

Using liquid handling systems in the laboratory

Dispensing very small amounts of liquid is essential in all fields of natural science and medicine.

Pipetting in the microliter range is an integral part of a wide spectrum of experimental systems. New dispensing systems allowed experiments to be automated and simplified. At the same time, new technologies, such as genetics, place even higher demands on the construction and materials of the systems used.

Since the late 1950s, dispensing technology in the lab has been in a constant state of development and technical improvement.

A wide range of pipetting systems is now available, including air-cushion pipettes and positive-displacement pipettes as well as manual and electronic systems. Direct pipetting, "reverse" pipetting and dispensing are all possible. New problems arise all the time in the lab. New methods or legal restrictions lead to even more complex demands being made.

This issue of the "Applications" series has been designed as a lab poster, providing an overview of pipetting in the lab and of frequently-occurring problems.

Examples include:

- Which basic rules should be observed when pipetting?
- Which pipetting technique is recommended for which application?
- How can a pipette be tested and adjusted?
- Which measures should be taken for pipetting aggressive solvents?
- How can a pipette be decontaminated?

Unfortunately, it is not possible to deal with all problems in this issue. For further information, please read the instruction manual which accompanies your Eppendorf pipette.

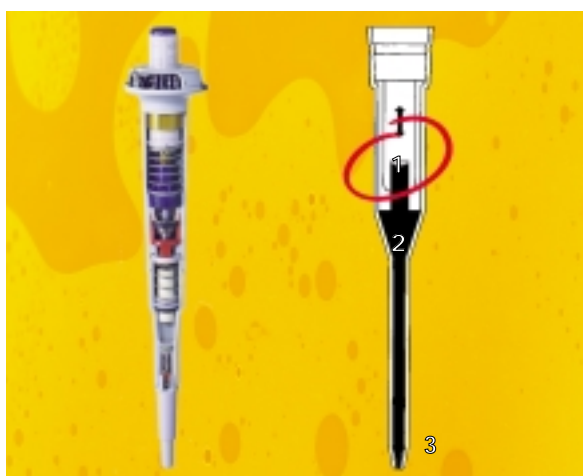


Construction principle of pipetting and dispensing systems

There are two fundamental principles: Air-cushion pipettes (piston-stroke pipettes) and positive-displacement systems.

The fundamental differences are shown in the two diagrams.

Positive-displacement systems function with virtually no air cushion, since an integrated piston in the pipette tip comes into direct contact with the sample solution. The piston is replaced after every pipetting process. Many dispensers also function according to this principle.



Air-cushion pipette
The seals and spring systems are clearly visible.

Positive-displacement system
The piston which is integrated in the tip (2) and which has a leakage seal (3) is reversibly connected to the piston of the pipette (1).



Important applications include pipetting viscous solutions, solutions with a high vapor pressure or high density, tenside solutions, as well as preventing contamination when working with radioactive or aggressive substances and biomolecules, such as nucleic acids in PCR*.

Piston-stroke pipettes have an air-cushion which moves between the piston and the sample liquid, and which aspirates and dispenses the sample. It functions like an elastic spring. A series of other factors must be taken into consideration, e.g. the dead volume, heat from the user's hand and the shape of the pipette tip.

One advantage is that pipetting results are almost totally unaffected by the vapor pressure, density, viscosity or wetting behavior of the solution. Furthermore, cross-contamination as a result of aerosols being carried over is impossible.

This system guarantees high accuracy and precision and low costs for most applications. Filtertips can also be used to prevent contamination from aerosols.

* PCR is patented by Hoffmann-La Roche

Applications

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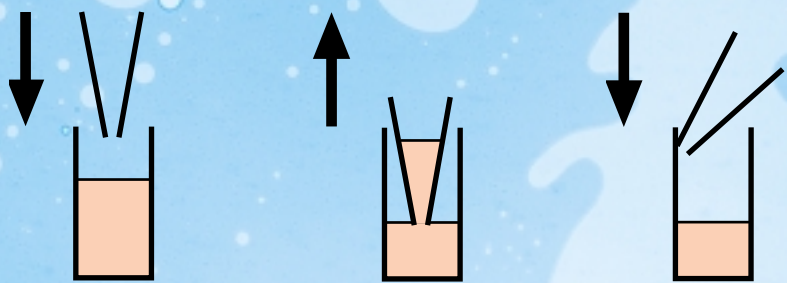
Liquid Handling No. 1

