- M. R. Drury and J. D. F. Fitz Gerald, *Geophys. Res. Lett.* 23, 701 (1996); R. Wirth, *Contrib. Mineral. Petrol.* 124, 44 (1996).
- M. R. Carroll, D. S. Draper, R. A. Brooker, S. Kelley, in Noble Gas Geochemistry and Cosmochemistry, J. Matsuda, Ed. (Terra Scientific, Tokyo, 1994), pp. 325–341.
- T. Matsumoto, M. Honda, I. McDougall, S. Y. O'Reilly, Geochim. Cosmochim. Acta 62, 2521 (1998).
- 26. T. J. Dunai and H. Baur, Geochim. Cosmochim. Acta 59, 2767 (1995).
- 27. G. L. Davies, *Carnegie Inst. Wash. Yearb.* **76**, 631 (1977).
- C. R. Neal et al., in Large Igneous Provinces: Continental, Oceanic, and Planetary Flood Volcanism (Geophys. Monogr. 100, American Geophysical Union, Washington, DC, 1997), pp. 183–216.
- 29. A. D. Beard et al., Lithos **39**, 93 (1996).

# A Potent Greenhouse Gas Identified in the Atmosphere: SF<sub>5</sub>CF<sub>3</sub>

### W. T. Sturges,<sup>1</sup> T. J. Wallington,<sup>2</sup> M. D. Hurley,<sup>2</sup> K. P. Shine,<sup>3</sup> K. Sihra,<sup>3</sup> A. Engel,<sup>4</sup> D. E. Oram,<sup>1</sup> S. A. Penkett,<sup>1</sup> R. Mulvaney,<sup>5</sup> C. A. M. Brenninkmeijer<sup>6</sup>

We detected a compound previously unreported in the atmosphere, trifluoromethyl sulfur pentafluoride (SF<sub>5</sub>CF<sub>3</sub>). Measurements of its infrared absorption cross section show SF<sub>5</sub>CF<sub>3</sub> to have a radiative forcing of 0.57 watt per square meter per parts per billion. This is the largest radiative forcing, on a per molecule basis, of any gas found in the atmosphere to date. Antarctic firm measurements show it to have grown from near zero in the late 1960s to about 0.12 part per trillion in 1999. It is presently growing by about 0.008 part per trillion per year, or 6% per year. Stratospheric profiles of SF<sub>5</sub>CF<sub>3</sub> suggest that it is long-lived in the atmosphere (on the order of 1000 years).

The Kyoto Protocol highlighted the need to assess a broad range of greenhouse gases for their contribution to radiative forcing. Whereas most attention has been placed on the major greenhouse gases (CO<sub>2</sub>, CH<sub>4</sub>, and N<sub>2</sub>O), it is clear that some gases that are present at much lower concentrations can contribute to global warming because of their exceptionally large infrared (IR) absorption. A notable example is  $SF_6$ , which is closely chemically related to the "new" molecule SF<sub>5</sub>CF<sub>3</sub>. SF<sub>6</sub> is of almost exclusively anthropogenic origin, currently present in the atmosphere at only ~4 parts per trillion (ppt), but with a global warming potential (GWP) of 22,200 relative to CO<sub>2</sub> over a 100-year time horizon (1).  $SF_6$  is one of the greenhouse gases that the Kyoto Protocol seeks to control.

We first noted  $SF_5CF_3$  as an unidentified chromatographic peak that eluted shortly after  $SF_6$  in gas chromatography–mass spectrometry (GC-MS) analysis of stratospheric air samples. The identity of the peak was determined by the presence of ions with masses of 68.995 ( $CF_3^+$ ), 88.967 ( $SF_3^+$ ), and 126.964 (SF<sub>5</sub>CF<sub>3</sub><sup>+</sup>) in their correct relative abundances, and the identity was then confirmed by an exact GC retention time match with a sample of pure compound (from Flura Corporation, Newport, Tennessee) diluted in nitrogen. The mass spectrum (comprising almost exclusively the three above-mentioned ions) was obtained from the pure compound because no library spectrum could be found.

