



nano-TA™: Nano Thermal Analysis

Application Note #5

Heated tip-AFM of nanocomposite Polymer Membranes

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Introduction

Nanocomposite Polymer Membranes: Unlike traditional composite materials whose properties are influenced largely by the constituent components, nanocomposites exhibit such large specific surface areas that interfacial properties can significantly affect bulk behaviour. Poly trimethyl silyl propyne (PTMSP) has garnered attention since its discovery due to its very high permeability, high T_g and exhibited reverse selectivity[1, 2]. However, it was the discovery that the permeability and reverse selectivity could be significantly enhanced by the addition of silica nanoparticles that has generated most recent interest in the material[3-7]. This enhancement was particularly significant because it contradicted the Maxwell equation which predicts a reduction in permeability with the addition of an impermeable phase. Further studies on filler size and surface chemistry suggested that the addition of silica resulted in an increased fractional free volume in the polymer[8]. While the importance of the nanoscale PTMSP-Silica interface has been acknowledged, most of the data currently published is based on macroscopic and microscopic measurements and little is known about the local nanoscale properties of the material. Complex organic systems such as these are particularly well suited for bottom-up, molecular design approaches. In solid state form, these systems are known to exhibit bulk deviating material and transport properties due to finite size limitations and interfacial constraints. Practical examples are found in photonics, light emitting diode materials, nanoelectromechanical systems (NEMS), and organic and inorganic hybrid systems, such as nanocomposites.

A common way to gain fundamental insight into material and transport properties is to perform both a structural and a relaxation analysis. However a big challenge lies in obtaining convenient access to the molecular mobility of material arrangements that are interfacially-constrained. Generally, the molecular mobility is not directly accessible for constrained systems with spectroscopic methods. The availability of the nanoscale thermal probes enables us to directly

access the interface and characterize the polymer via the Heated Tip-AFM mode (a term that we define below).

The aim of this work was to use HT-AFM for the following:

1. To use nano-thermomechanical analysis to locally probe the heterogeneous nanocomposite membrane films.
2. To characterize the interfacial interaction between the dispersed nanoparticles and the polymer matrix by determining the shear forces

Heated Tip AFM (HT-AFM) refers to any AFM operation where a heated tip is used instead of a normal tip. Nearly any AFM mode (tapping/contact/Force-Volume etc.) can accommodate a heated tip to yield new information tied to the thermal properties of the sample. Compared to substrate heating methods, HT-AFM offers some distinct advantages. Foremost is the ability to heat to higher temperatures. With substrate heating, significant warming of the materials in contact with the sample occurs which can include the piezoelectric crystal used for scanning. This can cause problems with the calibration of the scan as well as the possibility of damage to the scanner at high temperatures. Another advantage with the heated tip is that it has much less thermally induced drift. With substrate heating, spatial displacements of many microns are observed over relatively small temperature ranges. With the heated tip, displacements of less than a micron are observed over hundreds of degrees. Other advantages include the ability to operate at very high heating rates due to the low thermal mass, and the ability to perform multiple measurements on specimens that are highly sensitive to thermal history.

nano-TA is a local thermal analysis technique which combines the high spatial resolution imaging capabilities of atomic force microscopy with the ability to obtain understanding of the thermal behaviour of materials with a spatial resolution of sub-100nm. The conventional AFM tip is replaced by a special nano-TA probe that has an embedded miniature heater and is controlled by the specially designed nano-TA hardware and software. HT-AFM enables a surface to be visualised at nanoscale resolution with the AFM's routine imaging modes which enables the user to select the spatial locations at which they would like to investigate the thermal properties of the surface. The user then obtains this information by applying heat locally via the probe tip and measuring the thermomechanical response.

Experimental Setup

temperature at the surface depends on contact resistance, tip temperature, and scan speed. Figure 2 below shows a series of heated tip scans on a PTMSP composite taken at tip temperatures above, but with equilibrium scan temperatures below the T_g of the matrix. The HT-AFM is operated over a region with exposed silica particles, while increasing temperature in 10°C increments with each new scan. When a critical temperature is reached (290°C for the series under investigation), the forces acting on the particle, coupled with potential thermally induced instabilities, are sufficient to debond the particle from the matrix. The lack of matrix deformation around the particle suggests that the debonding occurred below T_g and most likely in an elastic manner.

