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Title of Project:

Analysis of Minerals Containing Dissolved Traces of the Fluid Phase Components "Water" and "Carbon Dioxide"

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I.1 Summary Abstract

Substantial progress has been made towards a better understanding of the dissolution of common gas/fluid phase components, notably H$_2$O and CO$_2$, in minerals. It has been shown that the dissolution mechanisms are significantly more complex than currently believed by the majority of researchers in the field. By judiciously combining various solid state analytical techniques convincing evidence has been obtained that traces of dissolved gas/fluid phase components undergo, at least in part, a redox conversion by which they split into reduced H$_2$ and reduced C on one hand and oxidized oxygen, O$^-$, on the other hand. Analysis for H$_2$ and C as well as for any organic molecules which may form during the process of co-segregation are still impeded by the omnipresent danger of extraneous contamination. However, the presence of O$^-$, an unusual oxidized form of oxygen, has been proven beyond reasonable doubt. The presence of O$^-$ testifies to the fact that a redox reaction must have taken place in the solid state involving the dissolved traces of gas/fluid phase components. Detailed information about the techniques used and the results obtained are given in the individual chapters of this Final Report.

I.2 Introduction

The work to be performed under the NASA/SETI Cooperative Agreement NCC 2-446 was concerned with traces of dissolved gases such as H$_2$O, CO$_2$ and possibly N$_2$, in minerals.

H$_2$O, CO$_2$ and N$_2$ are present as gas phase or liquid phase components in all terrestrial magmatic and metamorphic environments where minerals crystallize from melts or recrystallize at high temperatures and pressures under near-equilibrium conditions. The same gas/fluid phase components are present on other rocky solar system bodies or were present during their formation. It can further be said that gases of the C–H–O–N system were prevalent in the solar nebula and therefore in touch with every solid matter which condensed prior to the formation of our planetary system. Finally, it can be said with a high degree of confidence that C–H–O–N gases dominate in the entire known universe wherever star formation processes occur amidst molecular clouds seeded with interstellar dust.

Before this universal background of gas–solid interaction the NCC 2-446 project addressed one fundamental question: How do the gas/fluid phase components H$_2$O, CO$_2$ and N$_2$ interact with solid minerals, in particular oxide and silicate minerals? The main objective of the Project was to improve our understanding of the dissolution mechanisms of these gas/fluid components in relation to possible reaction mechanisms leading to the formation of organic molecules.

In the case of H$_2$O there is general agreement among researchers that traces of H$_2$O always dissolve in the matrices of those minerals which crystallize or recrystallize in the presence of H$_2$O,
either as vapor or fluid. There is also general agreement that the H₂O dissolution mechanism entails the ubiquitous formation of OH⁻ anions associated with defect sites in the mineral matrices.

However, in the case of CO₂ there is no general agreement whether this common gas/fluid phase component dissolves in the minerals. This is despite the fact that, on the basis of thermodynamic laws, one is forced to expect that every gas, in particular every reactive gas, dissolves in every mineral that forms in their presence. A number of authors whose work will be commented upon below have maintained over the past years that CO₂ does not dissolve in minerals even if they crystallized or recrystallized under high CO₂ pressure. I have been embroiled in a long-standing discussion, even controversy, with those who negate any dissolution of CO₂ in such mineral matrices. One of the objectives of the NASA/SETI Cooperative Agreement NCC 2-446 project was to help clarify some of the fundamental issues involved in this controversy and, if possible, offer solutions which will lead to an unambiguous answer.

These objectives have been met, though along research lines not envisioned at the time of submission of the NASA/SETI Cooperative Agreement NCC 2-446. In the following section I shall briefly review the premisses which formed the basis of this Project. I shall not at this point engage in a detailed discussion which includes literature references to my own earlier work and to the work of others in the field. This detailed discussion is given in some of the papers which have either already been published or are submitted or are being now prepared for publication.