Air samples were analyzed with the GC-MS system operated in single-ion mode (a mass of 88.967 was used to monitor SF<sub>5</sub>CF<sub>3</sub>), giving a detection limit of ~1 part per 10<sup>15</sup> for 800-ml air volumes (2, 3). SF<sub>6</sub> concentrations were referenced to a calibrated ambient air standard (Colorado Mountains, 1994; SF<sub>6</sub> = 3.39 ppt) (4). SF<sub>5</sub>CF<sub>3</sub> was calibrated by two separate methods (5), giving a mean concentration in the same air standard of 0.094  $\pm$  0.020 ppt.

Samples of air were pumped out of deep consolidated snow (firn) at Dome Concordia in eastern Antarctica (75°S, 123°E; 3233 m above sea level) in January 1999 (6) at various depths from the surface to pore close-off at ~100 m. Mean equilibration times between changing surface concentrations and air at depth in the firn can be on the order of years or even decades, depending on the diffusivity of the particular gas and its concentration gradient. This was calculated with the aid of a diffusive transport model constrained by measured profiles of gases with wellknown atmospheric trends (e.g., CO<sub>2</sub>) (7).

Profiles of  $SF_5CF_3$  and  $SF_6$  with depth in

- 30. We thank G. Pearson (Durham University, UK) for kindly providing the samples from Malaita and H. Downes (Birkbeck College, University of London) and P. Kempton (UK National Environment Research Council Isotope Unit) for providing the sample from Elovy Island. S.P.K. and J-A.W. greatly appreciate a grant from the Leverhulme Trust that allowed this work to be undertaken.
  - 21 April 2000; accepted 12 June 2000

the firn are shown in Fig. 1. They are remarkably similar in form, leading to the suspicion that their growth rates are closely related. Published measurements and emission estimates for  $SF_6$  were combined (8) to give the global SF<sub>6</sub> trend shown in the inset of Fig. 1. Modeling this trend gave an excellent match to the observed SF<sub>6</sub> concentrations in the firn, except for a possible slight overestimate from extrapolating the trend from 1997 to 1999. A hypothetical trend for SF5CF3 was then constructed with the SF<sub>6</sub> trend scaled to the mean relative concentration of SF5CF3 to SF6 in surface air at Dome Concordia (0.122 to 4.00 ppt) (Fig. 1). This trend was modeled with a diffusion coefficient for SF5CF3 in air estimated from molecular volume calculations (9). The resulting fit to the observed SF<sub>5</sub>CF<sub>3</sub> profile is very good (Fig. 1), with most measurements falling within an envelope of concentrations 10% above and below the mean. We therefore think that this time trend for SF<sub>5</sub>CF<sub>3</sub> is a good representation of its actual atmospheric growth and suggests that emissions began in the late 1950s.

Air was also collected from the stratosphere by using a balloon-borne liquid neonbased cryogenic sampler (10). Vertical profiles of SF<sub>5</sub>CF<sub>3</sub> are shown in Fig. 2 for both mid-latitude and Arctic vortex flights, launched from Aire sur l'Adour, France (44°N), and Kiruna, Sweden (68°N), respectively. At both latitudes, the concentrations declined monotonically with altitude. Measurements of SF<sub>6</sub> were used to determine the mean age of air (11) for each sample. The mean ages at maximum altitude were 4 and 7 years for the mid-latitude and Arctic vortex flights, respectively. The greater decline in SF<sub>5</sub>CF<sub>3</sub> with altitude observed for the highlatitude flight can be explained by subsidence of older air in the polar vortex. The SF<sub>5</sub>CF<sub>3</sub> concentrations expected on the basis of the mean age of air and the atmospheric trend of  $SF_5CF_3$  (from Fig. 1) are shown as thick lines in Fig. 2. Agreement between the predicted and measured SF<sub>5</sub>CF<sub>3</sub> is very good within the errors of the measurements, at least at lower altitudes. We therefore expect the upper limit of the lifetime of SF<sub>5</sub>CF<sub>3</sub> to approach that of  $SF_6$  [3200 years (12)]. By analogy with  $SF_6$ , we do not expect any substantial sinks of  $SF_5CF_3$  in the troposphere nor do we expect uptake by soils, plants, or the ocean (12).