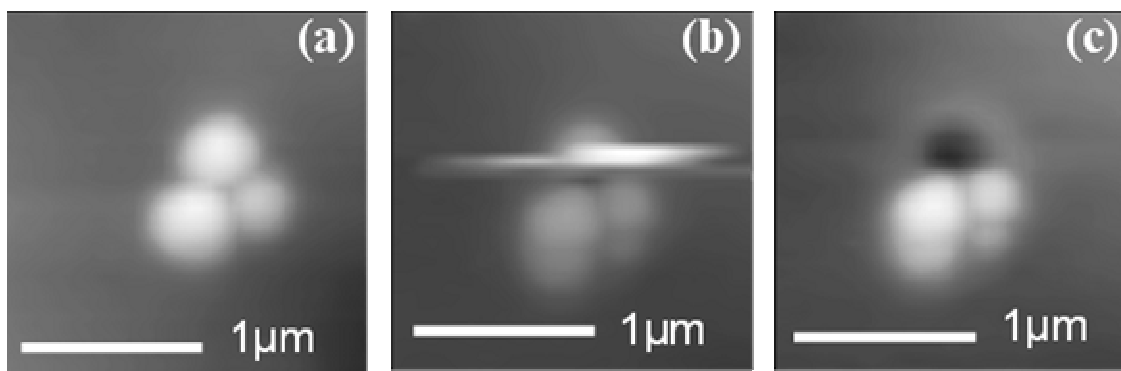


Fig. 2. Series of scanning heated tip images taken at tip temperatures at 220°C (a), 290°C (b) and 300°C (c). The streaking in image b shows the debonding event and subsequent dragging of the particle.

The lateral forces associated with debonding were determined by investigation of the lateral force microscopy signal. Figure 3 shows a characteristic lateral force scan. The large peak is associated with the torsional deflection of the cantilever as the tip contacts the particle. Eventually, the tip slides past the contact point and a slight decrease in friction relative to the matrix is observed. The dependence of the lateral force on temperature, for another thermally debonded particle, was determined by finding the local L-R signal maximum at the polymer-particle interface from the series temperature scans. As shown in Figure 4, increasing temperature leads to an increasing peak lateral force on the particle. Impact force is relatively constant below T_g , but increases significantly as the tip temperature is increased above T_g . The maximum observed force at 10°C below the debonding temperature was 220nN . The temperature dependence is likely associated with the increased time the tip spends at the particle interface as a result of the deforming lever. This increased time would allow for a locally higher equilibrium temperature at the interface, and could be a sufficient initiator for debonding.

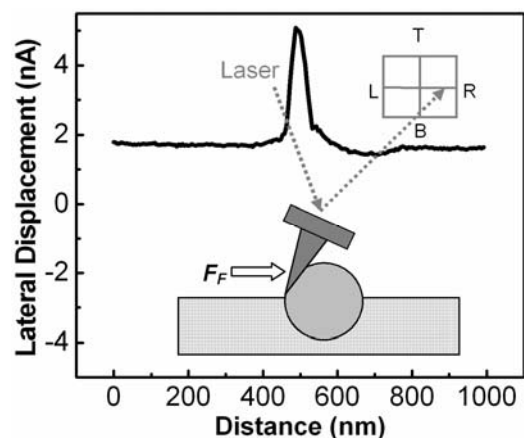


Figure 3: Characteristic forward friction scan across stable silica particle when scanned with heated tip.

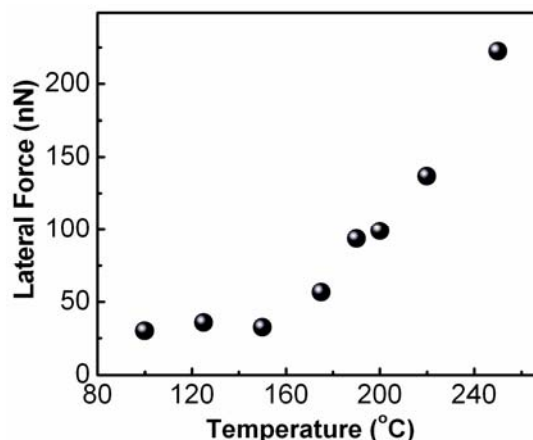


Figure 4: Maximum lateral force subjected to particle as a function of temperature.

Conclusions

HT-AFM was used to identify the matrix Tg of the composite film, while also acting as a manipulating tool for investigating the lateral forces exerted during particle-matrix debonding. By studying the particle matrix adhesion, it may be possible to design the interface for optimum reverse selective membrane properties.

References:

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