Reversible Decrease of Gel-Solvent Friction

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regions of the CH₄-N₂ and CH₄-CH₄ absorption.

Comparative planetology has appropriately focused on understanding the relation between the so-called terrestrial planets: Venus, Earth, and Mars. The large planets in the outer solar systems seem to represent a different type of object, not akin to Earth from an atmospheric science point of view. However, Titan does have an atmosphere of Earth-like composition and pressure. Preliminary studies of the data returned from the Voyager 1 flyby of Titan have allowed us to explore first the greenhouse effect of atmospheric gases other than CO₂ and H₂O and now the antigreenhouse effect due to organic haze particles. Further investigation of Titan in the upcoming NASA-European Space Agency Cassini Orbiter and Huygens Probe mission should give us more understanding of Titan's atmosphere, one that is both strange and yet similar to our own.

REFERENCES AND NOTES
6. Toon et al. (2) have determined that the cloud opacity is less than 2 at 200 cm⁻¹ and that the particles of the cloud are probably larger than tens of micrometers. This implies that the visible optical depth is also about 2. McKay et al. (5) have shown that only large particles and optically thin clouds are consistent with both the Voyager data and the surface energy balance.

7. The existence of an ocean on Titan was suggested by J. J. Lunine, D. J. Stevenson, and Y. L. Yang (Science 222, 1229 (1983)) on the basis of the CH₄ photochemistry. However, recent radar data suggest that only small lakes may exist on Titan (D. O. McElhennan, A. W. Grossman, R. J. Butler, M. A. Slade, ibid. 248, 975 (1990)). In either case atmospheric CH₄ is determined by equilibrium with this liquid reservoir.
8. Our model is described in McKay et al. (5); the parameters in the current best-fit model are as those listed in table III of McKay et al. under the no-cloud model except that: pressure P₉ = 10⁻⁶ mbar (correcting the misprint in McKay et al.), relative humidity RH₄CH₄ is taken to be 66% following Lellouch et al. (4), and the haze asymmetry scaling factor < 0.1 is 1.05. This model comes close to reproducing the temperature profile of Titan's atmosphere. The computed surface temperature is 93.67 K compared to the reported value of 93.9 K (4).

11. We assume that in this idealized antigreenhouse the layer is isothermal because this simplifies the analysis. Isothermal conditions could be maintained by adiabatic motions if the layer were physically thin. However, in general the layer would not be isothermal and the emissions from the bottom of the layer would depend on the temperature at that point. This temperature would be determined by the internal greenhouse and antigreenhouse effects within the layer itself.

12. The effective temperature is the temperature at which a blackbody would emit the same total thermal infrared radiation as the surface and atmosphere of Titan. This must balance the total solar radiation absorbed by the surface and atmosphere. In our model calculations, the effective temperature of Titan is 82 K (5).

13. The factor (1/2)²⁻¹ in temperature arises as follows: the idealized isothermal (11) antigreenhouse layer absorbs all the incoming solar radiation and emits half back to space and half to the surface. Thus, the flux reaching the surface is reduced by a factor of 1/2. Temperature is proportional to the fourth root of the flux.


16. Below the tropopause it is likely that the haze particles are coated with condensing hydrocarbons, principally CH₄, and are thereby swept from the atmosphere. See (2); C. F. Frère, thesis, University of Paris XIII (1989).

17. The troposphere does not emit as a graybody but the opacity of N₂-N₂ dominates the spectrum (Fig. 1), and an optical depth of unity at the peak of the Planck function occurs near the tropopause. At 130 cm⁻¹, near the emission peak for 73 K, an optical depth of unity occurs at an altitude of 25 km and a temperature of 73 K, in agreement with our simple estimation.

18. We thank R. E. Samuelson for a careful and incisive review that considerably improved the final paper. This work was supported by NASA's Planetary Atmospheres Program.

15 April 1991; accepted 25 June 1991

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The friction between water and the polymer network of a gel is found to decrease reversibly by three orders of magnitude and appears to diminish as the gel approaches a certain temperature at constant volume and network structure.

When water passes through a polymer network, a frictional resistance arises between the water and the network. For a permanently cross-linked network, the friction, normalized by the viscosity of water, was expected to be independent of temperature. It was found, however, that the friction decreases reversibly by three orders of magnitude and appears to diminish as the gel approaches a certain temperature. This phenomenon occurs in spite of the fact that the network structure and the total volume of the gel, and thus the network density, are unchanged.

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The configuration for measuring the friction coefficient of a gel is schematically shown in Fig. 1. Water is pushed through a gel slab of thickness d by a small pressure p. The proportionality between the velocity of the water v and the pressure determines the friction coefficient:

\[ f = \frac{p}{d} \]

To avoid any leak of water and to maintain constant gel volume, most of the gel surfaces were chemically attached to the surfaces of a pair of gel bonding plastics (Bio-Rad) by covalent bonding, except the small circular portions left open for the water flow. The open portion was mechanically pressed by rigid paper filters to prevent swelling. Small shrinkage should have oc-

Fig. 1. Schematic illustration of the measurements of the friction coefficient between a polymer network and water. In order to avoid any leak of water and to fix the gel volume, most of the gel surfaces (diameter 10 mm) were chemically attached to the surfaces of a pair of gel bonding films (Bio-Rad) by covalent bonding, except the small circular portions left open for the water flow. The small open portion (diameter 2 mm) was mechanically pressed by rigid paper filters to prevent swelling.
During the friction experiment a test tube containing a gel without a mold was placed in the same water bath and the appearance of the gel was continuously monitored. At the highest temperature of our experiment, the decrease in the free gel diameter from the isochore diameter was less than 10%. These observations confirmed that the gel swelling or shrinking as a whole should not have a significant effect on the friction measurement.