Toward the top of the stratospheric profiles (Fig. 2), there is evidence of a possible divergence toward observed  $SF_5CF_3$  concentrations

<sup>&</sup>lt;sup>1</sup>School of Environmental Sciences, University of East Anglia, Norwich NR4 7TJ, UK. <sup>2</sup>Ford Motor Company, Mail Drop SRL-3083, Dearborn, MI 48121–2053, USA. <sup>3</sup>Department of Meteorology, University of Reading, Reading RG6 6BB, UK. <sup>4</sup>Institute for Meteorology and Geophysics, Johann Wolfgang Goethe University of Frankfurt, D-60325 Frankfurt, Germany. <sup>5</sup>British Antarctic Survey, Natural Environment Research Council, Cambridge CB3 0ET, UK. <sup>6</sup>Atmospheric Chemistry Division, Max Planck Institute for Chemistry, D-55060 Mainz, Germany.

Fig. 1. Measurements of  $SF_5CF_3$  (triangles) and  $SF_6$  (circles) in air extracted from firn at Dome Concordia, Antarctica, in January 1999 and firn modeling (SF<sub>5</sub>CF<sub>3</sub>, thick solid line; SF<sub>6</sub>, thin solid line) of expected expected depth profiles based on the atmospheric scenarios shown in the inset. The dotted lines shown for SF5CF3 denote concentrations that are  $\pm 10\%$  of the modeled line. Error bars on the symbols



are 1 $\sigma$  of the total accumulated analytical precision values (the bars are smaller than the symbols for SF<sub>6</sub>). The SF<sub>6</sub> trend for 1978 to 1997 was derived from the quadratic fit to measurements made at Cape Grim, Tasmania (8), extrapolated to 1999. To estimate the trend before 1978, we converted global emission estimates of SF<sub>6</sub> from 1953 to 1985 to global average concentrations using SF<sub>6</sub> concentration (ppt)  $\approx$  metric tons of SF<sub>6</sub>/25000 (8). These values were adjusted to equivalent Cape Grim concentrations using the calculated annual growth rate and a time lag of half the interhemispheric exchange time of 1.35 years (17). A third-order polynomial fit to the data yielded SF<sub>6</sub> concentrations for 1953 to 1978.

that are lower than those predicted. This is, however, very sensitive to the accuracy of the reconstructed atmospheric trend and the assumption of a constant  $SF_5CF_3$ -to- $SF_6$  ratio over time (the small offset between measured and predicted concentrations at all altitudes in the later flight suggests some deviation from these assumptions). In the top few kilometers of the profiles, measured N<sub>2</sub>O concentrations averaged ~50 parts per billion (ppb). This means that ~80 to 90% of the N<sub>2</sub>O that originally entered the stratosphere [~310 ppb in the early



**Fig. 2.** Measurements of SF<sub>5</sub>CF<sub>3</sub> from stratospheric balloon flights launched from (**A**) Kiruna, Sweden (68°N), in February 1997 (diamonds) and (**B**) Aire sur l'Adour, France (44°N), in May 1999 (squares). Error bars are 1 $\sigma$  of the total accumulated analytical precision values. Predicted values are shown as thick lines, based on scaling measured SF<sub>6</sub> concentrations to the constant ratio used to derive the SF<sub>5</sub>CF<sub>3</sub> trend in Fig. 1. Because both compounds are growing at essentially the same rate, it is not necessary to correct for nonlinear growth (22). Envelopes of  $\pm$ 10% the predicted concentrations (compare Fig. 1) are shown as thin lines.

1990s (1)] had been destroyed in this "old" air. If measured  $SF_5CF_3$  concentrations were, on average, 10% lower than those predicted from  $SF_6$ , then by analogy to N<sub>2</sub>O [lifetime of ~120 years (1)], the stratospheric lifetime of  $SF_5CF_3$ would be on the order of 1000 years. We therefore expect the lifetime of this molecule to be somewhere between several hundred and a few thousand years.

The radiative forcing due to  $SF_5CF_3$  has not previously been evaluated. The IR absorption spectrum of  $SF_5CF_3$ , measured over the range from 600 to 3800 cm<sup>-1</sup> with a Fourier transform IR spectrometer (FTIR), is shown in Fig. 3. IR bands were observed at 755, 884, 889, 903, 1172, and 1257 cm<sup>-1</sup>. IR absorption increased linearly with  $SF_5CF_3$ concentration. Absorption cross sections (base e, in units of square centimeters per molecule of  $7.6 \times 10^{-18}$  at 903 cm<sup>-1</sup> and

Fig. 3. IR spectrum of SF<sub>5</sub>CF<sub>3</sub>. Samples (from Flura Corporation, Newport, Tennessee) of 0.7 to 3.1 mtorr of SF5CF3 purified by repeated freeze-pump-thaw cycling to remove an  $\sim 2\%$ impurity,  $SF_6$ were mixed with 700 torr of ultrahigh purity synthetic air  $(N_2/O_2)$  and introduced into a 140-liter, 2-m-long, evacuable Pyrex chamber (13). The FTIR was operated at a resolution of 0.5 cmand measurements were made at 296  $\pm$  2 K.  $\sigma_{\rm e^{\prime}}$ absorption cross section (base e).