No. 10

1. Pipetting techniques

1.1 Air-cushion pipettes

Forward pipetting

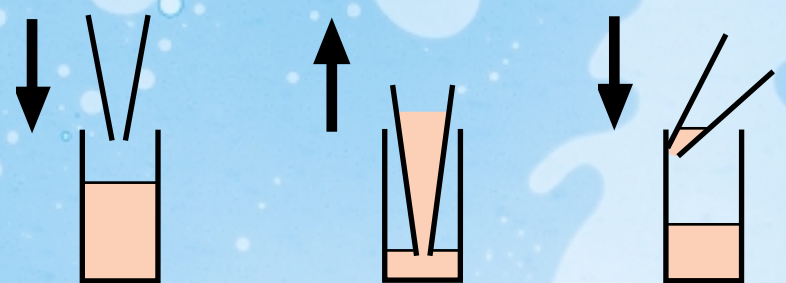


1. Press down button to first stop. Immerse tip a few millimetres into liquid.
2. Release button slowly. Tip fills up.
3. Dispense liquid by pressing down button to first stop. Then blow out remaining liquid by pressing button down to second stop.

Prewet tip when working with volumes greater than 10 μ l

Application is recommended for:
Standard solutions, such as water, buffer, diluted saline solutions and diluted acids and alkalis.

Reverse pipetting



1. Press down button to second stop. Immerse tip a few millimetres into liquid.
2. Release button slowly. Tip fills up.
3. Dispense liquid by pressing down button to first stop. Some liquid remains in the tip!

Application is recommended for:
Viscous solutions, solutions with a high vapor pressure, wetting solvents.

1.2 Positive-displacement systems

When dispensing liquids with high vapor pressure it is preferable to use a direct-displacement system instead of reverse pipetting. For this application it is recommended to prewet the tip of the direct-displacement pipette. Dispensing hexane using the Multipipette® plus / Combitips plus

system can be used as an example of prewetting a positive-displacement system. Hexane is aspirated once or twice into the Combitip plus and immediately dispensed. This allows the small bubble of air remaining in the Combitip plus to be saturated with hexane vapor.

Highly accurate dispensing with exact drip separation is then possible. This is very important for sample preparation in HPLC and other such applications. If the Combitip is not prewetted, the solution gradually evaporates and thus expels the hexane from the Combitip, i.e. the

Combitip drips. When pipetting with an air-cushion pipette, it is virtually impossible to saturate the entire air space above the liquid with vapor, since the volume of air is considerably larger than that in a positive-displacement tip.

2. Troubleshooting and solutions

Handling errors, damaged or contaminated devices and external factors can cause significant deviations to desired pipetting volumes. The cause of, and solutions to, some important errors are listed below.

Error	Cause	Solution
Pipette drips or leaks.	Tip is loose. Tip does not fit correctly.	Use original tip. Press on tightly.
	Nose cone is scratched.	Replace nose cone.
	Seal of nose cone leaks.	Replace nose cone or seal.
	Piston contaminated by reagent deposits.	Clean and lubricate piston. Replace seal.
	Piston damaged.	Replace piston and piston seal.
	Piston seal damaged.	Replace seal and lubricate piston.
Pushbutton does not move smoothly.	Piston scrapes and is contaminated.	Clean and lubricate piston.
	Seal swollen by reagent vapors.	Open pipette and ventilate. Lubricate piston if necessary.
	Piston visibly damaged or coated with insoluble deposit.	Replace piston and piston seal.
Volume inaccurate.	Deviating pipetting conditions.	See table on right.
	Pipette leaks.	Check tightness, then proceed as above.
	Pipette misadjusted.	Recalibrate as described.

Causes of inaccurate volume	Maximum error (inaccuracy)	Solution
Pipette is held at an angle (30°).	+ 0.5 %	Hold pipette straight.
Pipette tip does not fit correctly.	> 0.4 %	Use original Eppendorf tips.
Density of medium (e.g. $\rho = 1.1 \text{ g/cm}^3$).	- 0.4 %	Adjust pipette.
Temperature differences (e.g. pipette 22 °C, sample 4 °C).	- 5.4 %	If possible, align pipette and medium.
Geographical location (e.g. 1,000 m above sea level).	- 0.4 %	Adjust pipette.

3. Checking pipettes for leaks

Testing for leaks is relatively easy with pipettes with a volume greater than 10 μ l. A transparent plastic tube of approximately 20 cm in length is required. After liquid has been aspirated through the tube, a mark is made on the meniscus. After one minute has expired, the level of the liquid should not have fallen below the mark. If this does occur, it should be assumed that there is a leak and the

The tube should have the following inner diameter:

for pipettes	
10–100 μ l	0.5–1.0 mm
100–500 μ l	1.5–2.0 mm
>500 μ l	5.0 mm

pipette should be serviced in accordance with the instruction manual.