The body of this report consists of reprints, preprints and as yet unpublished manuscripts. There are arranged according to the subtopics which they address. For the sake of uniformity of presentation all sections are formatted in the same way, e.g. even in cases where printed papers are available their texts are included here in manuscript form. The appropriate references for identification are given in the headers of each section.

### I.3 Outline of the Fundamental Questions Raised in this Project

The work to be performed under the NASA/SETI Cooperative Agreement NCC 2-446 relates to low-\(z\) element impurities, in particular H and C, in minerals which have grown in the presence of the common gases or fluids H₂O and CO₂. Contrary to widely held belief there were strong indications* that structurally dissolved gases, specifically H₂O and CO₂, in minerals undergo (at least in part) a redox conversion by which they split into reduced H₂ plus C and oxygen that is oxidized from the 2⁻ to the 1⁻ oxidation state. This redox conversion is independent of changing transition metal oxidation states but may be coupled to them in as yet to be determined ways. If generally true and applicable to a wide range of mineral phases, such a redox conversion would have far-reaching implications for many areas of scientific endeavor.

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* From the P.I.'s earlier work as described in the enclosed reprints and preprints and from other researchers.
Since any mineral dissolving gaseous components at high temperatures is likely to become supersaturated upon cooling, the diffusively mobile H₂ plus C are expected to segregate to the crystal surface, thereby producing conditions where solute H₂ would react with solute C to give organic molecules or precursors thereof.

After segregation of the reduced moieties H₂ and C the bulk of the minerals would still contain the oxidized oxygen as a "memory" of the redox conversion.

The interest of this work to NASA's Origin of Life program rests with the fact that the processes summarily described in (i)-(ii) represent an new and potentially important avenue towards the synthesis of organic molecules, in particular complex molecules. The hypothesis can be advanced that such processes might have played a role on the prebiotic Earth. As a result of tectonic and volcanic activities that have been a characteristic feature of the Earth since her earliest geologically recorded history huge amounts of minerals and rocks have passed through the global cycles: Rocks have solidified and crystallized at great depth, in the presence of the ubiquitous gas/fluid components H₂O and CO₂, and have been uplifted to the surface to be subjected to weathering. If these rocks released organic molecules as part of their weathering, they would have represented an untapped source of complex organic molecules.

The NCC 2-446 project started with an attempt to characterize step (i), e.g. the formation of organic molecules produced by the segregation of solute H₂ and solute C from within selected minerals. The work performed concentrated on single crystals of olivine, the dominant mineral phase in the upper mantle, ideally (Mg,Fe)₂SiO₄ with the Mg:Fe ratio approximately equal to the solar abundances. Since the upper mantle is a region where high temperatures and high pressures prevail — with CO₂ being the dominant fluid phase component followed by H₂O and other quantitatively less important gases — the choice seemed rational. However, the difficulties confronting the attempt of reliable analyses of solute H₂ and C are horrendous. Any laboratory study of organic molecules on mineral surfaces has to first demonstrate that these organic molecules are not the result of contamination by C-bearing gases from the environment. Given the omnipresence of C-bearing gases in natural and laboratory environments and given the extreme sensitivity with which the analyses for organic molecules have to be performed, it is a daunting task to establish proof beyond reasonable doubt of the absence of contamination. The only reasonable approach would be to prepare minerals in the presence of isotopically labelled gases, e.g. D₂O and ¹³CO₂, and then search specifically for isotopically labelled organic molecules. This task was not part of the NCC 2-446 project**.

In this Report I present as yet unpublished manuscripts describing experiments that were carried out as part of the NCC 2-446 project. The publication of the complex gas evolution observed upon impact fracture of mantle-derived olivine crystals (Chapter II) has been delayed due to objections raised by anonymous reviewers who seem to subscribe to the notion that the

** I have meanwhile submitted a Proposal to NASA's Life Science Division for carrying out such a new project.
postulated redox conversion of dissolved fluid phase components does not take place and that CO₂ does not dissolve in the solid state.