 $6.6 \times 10^{-18}$  at 1257 cm<sup>-1</sup> and integrated band strengths (units of centimeters per molecule of  $1.24 \times 10^{-17}$  (670 to 780 cm<sup>-1</sup>),  $1.45 \times 10^{-16}$  (840 to 960 cm<sup>-1</sup>), and 9.63 ×  $10^{-17}$  (1125 to 1325 cm<sup>-1</sup>) were derived. These are believed to be accurate to 5% (*13*). The strength of the IR absorption lines and the fact that 60% of the integrated cross section lies in the atmospheric IR "window" between 800 and 1200 cm<sup>-1</sup> make SF<sub>5</sub>CF<sub>3</sub> a very effective greenhouse gas on a per molecule basis.

Subsequently, the IR absorption spectrum was used to calculate an adjusted cloudy sky radiative forcing (14, 15) of 0.57 W m<sup>-2</sup> ppb<sup>-1</sup> for SF<sub>5</sub>CF<sub>3</sub>. This is the strongest radiative forcing, on a per molecule basis, of any molecule found in the global atmosphere to date (1). Indeed, only one of the more than 100 known or potentially occurring manmade gases surveyed in (1) has a stronger radiative forcing than SF<sub>5</sub>CF<sub>3</sub>. The forcing due to SF<sub>6</sub> is slightly weaker at 0.52 W m<sup>-2</sup> ppb<sup>-1</sup>.

Without better knowledge of the lifetime of SF<sub>5</sub>CF<sub>3</sub>, it is not possible to calculate the GWP with confidence, but assuming that it has the same lifetime as SF<sub>6</sub>, we estimate that its 100-year mass-normalized GWP relative to CO<sub>2</sub> (1) would be 18,000. If the lifetime were 1000 years, the GWP would be only marginally smaller at 17,500. SF<sub>6</sub>, which has a slightly higher GWP of 22,200, is the only molecule listed by the World Meteorological Organization/United Nations Environment Programme with a stronger GWP than SF<sub>5</sub>CF<sub>3</sub> (1).

It is remarkable that the trends of  $SF_6$  and  $SF_5CF_3$  have tracked each other so closely over the past 30 years.  $SF_5CF_3$  may be a by-product of the manufacture of  $SF_6$ , but we have not detected any in a sample of pure  $SF_6$ . One of the dominant uses of  $SF_6$  is in gas-insulated switchgear, transformers, accelerators, and other high-voltage equipment.



These uses have accounted for an almost constant 80% of SF<sub>6</sub> sales ( $\delta$ ); the increasing use by the electronics industry has been largely compensated by the decreasing use in magnesium smelting ( $\delta$ ). We speculate that SF<sub>5</sub>CF<sub>3</sub> originates as a breakdown product of SF<sub>6</sub> in high-voltage equipment. Such systems likely contain fluoropolymers, which provide a source of CF<sub>3</sub> groups that may be attacked by SF<sub>5</sub> radicals formed by high-voltage discharges.