4. Chemical stability of plastics used in Eppendorf pipettes

The parts of the pipette which may come into contact with solvent vapors are made of PP or PVDF. The detachable tip ejector of the Reference pipette is made of PEI.

If necessary, the suitability of the pipette should be examined. Cleaning the pipette after aggressive solvents have been used is highly recommended. Due to individual differences in the conditions of use, the accuracy of the details cannot be guaranteed.

Substance	Density [mg/μl]	Vapor pressure [mbar]	Concentration [%]	Temperature [°C]	PP	PVDF	PEI
Acetic acid	1.06	15.4	100	20	+	+	+
Acetone	0.79	233.0	100	20	+	o	+
Ammonia			25	20	+	+	+
Butanol	0.81	6.7	100	20	+	+	+
Chloroform	1.47	213.0	100	20	o	+	-
Ethanol	0.79	59.0	96	20	+	+	+
Formic acid	1.23	42.0	10	20	+	+	+
Glycerine	1.26		100	60	+	+	+
Hydrochloric acid	1.15		30	20	+	+	+
Hydrofluoric acid	1.13		60	20	+	+	+
Methanol	0.79	128.0	100	20	+	+	+
Nitric acid	1.41		70	20	-	+	o
Phenol	1.06		10	20	+	+	+
Phosphoric acid	1.88		85	20	+	+	+
Potassium hydroxide solution	1.29		30	20	+	o	+
2-propanol (iso-)	0.78	42.5	100	20	+	+	+
Sodium hydroxide solution	1.33		30	20	+	o	+
Sulfuric acid	1.84		98	20	o	+	o
Trichloroacetic acid	1.62		50	20	+	+	o

- + : Stable, remains unchanged even after long periods of contact
- o : Partly stable, can only be used for short periods of contact
- : Unstable, changes in the material can occur after short periods of contact

5. Decontaminating air-cushion pipettes when working with various liquids

Liquid	Handling, Special features	Decontamination
Aqueous solutions and buffers	Pipette is calibrated with distilled water. Results are extremely accurate.	Open pipette, rinse contaminated parts well with distilled water, allow to dry at maximum 60 °C in drier compartment. Lubricate piston if necessary.
Inorganic acids	It is advisable to occasionally rinse the pipette lower part with distilled water if high-concentration acids are pipetted frequently. Using Filtertips is also recommended.	The plastics used in Eppendorf pipettes are acid-resistant, as are the ceramic pistons (except to hydrofluoric acid). However, aerosols from the acids can enter the pipette lower part and affect the performance of the pipette. Clean as described above in "Aqueous solutions".
Alkalis	It is advisable to occasionally rinse the pipette lower part with distilled water if high-concentration alkalis are pipetted frequently. Using Filtertips is also recommended.	The plastics used in Eppendorf pipettes are alkali-resistant, as are the ceramic pistons. However, aerosols from the alkalis can enter the pipette lower part and affect the performance of the pipette. Clean as described above in "Aqueous solutions".
Potentially infectious liquids	To avoid contamination, Filtertips should be used. Alternatively, positive-displacement systems can be used.	Autoclave the contaminated parts at 121 °C for 20 min (the Eppendorf Reference can be completely autoclaved. It must be disassembled before hand by unscrewing twice) or immerse the lower parts in normal laboratory disinfectants, rinse with distilled water and allow to dry as described above. Proceed as described above in "Potentially infectious liquids".
Cell cultures Organic solvents	To guarantee sterility, Eppendorf Filtertips should be used. 1. Density is different to that of water. Therefore it is necessary to adjust the pipette. 2. Pipetting should be carried out rapidly, due to the high vapor pressure and the changes in the wetting behaviour. 3. After pipetting has been finished, open the pipette and allow the liquid to evaporate.	This evaporation process is normally sufficient for liquids with a high vapor pressure. Alternatively, immerse the contaminated parts in detergent, rinse well with distilled water and dry as described above.
Radioactive solutions	To avoid contamination, Filtertips should be used. An alternative would be to use positive-displacement systems.	Open pipette and place contaminated parts in complex solutions or special cleaning solutions, rinse well with distilled water and dry as described above
Nucleic acids/Proteins	To avoid contamination, Filtertips should be used. An alternative would be to use positive-displacement systems.	1. Proteins: Open pipette, rinse pipette with detergent. Rinse and dry as described above. 2. Nucleic acids: Decontaminate by boiling in glycine/HCl buffer (pH 2) for 10 minutes (this ensures that no more DNA can be detected on an agarose gel). Rinse well with distilled water and dry as described above. Lightly lubricate piston.

