

Method Development in Solid Phase Extraction using ISOLUTE® 101 SPE Columns for the Extraction of Aqueous Samples

This Chemistry Data Sheet includes specifics on the use of ISOLUTE® 101 for the extraction of polar analytes from aqueous samples (page 1), ordering information (page 2) and a general discussion on non-polar sorbents (appendix page 3).

The non-selective nature of the resin-based sorbent is ideal for screening applications where a broad range of analytes are to be extracted. The simultaneous extraction of acidic, basic and neutral drugs from biological fluids is a useful application of ISOLUTE 101¹.

In method development using ISOLUTE 101, the following points are important:

Sample Pre-treatment

Due to the extremely hydrophobic nature of ISOLUTE 101, sample pre-treatment is often unnecessary. For viscous samples, dilute with deionized distilled water to reduce the viscosity.

Suppressing ionization of the analytes by pH control can enhance the recovery of polar ionizable molecules using ISOLUTE 101 columns. Using the "2 pH unit rule", samples containing acidic compounds should be adjusted 2 pH units below the lowest pK, while samples containing basic compounds should be adjusted at least 2 pH units above the highest pK. (see **Figure 1**)

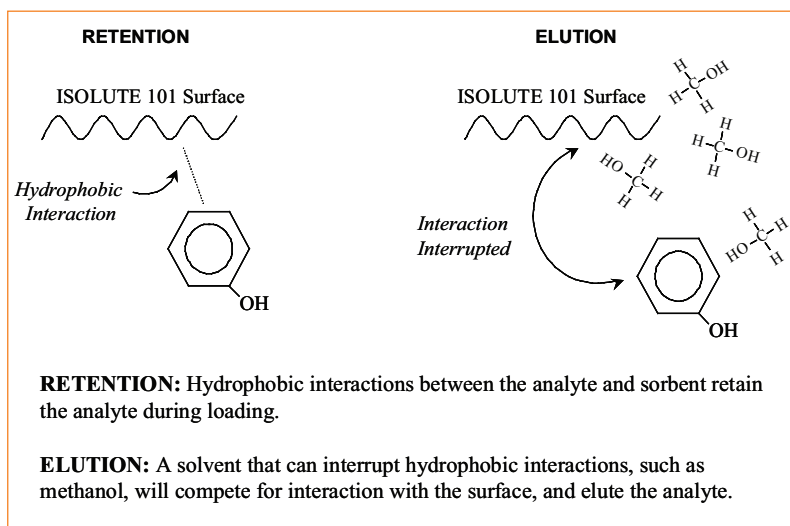


Figure 1. Retention and elution using ISOLUTE 101

Column Solvation and Equilibration

ISOLUTE 101 should be solvated with a water miscible solvent (e.g., methanol, acetone, acetonitrile) prior to sample loading. The solvation step should be followed with a distilled water equilibration rinse to remove excess solvent. If pH control is required, the column should be equilibrated using a buffer of the same pH during the equilibration step.

Sample Loading

When developing a method using ISOLUTE 101, good starting points for flow rates are 3 mL/min for 3 mL columns and 7 mL/min for 6 mL columns. It is likely that loading rates can be increased after method chemistry is established. Evaluation of analyte recovery versus increasing flow rate is a useful exercise to maximize sample throughput. It may be necessary to add 0.5 to 2% (v/v) wetting agent (e.g. methanol, isopropanol) to large volume samples (> 100 mL) to maintain an active sorbent surface.

Interference Elution

A typical solvent for interference elution is distilled deionized water.

If control of sample pH is necessary to maximize retention of the analytes, maintaining the pH of the interference elution solvent at the same pH is often necessary to prevent analyte breakthrough during this step. To improve the purity of the extract, it is sometimes possible to add a water miscible organic solvent such as methanol to the aqueous interference elution solvent without eluting the analyte(s). When optimizing the concentration of organic solvent in the interference elution solvent, it is vitally important to monitor for losses in analyte recovery.

When analytes are non-ionizable, but interference compounds are ionizable, pH control of the interference elution step may be beneficial. To minimize retention of acidic interferences, use a high pH. To minimize retention of basic compounds, use a low pH.

Analyte Elution

This unmodified resin-based sorbent allows analyte elution using a pure organic solvent such as methanol, tetrahydrofuran (THF), isopropanol, acetonitrile, acetone or ethyl acetate.

To minimize the analyte elution solvent volume, allow the elution solvent to "soak" the sorbent bed for a period of time. For users of SPE automation equipment, this soak step can be programmed into the method. Determine analyte recovery versus elution solvent flow rate through the column to maximize recovery. Gravity flow of some elution solvents is sometimes a practical option.

If eluting using a water immiscible solvent, ensure the column is dried before elution. Where pH control is necessary, it is recommended that the addition of a modifier (e.g. formic acid or ammonia) be added to improve the solubility of the analyte(s) in the chosen elution solvent.