Despite the large GWP of SF5CF3, the amount currently in the atmosphere is so small that the contribution of this molecule to overall radiative forcing is very minor ( $<10^{-4}$  W  $m^{-2}$ ). The present-day burden of  $SF_5CF_3$  is  $\sim 3.9 \times 10^3$  metric tons, with emissions rising by  $\sim 270$  metric tons year<sup>-1</sup> (from the trend shown in Fig. 1 and assuming negligible atmospheric loss). This emission is equivalent in GWP terms (for a 100-year time horizon) to  $\sim 1\%$  of the annual UK emission of CO<sub>2</sub> (16). SF<sub>5</sub>CF<sub>3</sub>, however, has a much longer lifetime than that of CO<sub>2</sub> (atmospheric CO<sub>2</sub> equilibration time of 50 to 200 years), and its rate of growth may be accelerating. There is, therefore, potential for an almost irreversible accumulation of this gas in the atmosphere. SF<sub>5</sub>CF<sub>3</sub>, unlike CO2, has no natural sources and therefore has the potential for substantial emission control after its sources have been identified. If it is indeed formed in high-voltage equipment, then there may already be substantial amounts "stored" in electrical installations worldwide. We think that it is important to continue monitoring the atmospheric concentration of  $SF_5CF_3$  in order to determine and control its sources and to guard against an undesirable accumulation of this strong greenhouse gas in the atmosphere.

#### **References and Notes**

- Scientific Assessment of Ozone Depletion: 1998, Global Ozone Research and Monitoring Project, Rep. 44 (World Meteorological Organization, Geneva, 1999).
- 2. Air samples (200 to 800 ml) were cryofocused at -186°C with liquid argon, desorbed at 90°C, and separated on a temperature-controlled alumina PLOT capillary column. The Micromass "AutoSpec" configuration comprised an electron ionization source, a magnetic field sector, pre– and post–electric field focusing sectors, and a photomultiplier detector.
- P. J. Fraser, D. E. Oram, C. E. Reeves, S. A. Penkett, A. McCulloch, J. Geophys. Res. 104, 15985 (1999).
- The air standard was obtained from the U.S. National Oceanic and Atmospheric Administration (NOAA), Boulder, CO. The NOAA scale agrees to within 1% of that of the University of Heidelberg, Heidelberg, Germany (1, 17).
- 5. First, pure SF<sub>5</sub>CF<sub>3</sub> (Flura Corporation, Newport, TN) was successively diluted in large stirred aluminum drums to 1 part per billion by volume, followed by injection of microliter amounts into the GC-MS using a small sample loop filled to varying pressures to span the range of concentrations that we measured in ambient air. These measurements were referenced to a synthetic calibrated 10-ppb SF<sub>6</sub>-in-nitrogen standard ["Heidelberg" scale (17)] that was prepared and analyzed in the same way. Second, the relative abundances of two ions common to both SF<sub>6</sub> and SF<sub>5</sub>CF<sub>3</sub> (masses of 89 and

#### REPORTS

127) were determined from full-scan mass spectra of diluted pure gases and compared with the relative peak areas in the NOAA standard. The second method is independent of errors arising from dilution effects, but it assumes invariant ionization efficiency. The first method yielded a concentration in the NOAA standard (collected at Niwot Ridge, CO, in spring 1994) of 0.072  $\pm$  0.009 ppt, whereas the second method gave a concentration of 0.116  $\pm$  0.018 ppt.

- 6. Air was extracted by drilling a bore hole to successive depths, each time letting down an inflatable sleeve to seal off the hole near the bottom, and then pumping air out from beneath the seal (18) into fused silicalined stainless steel sample flasks (SilcoCan canisters, Restek Corporation, Bellefonte, PA). Integrity of the pumped air was assured by continuous in situ monitoring of CO<sub>2</sub>.
- A. Fabre, J.-M. Barnola, L. Arnaud, J. Chappellaz, *Geophys. Res. Lett.* 27, 557 (2000).
- M. Maiss and C. A. M. Brenninkmeijer, *Environ. Sci. Technol.* **32**, 3077 (1998).
- E. N. Fuller, P. D. Schettler, J. C. Giddings, Ind. Eng. Chem. 58, 19 (1966).
- A. Engel, U. Schmidt, R. A. Stachnik, J. Atmos. Chem. 27, 107 (1997).
- 11. M. Strunk et al., Geophys. Res. Lett. 27, 341 (2000).
- A. R. Ravishankara, S. Solomon, A. A. Turnipseed, R. F. Warren, Science 259, 194 (1993).
- S. Pinnock, M. D. Hurley, K. P. Shine, T. J. Wallington, T. J. Smyth, J. Geophys. Res. 100, 23227 (1995).
- Intergovernmental Panel on Climate Change, Climate Change 1995: The Science of Climate Change, J. T. Houghton et al., Eds. (Cambridge Univ. Press, Cambridge, 1996).
- 15. The radiative forcing was calculated with a 10 cm<sup>-1</sup> narrow band radiation scheme in which the SF<sub>5</sub>CF<sub>3</sub> forcing was calibrated to agree exactly with line-by-line calculations [using an irradiance version of the Reference Forward Model (19)] for clear skies (20). Well-mixed vertical profiles were assumed, and the global-

mean forcing was derived by averaging three profiles representing the tropics and extratropics (21).