6. Testing the accuracy and precision of pipetting systems

A. Gravimetric testing

The process is described in detail in the Eppendorf "Standard Operating Procedure" (SOP), which is available free-of-charge or can be downloaded (www.eppendorf.com). The following preconditions must be fulfilled for gravimetric testing (see photo):

- Evaporation protection for the weighing chamber, with the aid of moist blotting paper or an evaporation trap.
- No drafts, direct sunlight or heat.
- Distilled or deionized and degassed water must be used as a test-solution.

- The temperature of the room, the water and the pipette must be maintained at a constant 15 – 30 °C, ± 0.5 °C.

The mass of the water is obtained by weighing; this must be converted into the volume (volume = mass / density). Temperature and density are also factors in this conversion.

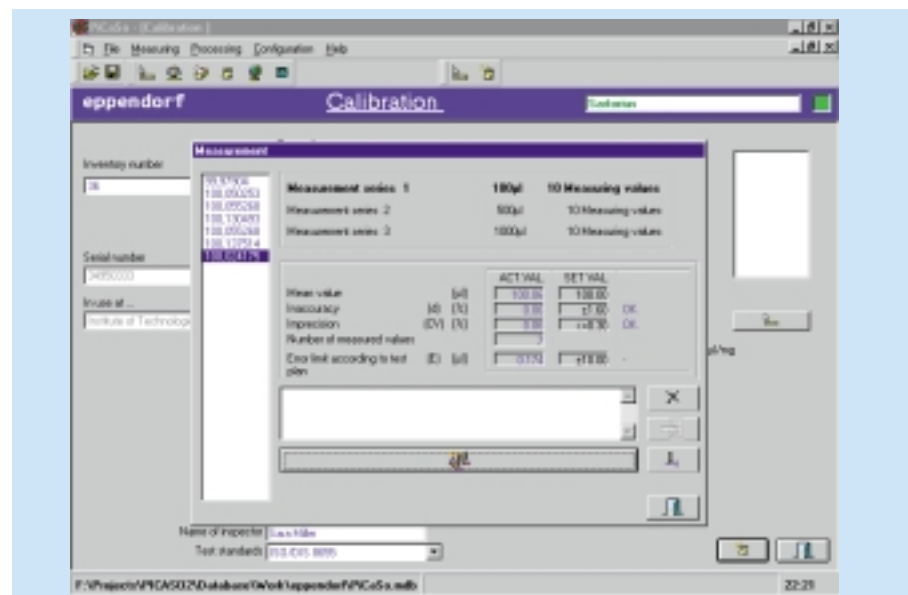
Example: Mass obtained: 99.3 mg.
At 23 °C, the density ρ of water is 0.997 mg/μl. The actual volume is 99.6 μl.
Further details can be found in the Eppendorf SOP.

Volume tested	2 to 50 μl	Greater than 50 μl
Analytical balance with a sensitivity of	0.001 mg	0.01 mg
Weighing vessel max.	6 ml	20 ml
Room thermometer with a resolution of	0.1 °C	0.1 °C
Thermometer for sample with a resolution of	0.1 °C	0.1 °C

B. Photometric pipette test

Testing pipettes with a volume of less than 1 μl is virtually impossible using an analytical balance. A photometric test has








been developed for these pipettes, which is described in detail in the special issue from Eppendorf.



Pipette/Tip combination

Eppendorf offers the ideal pipette tips for applications in all volume ranges. Each tip has been specially developed to suit a specific volume range in order to guarantee the

best possible dispensing results with the appropriate pipettes. In this way, Eppendorf pipettes combine with Eppendorf pipette tips to form a perfect system.

