



Gri-bag®

Specifically designed
for IV compounding

Gri-bag® is a sterile, non-pyrogenic bag for the preparation and administration of intravenous solutions.

Why use Gri-bag®?

Safety

Gri-bag® is needle-free, and therefore offers improved safety for pharmacy technicians by reducing the risk of exposure to hazardous drugs. Gri-bag®, used in conjunction with Gri-fill®, also improves accuracy when mixing, helping ensure that the patient is given the right dose for the treatment.

Sterility

Gri-bag® has a 0.2 µm pore filter [USP*, EP†] incorporated in the filling line that maintains the sterility of the mixture throughout the process. The twist-off dispensing port helps ensure that the mixture has not been altered.

Stability

Gri-bag® has been designed to minimize the risk of biological contamination, and to maintain physical and chemical stability therefore extending the beyond use date [BUD] for each mixture.

*USP: United States Pharmacopeia, † EP: European Pharmacopoeia



Gri-bag® is the perfect complement to Gri-fill®

When a Gri-bag® is filled by a Gri-fill® System, the process includes filtration through a sterilizing 0.2 µm pore filter [USP, EP] and a filter integrity control after filling every unit. The determination of the bubble point allows the system to apply air pressure in order to check that the filter maintains the integrity after every filling. The result of the test is printed on the final label of the compounded unit.

By using Gri-bag® together with Gri-fill®, we can extend the beyond use date [BUD] of the mixture, because we can assure and document that the sterility has been maintained throughout the process.



GRIFOLS

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