VTI-SA+ Vapor Sorption Analyzer is a continuous vapor flow sorption instrument for obtaining precision water and organic vapor isotherms at temperatures ranging from 5 °C to 150 °C at ambient pressure. The VTI-SA+ combines the features of VTI’s original SGA design with almost two decades of field-proven performance: the isothermal aluminum block construction, the three isolated thermal zones and chilled-mirror dew point analyzer for primary humidity measurements with the field-proven TA Instruments thermobalance technology. All to provide precise and accurate gravimetric measurements with excellent temperature and RH stability.

### Maximum Sample Weight
750 mg / 5 g

### Dynamic Range
100 μg / 500 μg

### Weighing Accuracy
±0.1%

### Weighing Precision
±0.01%

### Sensitivity
0.1 μg / 0.5 μg

### Signal Resolution
0.01 μg / 0.05 μg

### Temperature Control
Peltier Elements, Resistance Heaters

### Experimental Temperature Range
5 to 150 °C

### Isothermal Stability
±0.1 °C

### Relative Humidity Control Range
See Figure Below

### Accuracy
±1% RH

### Humidity Control
Closed loop, Dew Point Analyzer

### Organic Solvent Capability
Optional

### Camera/2.5X Microscope Accessory
Optional

### Raman Probe Accessory
Optional

![Graph showing temperature and relative humidity control](image)
VTI-SA+ TECHNOLOGY

Symmetrical Microbalance Design
The VTI-SA+ Analyzer is a symmetrical vapor sorption instrument where both the sample and reference chambers are exposed to the same conditions of
 temperature and humidity. In this symmetrical design, any water or organic vapor sorption onto the hangdown wires and sample holder is differentially eluted and the resultant data represents the uptake by the sample alone. This eliminates the need for background subtraction experiments and associated uncertainty typical in competitive, asymmetrical systems.

Resolution and Stability of the Microbalance
The standard VTI-SA+ boasts a microbalance that has a resolution of 0.1 microgram sensitivity optimized for pharmaceutical applications. A higher mass capacity version (e.g., 5 g, 0.5 microgram sensitivity) is also available. To ensure effective work in pharmaceutical studies, the design provides an enhanced stability by maintaining the balance compartment at a constant temperature, independent of the sample temperature. Because the balance is maintained at constant temperature, the user has the option of drying the sample at temperatures other than the experimental temperature or to run different temperature and RH profiles without removing the sample.

Precision Humidity Measurements
As part of our standard design, the VTI-SA+ employs a chilled mirror dew point analyzer (a NIST-traceable standard for humidity) to determine the absolute relative humidity at the sample. In applications where RH control is critical (as in most pharmaceutical studies), chilled-mirror dew point analyzers are the preferred method, because of the absence of drift and long term stability.

Sorption Testing Using an Organic Vapor
The VTI-SA+ can also be configured for organic vapor sorption. In the VTI-SA+, the concentration of the organic vapor in the gas stream reaching the sample is determined by the fraction of gas going through the organic solvent evaporator and the fraction of dry gas.

In competitive systems, assumptions are made that the evaporator is 100% efficient and that the temperature of the evaporator is constant from low to high concentrations. The VTI-SA+ system measures the temperature of the organic solvent in the evaporator and uses this information together with the Wagner equation to control the organic vapor concentration in the gas phase. This method solves the issue of the adiabatic cooling of the solvent, a major source of error in competitive systems.

The solvent containers/evaporators are easily removed and exchanged so there is no need for decontamination or cleaning of the system when changing organic solvents or reverting to water adsorption experiments. For safety, the evaporator compartment is purged with dry nitrogen and fitted with a combustible gas sensor with an audible alarm, that when triggered, shuts down the power to the analyzer.
Simultaneous Microscope Camera or Raman Measurement
The VTi-SA+ is fully compatible with simultaneous optical measurements, including a high-resolution CCD camera/2.5X microscope or a universal Raman Probe*. These optional accessories are interchangeable and field installable, providing the highest level of flexibility for your measurements.

Sample Chamber Design
In the VTi-SA+ Analyzer, the sample and reference chambers are located within an aluminum block maintained at constant temperature (within ±0.1°C) by precise Peltier control elements. Our unique aluminum block design has two distinct advantages. First, due to the high thermal conductivity of the aluminum, thermal gradients within the chambers are minimized. Secondly, because the chamber is a metal block, the issues of static electricity are eliminated. This feature is especially useful when analyzing fine divided powders, as is often the case with pharmaceuticals. The sample temperature is measured with a highly accurate, calibrated platinum resistance thermometer. When higher temperatures are required, the block can be heated to 150 °C using installed resistance cartridge heaters.