- J. W. L. Goodwin, A. G. Salway, T. P. Murrels, C. J. Dore, H. S. Eggleston, UK Emissions of Air Pollutants 1970–1997: A Report of the National Atmospheric Emissions Inventory (AEA Technology, Harwell, UK, 1999).
- M. Maiss and I. Levin, Geophys. Res. Lett. 21, 569 (1994).
- 18. J. Schwander et al., J. Geophys. Res. 98, 2831 (1993).
- A. Dudhia, Reference Forward Model v3 Software User's Manual, ESA Document PO-MA-OXF-GS-0003 (Department of Atmospheric, Oceanic and Planetary Physics, University of Oxford, Oxford, 1997).
- N. Christidis, M. D. Hurley, S. Pinnock, K. P. Shine, T. J. Wallington, J. Geophys. Res. **102**, 19597 (1997).
- R. S. Freckleton et al., Q. J. R. Meteorol. Soc. 124, 2099 (1998).
- 22. C. M. Volk et al., J. Geophys. Res. 102, 25543 (1997).
- This work was supported by the European Commission 23. (EC) through the FIRETRACC/100 (Firn Record of Trace Gases Relevant to Atmospheric Chemical Change over 100 Years) and HALOMAX (Mid- and High-Latitude Stratospheric Distribution of Long- and Short-lived Halogen Species During the Maximum of the Chlorine Loading) projects; by the UK Natural Environment Research Council; and by the UK Department of the Environment, Transport and the Regions. The Antarctic field work was also supported by the European Science Foundation/EC European Project for Ice Coring in Antarctica (EPICA) program, by the French Polar Institute, and by the Ente per le Nuove Tecnologie, l'Energia e l'Ambiente (ENEA) Antarctic Project (Italy). We particularly thank L. Arnaud and A. Manouvrier [CNRS-Laboratoire de Glaciologie et de Géophysique de l'Environnement (LGGE)] for firn sampling, H. McIntyre (University of East Anglia, Norwich, UK) for GC-MS analyses, and J. Chappellaz and J.-M. Barnola (CNRS-LGGE) for the firn model and supporting data

7 March 2000; accepted 9 June 2000

# CLAVATA3, a Multimeric Ligand for the CLAVATA1 Receptor-Kinase

### Amy E. Trotochaud, Sangho Jeong, Steven E. Clark\*

The CLAVATA1 (CLV1) and CLAVATA3 (CLV3) proteins form a potential receptor and ligand pair that regulates the balance between cell proliferation and differentiation at the shoot meristem of *Arabidopsis*. CLV1 encodes a receptorkinase, and CLV3 encodes a predicted small, secreted polypeptide. We demonstrate that the CLV3 and CLV1 proteins coimmunoprecipitate in vivo, that yeast cells expressing CLV1 and CLV2 bind to CLV3 from plant extracts, and that binding requires CLV1 kinase activity. CLV3 only associates with the presumed active CLV1 protein complex in vivo. More than 75% of CLV3 in cauliflower extracts is bound with CLV1, consistent with hypotheses of ligand sequestration. Soluble CLV3 was found in an approximately 25-kilodalton multimeric complex.

Although several biologically active putative ligands and putative receptors have been identified in plants (1), no proteinaceous ligand and cognate receptor pair have been identified at present. The CLV1 and CLV3 gene products

are likely candidates for such a pair. The *CLV1* and *CLV3* genes function in the same pathway, and *clv1* and *clv3* mutations exhibit dominant interactions (2). *CLV1* encodes a predicted receptor-like protein kinase with an extracellular domain composed of 21 tandem leucine-rich repeats (LRRs) and an intracellular protein kinase domain that has been shown to act as a serine kinase (3–5). *CLV3* encodes a predicted small, secreted protein (6).

Department of Biology, University of Michigan, Ann Arbor, MI 48109–1048, USA.

<sup>\*</sup>To whom correspondence should be addressed. E-mail: clarks@umich.edu