Model	Volume-range	Standartip/Eurotips							GELoader Tip	Filtertip	Biopur
		Nanotip 2.5 µl	Cristaltip 20 µl	100 µl	300 µl	1,000 µl	1.25 ml	2.5 ml			
	0.1 µl–2.5 µl	•								2.5 µl	
	0.5 µl–10 µl	•	•						•	10 µl	20 µl
	2 µl–20 µl			•						100 µl	100 µl
	10 µl–100 µl			•						100 µl	100 µl
	20 µl–200 µl			•	•					250 µl	100/300 µl
	100 µl–1,000 µl					•				1,000 µl	1,000 µl
	500 µl–5,000 µl							•			
	0.1 µl–2.5 µl	•								2.5 µl	
	0.5 µl–10 µl	•	•						•	10 µl	20 µl
	2 µl–20 µl		•						•	10 µl	20 µl
	2 µl–20 µl			•						100 µl	100 µl
	10 µl–100 µl			•						100 µl	100 µl
	50 µl–200 µl			•	•					250 µl	100/300 µl
	100 µl–1,000 µl					•				1,000 µl	1,000 µl
	500 µl–2,500 µl							•			2.5 ml
	1 µl–10 µl	•	•						•	10 µl	20 µl
	10 µl–100 µl			•						100 µl	100 µl
	200 µl–1,000 µl					•				1,000 µl	1,000 µl
	1,500 µl–2,500 µl							•			2.5 ml
	10 µl–100 µl			•						100 µl	
	200 µl–1,000 µl					•				1,000 µl	
	0.5 µl–10 µl	•	•						•	10 µl	20 µl
	10 µl–100 µl			•						100 µl	
	10 µl–100 µl			•						100 µl	100 µl
	30 µl–300 µl				•					250 µl	300 µl
	0.5–10 µl	•	•						•	10 µl	20 µl
	5–100 µl			•						100 µl	100 µl
	20–300 µl				•					250 µl	300 µl
	50–1,000 µl (50–1,200 µl)					•	•			1,000 µl	1,000 µl
	100–5,000 µl							•			

Technical specifications subject to change.

-  Manual
-  Electronic
-  Single-channel
-  Multi-channel
-  Adjustable
-  Fixed
-  Pipetting
-  Dispensing

Ordering information

Manual single-channel pipettes

Eppendorf Reference® 4910

Volume range	Button color	Order number
0.1–2.5 µl	anthracite	4910 000.085
0.5–10 µl	gray	4910 000.018
2.0–20 µl	gray	4910 000.026
2.0–20 µl	yellow	4910 000.034
10–100 µl	yellow	4910 000.042
50–200 µl	yellow	4910 000.093
100–1000 µl	blue	4910 000.069
500–2500 µl	red	4910 000.077

Eppendorf Research® 3111

Volume range	Button color	Order number
0.1–2.5 µl	anthracite	3111 000.017
0.5–10 µl	gray	3111 000.025
2.0–20 µl	yellow	3111 000.033
10–100 µl	yellow	3111 000.041
20–200 µl	yellow	3111 000.050
100–1000 µl	blue	3111 000.068
500–5000 µl	violet	3111 000.076

Eppendorf Reference® 4900 and Eppendorf Research® 3112 are available as fixed-volume pipettes in many different sizes.

Eppendorf Biomaster® 4830, 1–20 µl

4830 000.017

Eppendorf Mastertips® (positive-displacement system)

0030 001.320

Eppendorf Varipette® 4720, 1–10 ml

4720 000.011

Eppendorf Varitip® P (positive-displacement system)

0030 048.130

Eppendorf Varitip® S (dispensing part / Maxitip)

0030 050.533 / 0030 050.568

Manual multi-channel pipettes

Eppendorf Research® multi-channel

Volume range	Channels	Order number
0.5–10 µl	8	3144 000.018
10–100 µl	8	3144 000.034
30–300 µl	8	3144 000.050
0.5–10 µl	12	3144 000.026
10–100 µl	12	3144 000.042
30–300 µl	12	3144 000.069

Eppendorf Research® pro 4860

Volume range	Channels	Order number
0.5–10 µl	1	4860 000.011
5–100 µl	1	4860 000.020
20–300 µl	1	4860 000.038
50–1000 µl	1	4860 000.046
100–5000 µl	1	4860 000.054
0.5–10 µl	8	4860 000.018
5–100 µl	8	4860 000.534
20–300 µl	8	4860 000.550
50–1200 µl	8	4860 000.577
0.5–10 µl	12	4860 000.526
5–100 µl	12	4860 000.542
20–300 µl	12	4860 000.569

Manual hand dispensers

Eppendorf Multipette® plus

4981 000.019

Eppendorf Combitips® plus

(positive-displacement system) assortment 0030 069.285

Eppendorf Combitips® plus are available as standard products, and in

Eppendorf Biopur® quality, each in different sizes.

Bottle-top dispenser

Eppendorf Varispenser® plus

Size	Volume range	Order number
1	0.5–2.5 ml	4961 000.012
2	1.0–5.0 ml	4961 000.020
3	2.0–10.0 ml	4961 000.039
4	5.0–25.0 ml	4961 000.047
5	10.0–50.0 ml	4961 000.055
6	20.0–100.0 ml	4961 000.063

Eppendorf Varispenser®

Size	Volume range	Order number
1	0.5–2.5 ml	4960 000.019
2	1.0–5.0 ml	4960 000.027
3	2.0–10.0 ml	4960 000.035
4	5.0–25.0 ml	4960 000.043
5	10.0–50.0 ml	4960 000.051
6	20.0–100.0 ml	4960 000.060

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