*Raman Spectrometer Required
The patented Q5000 SA delivers the performance and reliability required in a leading sorption analyzer in a compact, user-friendly design. The Q5000 SA is designed for manual or automated sorption analysis of materials under controlled conditions of temperature and relative humidity (RH). Its design integrates our latest high sensitivity, temperature-controlled thermobalance with an innovative humidity generation system, multi-position autosampler, and powerful Advantage™ software with technique specific programs and Platinum™ features.

**SPECIFICATIONS**

- **Temperature Controlled Thermobalance**: Included
- **Dynamic Range**: 100 mg
- **Weighing Accuracy**: +/– 0.1%
- **Weighing Precision**: +/– 0.01%
- **Sensitivity**: < 0.1 μg
- **Baseline Drift**: < 5 μg
- **Signal Resolution**: 0.01 μg
- **Temperature Control**: Peltier Elements
- **Temperature Range**: 5 to 85 ˚C
- **Isothermal Stability**: +/– 0.1 ˚C
- **Relative Humidity Control Range**: 0 to 98 %RH
- **Accuracy**: +/– 1 %RH
- **Autosampler – 10 samples** Included
- **Platinum™ Software**: Included
- **Sample Pans**
  - Metal Coated Quartz 180 μL
  - Platinum 50, 100 μL
  - Aluminum Sealed Pan 20 μL

* Over 24 hours at 25 ˚C and 20 %RH with empty metal coated quartz pans
** Optional tray accommodates 25 samples for use with platinum and sealed aluminum pans
The Q5000 SA is a compact, benchtop instrument that delivers the performance and reliability required in a leading sorption analyzer designed for the study of materials under controlled conditions of temperature and relative humidity. Its modern, user-friendly design features a high sensitivity, temperature-controlled thermobalance, an innovative humidity generation system, a 10-position autosampler, and our latest Advantage™ software with Platinum™ features.

Humidity Control Chamber

The patented design features a pair of mass flow controllers (MFCs) that accurately meter and proportion gas to a symmetrical, well-insulated, aluminum block. The block contains a humidifier, gas transmission and mixing lines, plus easily accessible, identically arranged, sample and reference measurement chambers. Temperature regulation of the block interior from 5 °C to 85 °C is performed by four thermoelectric (Peltier) devices in conjunction with a thermistor in a closed-loop system. The mass flow controllers adjust the amounts of wet (saturated) and dry gas to obtain humidities from 0 to 98 %RH. Identical sensors are located adjacent to the sample and reference crucibles, and provide a continuous indication of humidity. Benefits of the design include precise temperature control and highly consistent atmosphere within the sample and reference chambers.
The heart of the Q5000 SA is our latest high performance thermobalance maintained at a constant temperature (±0.01˚C) by three symmetrically arranged heaters in a well-insulated, gas-purged chamber isolated from the furnace by a water-cooled plate. The sensitive, null-balance design features the latest in precision weighing technology. A key feature of the design for sorption analysis operation is the perfect symmetry of the balance assembly. Customer benefits of the patented design include sensitive, reliable operation with superior baseline flatness and exceptional accuracy and precision in weight change detection, factors that are critical for proper gravimetric sorption analysis performance and are totally free from any vapor condensation or electrostatic forces.

The integral Q5000 SA Autosampler features a programmable multi-position sample carousel that permits automated analysis of up to 10 samples using semi-spherical quartz or metal-coated quartz crucibles, and 25 samples using the optional Q5000 IR tray and platinum or sealed aluminum pans. The design provides smooth and efficient loading and unloading of the sample pans without disturbing the balance. All aspects of sample testing are automated and software controlled including pan taring and loading, sample weighing, autosampler movement, furnace movement, pan unloading, and furnace cooling. Autosampler productivity is maximized by our Advantage™ software which provides pre-programmed analysis, comparison, and presentation of results.

Semi-spherical metal-coated quartz crucibles (180 μL) and optional platinum (50 and 100 μL) TGA pans are available for use with the Q5000 SA. The former are commonly used in sorption analysis because of their anti-static capabilities, chemical inertness and ease of cleaning, while Platinum pans are generic for TGA analysis of most materials. Sealed aluminum pans are also an option for ensuring the integrity of materials which readily adsorb moisture or lose volatiles.
Vapor Sorption analysis is an established technique for determining the effect on materials of exposure to controlled conditions of temperature and humidity. Isotherm and Isohume™ experiments are the most commonly performed analyses.

