Antibiotics         G-418 Solution       20 ml 04 727 878 001         Hygromycin B       1 g (20 ml)       10 843 555 001         Transfection Reagents         X-tremeGENE 9 DNA Transfection Reagent       0.4 ml 06 365 779 001       06 365 787 001         Transfection Reagent       1 ml 06 365 809 001       06 366 511 001         Western Blotting Reagents         Lumi-Light <sup>PLUS</sup> Western Blotting Kit (1,000 cm² membrane)       12 015 218 001         Mouse/Rabbit)       1 roll (1,000 cm² membrane)       03 010 040 001         PVDF Western Blotting Membranes (30 cm × 3.00 m)       11 921 673 001         Western Blocking Reagent, Solution (10 blots, 100 cm²)       11 921 673 001         Reagent, Solution (10 blots, 100 cm²)       11 921 681 001         Cellular Analysis         RTCA Analyzer       05 228 972 001         RTCA SP Station       05 229 057 001         RTCA Control Unit 1.1       05 454 417 001         E-Plate 96 6 Units	hGH ELISA	1 kit (192 tests)	11 585 878 001
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Reagent, Solution       (10 blots, 100 cm²)       11 921 681 001         6 × 100 ml (60 blots, 100 cm²)       11 921 681 001         Cellular Analysis         RTCA Analyzer       05 228 972 001         RTCA SP Station       05 229 057 001         RTCA MP Station       05 331 625 001         RTCA Control Unit 1.1       05 454 417 001         E-Plate 96       6 Units       05 232 368 001         6 x 6 Units       05 232 376 001         E-Plate VIEW 96       6 Units       06 472 451 001			03 010 040 001
(60 blots, 100 cm²)       Cellular Analysis       RTCA Analyzer     05 228 972 001       RTCA SP Station     05 229 057 001       RTCA MP Station     05 331 625 001       RTCA Control Unit 1.1     05 454 417 001       E-Plate 96     6 Units     05 232 368 001       6 x 6 Units     05 232 376 001       E-Plate VIEW 96     6 Units     06 472 451 001		(10 blots, 100 cm <sup>2</sup> )	11 921 673 001
RTCA Analyzer       05 228 972 001         RTCA SP Station       05 229 057 001         RTCA MP Station       05 331 625 001         RTCA Control Unit 1.1       05 454 417 001         E-Plate 96       6 Units 05 232 368 001 05 232 376 001         E-Plate VIEW 96       6 Units 06 472 451 001			11 921 681 001
RTCA SP Station 05 229 057 001  RTCA MP Station 05 331 625 001  RTCA Control Unit 1.1 05 454 417 001  E-Plate 96 6 Units 05 232 368 001 6 x 6 Units 05 232 376 001  E-Plate VIEW 96 6 Units 06 472 451 001	Cellular Analysis		
RTCA MP Station       05 331 625 001         RTCA Control Unit 1.1       05 454 417 001         E-Plate 96       6 Units       05 232 368 001         6 x 6 Units       05 232 376 001         E-Plate VIEW 96       6 Units       06 472 451 001			05 228 972 001
RTCA Control Unit 1.1 05 454 417 001  E-Plate 96 6 Units 05 232 368 001 6 x 6 Units 05 232 376 001  E-Plate VIEW 96 6 Units 06 472 451 001	RTCA SP Station		05 229 057 001
E-Plate 96 6 Units 05 232 368 001 6 x 6 Units 05 232 376 001  E-Plate VIEW 96 6 Units 06 472 451 001			05 331 625 001
6 x 6 Units 05 232 376 001  E-Plate VIEW 96 6 Units 06 472 451 001	RTCA Control Unit 1.1		05 454 417 001
	E-Plate 96		
	E-Plate VIEW 96		

Product	Pack Size	Cat. No.
Cellavista Basic Magnification: 4x, 10x Illumination: Brightfield only		05 651 522 001
Cellavista Medium Magnification: 4x, 10x, 20x Illumination: Brightfield and Fluorescence, UV, Blue, Green		05 651 549 001
Cellavista High End Magnification: 2x, 4x, 10x, 20x, 40x Illumination: Brightfield and Fluorescence, UV, Blue, Cyan, Green, Amber, Red		05 651 557 001
Cedex XS Analyzer with Control Unit		05 926 432 001
Cedex Smart Slide package	15 x 8 measurements	05 650 801 001
CASY Model TT 45, 60, 150 µm		05 651 735 001

#### **Regulatory Disclaimer**

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