References

1. IST1069S SPE Method Recommendations for Toxicological Screening of Post Mortem Samples.

ISOLUTE 101 Product Ordering Information

Part Number	Description	Quantity
101-0010-B	100 mg/3 mL	50
101-0020-B	200 mg/3 mL	50
101-0020-C	200 mg/6 mL	30
101-0050-B	500 mg/3 mL	50
101-0050-C	500 mg/6 mL	30

Appendix

ISOLUTE non-polar sorbents: C2, C2(EC), C4, C6, C8, C8(EC), C18, C18(EC), MFC18, CH(EC), PH, PH(EC), ENV+ and 101.

The ISOLUTE family of non-polar sorbents is used to extract organic compounds from aqueous matrices.

ISOLUTE 101 and ENV+ are the most hydrophobic of all of the sorbents (see **Figure 2** for representation of ISOLUTE 101 surface). They are used primarily where the analytes are very water soluble, and extraction is difficult using a silica based sorbent. The resin-based sorbents are high capacity, highly cross-linked polystyrene based polymer columns capable of retaining analytes of a wide range of polarities. The very accessible high surface area of these non-polar sorbents provide retention of very polar and water soluble analytes. The optimized surface area / pore structure and the absence of fines provide high recoveries at high flow rates for many analytes.

The purity of the polystyrene-based polymers represents a dramatic improvement when compared with the first generation of commercially available styrene divinyl benzene polymer products. The absence of monomers ensures compatibility of the ISOLUTE 101 and ENV+ products with today's demanding low-level analytical applications.

ISOLUTE 101 has an unmodified surface and therefore does not exhibit any secondary interactions. It must be conditioned before use, and analyte elution with unmodified (pure) solvents is possible. ISOLUTE 101 is ideal for the extraction of polar analytes that are not adequately retained on C18 or C8 sorbents.

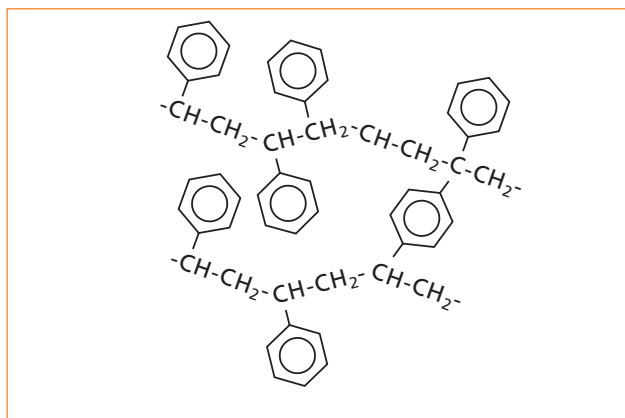


Figure 2. Structure of ISOLUTE 101

ISOLUTE ENV+

The sorbent has been specially derivatized to provide an easily wettable surface. It is used primarily where the analytes are very water soluble, and extraction is difficult using a silica based sorbent. This resin-based sorbent exhibits secondary interactions through the hydroxylated surface, which is particularly useful when extracting basic drugs from biological fluids, since modified elution solvents can be used, thus providing cleaner extracts.

Use of ISOLUTE ENV+ for many applications fields, in particular environmental analyses, is well documented.

ISOLUTE ENV+ is the sorbent of choice for extremely polar compounds.

C18, C18(EC), MFC18

The non-encapped trifunctional C18 sorbent has enhanced secondary silanol interactions (which can be very useful for example in the extraction of basic compounds from aqueous solution) compared to C18(EC). Non-encapped C18 has a lower carbon loading than the encapped sorbent. C18(EC) is also based on trifunctional silane chemistry, with many of the residual silanols on the silica surface end capped to minimize secondary silanol interactions. MF C18 (manufactured using monofunctional octadecyl silane) is non-encapped and like the non-encapped trifunctional C18, provides useful secondary silanol interactions. The accessibility of these silanol groups to analytes and solvents is increased in the monofunctional MF C18, compared to the trifunctional C18 sorbents.

C8, C8(EC), C6, C4, C2, C2(EC), CH(EC)

The non-polar characteristic of these sorbents decreases with carbon chain length. This can be advantageous when extracting non-polar analytes from aqueous matrices. Large, non-polar analytes, although well retained on C18 sorbents, can be difficult to elute as the non-polar interactions between analyte and sorbent are very strong. If a less retentive phase (such as C8, C6, C4, C2) is used, the analytes will still be retained, but can be eluted more easily, in minimal elution volumes. Sorbents which are encapped (C2(EC), C8(EC)) have fewer secondary interactions due to silanol groups than their non-encapped versions, and are therefore not recommended for the extraction of basic compounds.

PH, PH(EC)

These sorbents are generally considered to be less retentive than C18 sorbents, but exhibit different selectivities when extracting aromatic and non-aromatic analytes.



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