All TA Instruments sorption analyzers perform a range of essential sorption experiments such as time-courses, isotherms (constant temperature, variable RH), and isohumidity (Isohume™) experiments (constant RH, variable temperature). Complex protocols with step changes in temperature and RH can be defined and saved for later use. Also, multiple experiments can be run sequentially without further operator assistance.

In isotherm experiments, a weighed sample is “dried” externally, or preferably in the instrument, and exposed to a series of humidity step changes at constant temperature. The sample is staged at each humidity level until no further weight change is detected or a set time has elapsed. A data point is recorded; the humidity is changed in 5 or 10% controlled RH steps, and the process repeated in an increasing or decreasing procedure. Isohume experiments involve a series of temperature step changes at constant humidity and result in similar plots. They are used to determine how sample exposure to a given humidity results in a physiochemical change, such as a change in the sample’s hydration state. The curve shape provides useful information to this end.

TA Instruments analysis software offers Sorption Analysis, BET Analysis, and GAB programs. In addition, the full power and flexibility of our renowned Universal Analysis software provides for easy data manipulation, advanced reporting, plotting, and file exporting capabilities. In addition, advanced data reduction of VVTSA+ data can be performed using custom-designed data analysis packages. Analysis options include:

- Kinetic analysis for the determination of rate constant of adsorption
- Isotherm heat of adsorption using the Clausius-Clapeyron equation
- Surface area calculation using the BET equation for either water or organic vapors
**APPLICATIONS**

**Evaluation of Amorphous Structure**
Pharmaceutical scientists are often interested in determining the amount of amorphous material in a drug formulation. As the amorphous and crystalline forms are chemically identical, classical analytical techniques are often insensitive to amorphous content. The figure below shows the moisture sorption analysis of a parent drug in its amorphous and crystalline forms. As the amorphous form absorbs significantly more water, the Q5000 SA can be used to quantify relative amorphous content in drug mixtures.

**Analyzing Small Amounts of Pharmaceuticals**
When evaluating pharmaceuticals it is common for only small amounts of material to be available for conducting multiple analytical tests. Hence, the ability to work with small samples is critical. The low baseline drift of the Q5000 SA means that good results can be obtained on even 10-20 milligrams of a crystalline drug, such as prednisone, which adsorbs <0.1% moisture over a broad humidity range. The sorption results shown below represent about 15 micrograms of weight change full scale. The reversibility (lack of hysteresis) in the sorption/desorption profile for prednisone (as well as the low level of moisture adsorbed) indicates that the moisture picked up by the material is adsorbed on the surface of the material rather than being absorbed into its structure.

**Hydrate Formation**
The figure to the right contains the experimental results demonstrating the formation of a hydrate. The hydrate formation is characterized by a plateau in the desorption branch of the isotherm. In this example the hydrate is formed at around 43.5% RH: The sample adsorbs about 4.5% by weight water and does not lose the water of hydration until the RH is lowered below 25%. This hydrate would be considered as a labile or unstable hydrate.

**Characterization of Morphological Stability**
Exposure to elevated humidity can initiate morphological changes in some pharmaceutical materials, particularly in amorphous sugars. As the humidity is increased, the adsorbed water plasticizes the material and lowers the glass transition. When the glass transition temperature decreases to the experimental temperature, crystallization will typically occur. The data in the figure below shows the behavior of amorphous lactose at 25 °C under a constant increase in humidity. Note how the character in the measured weight signal is indicative of a variety of morphological changes including the glass transition and subsequent crystallization of the amorphous phase.
Packaging Film Analysis

In addition to evaluation of the actual pharmaceutical formulations, sorption analysis can also be valuable in comparing the polymeric films which are being considered for packaging the drugs and other materials. The figure to the right shows comparative profiles for two different packaging materials undergoing temperature and relative humidity cycling. Film A absorbs and desorbs moisture at a more rapid rate than the other film evaluated which suggests it may not be suitable for packaging moisture sensitive compounds.

Rate of Diffusion

The VTI-SA+ can be equipped with a diffusion cell which allows for the direct measurement of the permeability of a film or membrane for a particular solvent vapor. The cell consists of a canister that is filled either with a desiccant or absorber, a graduated lid for attaching the film to be tested, and a wire strip to hang the assembled cell on the hanging wire of the balance. Any vapor permeating through the film gets absorbed immediately and the weight of the cell will increase until steady-state conditions are reached. The normalized rate of permeation is obtained from the slope of this line (weight per unit time) and the diameter of the permeating film.