A slight opacity developed in the gel near the temperature at which the friction diminished, presumably due to the dynamic density fluctuations. As we shall see later the fluctuations seem to be responsible for the diminishing of the solvent-network friction.

The temperature dependence of the friction coefficient normalized by the viscosity of water, $f/\eta$, was measured (Fig. 3). The viscosity of water $\eta(T)$ is taken from a table. The temperature dependence of the friction of poly(acrylamide) gel is also shown in this figure [taken from Tokita and Tanaka (1)]. The normalized friction of the poly(acrylamide) gel is independent of the temperature, while the friction of the poly(N-isopropylacrylamide) gel decreases three orders of magnitude as the temperature approaches 33.6°C. The change is reversible.

The phenomenon of the reversible diminishing of friction may be intuitively explained as follows. The gel-solvent friction is primarily determined by the pore size of the polymer network and the viscosity of the solvent. The latter should not show any anomaly in our experiments. When the network is homogeneous the pore size should be given by the average distance between the nearest polymer-polymer contacts. Under certain conditions, however, the polymer network undergoes substantial density fluctuations in space and time. Some portions of the gel swell while the other portions shrink maintaining, the total gel volume constant. The effective pore size is then given by the distance over which the network density fluctuations are correlated. The water passes through the swollen open space avoiding the shrunken regions.

The pore size practically diverges as the gel passes the coexistence curve into the two-phase region, probably for one of the following reasons: First, the gel may remain in the metastable single phase as a superheated gel. The metastable state has been observed in the hysteresis of the swelling curves of various gels including NIPA gels. The temperature gap at a hysteresis can be several degrees (3). In this case the fluctuations are dynamic (time-dependent) and should diverge on the spinodal line. Second, the gel may undergo phase separation creating domains of swollen and shrunken phases. The density fluctuations are static (time-independent) and would diverge on or near the coexistence curve. In both cases the effective pore size diverges, making the friction diminish. The pore structure is not

![Fig. 2. The swelling curve (the isobar at zero osmotic pressure) of poly(N-isopropylacrylamide) gel in water. [Data taken from (3)]. The open circles are the critical point ($T_c$) and the temperature at which the friction diminishes ($T_{\alpha\beta}$). The friction measurement was carried out along the isochore at the volume of gelation. At the temperature $T_{1/2}$ the normalized friction $f/\eta$ becomes 1/2 that at 20°C.](image)

![Fig. 3. The friction coefficient, $f$, of water passing through gels as a function of temperature. Circles represent the poly(N-isopropylacrylamide) gel and squares represent the poly(acrylamide) gel of concentration of 8.8 g per 100 ml [taken from (1)]. Solid circles are used in the increasing of the temperature and open circles are used in the lowering of the temperature. For the NIPA gel, the friction is also plotted in the linear scale. The friction disappears at 33.6°C for the poly(NIPA) gel, whereas it is constant of temperature for the poly(acrylamide) gel.](image)

![Fig. 4. The log-log plot of the friction coefficient, normalized by the water viscosity $\eta(T)$, as a function of temperature difference from $T_{\alpha\beta}$. The viscosity of water was taken from a standard table. The exponent is 1/4, which is much smaller than the value 5/8 predicted for dynamic spinodal fluctuations.](image)
permanent, but is reversibly enlarged or reduced with temperature.

The temperature dependence of the dynamic contribution to pore size may be estimated by means of mode coupling theory, which views a gel as consisting of \( N \) pores of diameter \( \xi \) over which the density fluctuations are correlated \((6-8)\).

\[ f = N_0 \xi \]

The least-square analysis yields \( f \) = 1/4 and \( T_{\tau_{\text{m}}0} = 33.59^\circ \text{C} \) (Fig. 4).

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The theory further suggests

\[ N = N_0 \left( \frac{T_{\tau_{\text{m}}0}}{T} \right)^2 \]

where \( T_{\tau_{\text{m}}0} \) denotes the spinodal temperature (that is, the temperature at which the density fluctuations diverge). Thus

\[ f = \frac{6\pi N_0 \xi}{\eta} \left( \frac{T_{\tau_{\text{m}}0}}{T} \right)^2 \]

The least-square analysis yields \( v \) = 1/4 and \( T_{\tau_{\text{m}}0} = 33.59^\circ \text{C} \) (Fig. 4).

The determination of various critical exponents indicated that the phase transition of NIPA gels belongs to the universality class of three-dimensional Ising models \((4)\). The exponent 1/4 is much smaller than the theoretical critical exponent 5/8 of the three-dimensional Ising model \((9, 10)\). This discrepancy between the exponents may be because the isochore is not critical. Or it may be that the pore size diverged upon phase separation in the metastable region before the gel reached the spinodal line. In this case \( T_{\tau_{\text{m}}0} \) should be considered as the temperature at which the domain size grows to infinity rather than the spinodal temperature.

By choosing an optimal combination of solvent and temperature the phenomenon should be universally observed in any gel. A drastic and reversible change in the friction may have applications in separation technology and may be relevant to some biological transport phenomena.