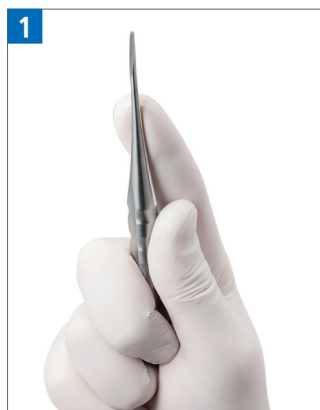




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For the best control, hold the elevator as illustrated.

P-LUX elevators, for axial luxation, are surgical instruments and should be used as such.

Do not use excessive force.



The instrument's tip is inserted into the sulcus towards the root in a most possible axial direction.

The tip is guided deeper along the root through gentle pressure and twisting movement in apical direction.



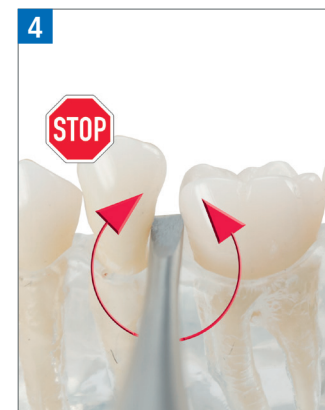
For optimal placement of the instrument, the tips are worked out more delicate.

Attention



Usage with leverage and non-planar pressure needs to be avoided.

This can cause breakage of the instrument tips.



Attention



We cannot be held responsible for damages caused by improper use.

P-LUX[↑]

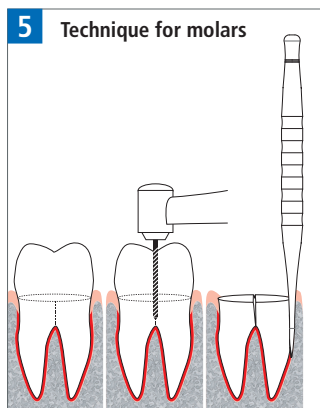
Elevators

REF 617-01 - 617-20

Dear valued customer!

We thank you very much for having chosen a high-quality instrument from A. Schweickhardt GmbH & Co. KG.

In order to enjoy this instrument for a long time, we kindly ask you to follow our application notes.



Note: These P-LUX elevators have been developed as an alternative for classic elevators. The instrument must not be used as a lever.

Furthermore the gentle removal of the teeth should be reached by **axial luxation** and cutting the scharpey's fibres.

Cleaning and Reprocessing

For all cleaning processes, please consider our application guidance of autoclavable instruments and accessories according to DIN EN ISO 17664, in the currently valid version.

See document "0703091007_REPROCESSING_AS_BA_e", downloadable from our website www.schwert.com.

CE

All medical devices indicated here bear the CE mark according to the Medical Device Directive 93/42/EEC.