I have therefore turned to step (ii) outlined above: The search for oxygen in the unusual (higher) oxidation state 1−. The rationale was that, if oxidized oxygen can be shown to be present in minerals crystallized in an overall reducing H₂O/CO₂ atmosphere, this was by itself a valid proof for the above-mentioned redox conversion. Because of the fundamental nature of this reaction the effort was focussed on a structurally and chemically simple compound: Magnesium oxide. MgO crystals are available in high purity grades, e.g. transition metal free, grown by a process known as carbon arc fusion from a melt (at 2800°C) that is saturated with the highly reducing atmosphere of an electric carbon arc. According to fundamental thermodynamic laws such crystals should carry a “memory” of the viciously reducing conditions that prevailed during their synthesis. The least that one would expect to find in them is oxygen in an oxidized state, e.g. O−, because such O− is the hallmark of highly oxidizing synthesis conditions.

The major difficulty with respect to O− (in contrast to O²−) was that nobody had ever tried to search for this unusual oxidation state of oxygen, except in the context of radiation damage of solids. Prior to the project NCC 2-446 no technique was known which could be used for investigating O− in situ in mineral matrices. Therefore I have embarked on an additional project to develop analytical tools to study O−. This effort led to the discovery of a new physical technique now known as Charge Distribution Analysis. CDA takes advantage of the fact that the O− represent mobile electric charge carriers, so-called positive holes, which occupy shallow acceptor levels near the valence band of wide band gap insulators. In essence, CDA consists of measuring contact-free the dielectric response in an electric field gradient at the limit of 0 Hertz. Since its inception in 1989 CDA has proven to be an extremely valuable and versatile technique. It has been thoroughly tested and demonstrated its applicability in a detailed study of MgO single crystals. CDA has been applied to a variety of materials, synthetic and natural, single crystalline, polycrystalline and glassy, as is documented in the collection of reprints and preprints which are assembled at the end of this Report.

I have also pursued another diagnostic property of the O−, namely the fact that O− is paramagnetic (while O²− is diamagnetic). By measuring the anomalous paramagnetism associated with the formation of O− in transition metal free MgO single crystals the conclusions drawn from the CDA data have been substantiated. It can safely be stated that the presence of O− in carbon arc fusion grown MgO single crystals has now been established by two fully independent physical techniques of which one, the paramagnetic measurements, gives a unique response which cannot be otherwise interpreted. In this sense the existence of O− in the crystals under study has been unambiguously proven. Thus, returning to the above two premisses, it can be firmly stated that with (ii) proven to hold, (i) must also be true.

Chapter III–V of this Report consist of a three-part manuscript submitted to the Journal of Geophysical Research but not yet available in the open literature.
GASES RELEASED FROM OLIVINE BY IMPACT FRACTURE

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Abstract

Mantle-derived olivine crystals from San Carlos, AZ, USA, were fractured and the gas emission was monitored by three techniques: fracto-emission mass spectroscopy (FEMS), surface analysis by laser ionization (SALI) and field ionization mass spectrometry (FIMS). The spontaneous emission of neutrals, measured for 1.5 sec after fracture, includes H₂, H₂O, CO/N₂ and CO₂, organic molecules producing fragments at 63, 65, 77, 91 and 105 amu plus Mg vapor at 24 amu. The emission of H₂O and CO₂ is often less intense than that of complex organic molecules. After fracture in air, followed by introduction into UHV and heating to 500°C, the crystals continue to release complex C-H, C-O-H and probably C-N-O-H molecules. The following compounds have been tentatively identified: benzene, toluene, benzaldehyde/xylene, naphthalene/azulene, biphenylene, fluorene, diphenylmethane, phenanthrene/anthracene, pyrene, chrysene, perylene/biphenylene, fluorine, benzopyrene, coronene, benzopyrene/dibenzopyrene. Contamination from external sources is unlikely. The results are interpreted assuming that, under upper mantle conditions, the fluid phase components H₂O, CO/N₂ and CO₂ enter into dilute solid solution with olivine, forming H₂ and reduced C- and/or N-bearing solute entities besides oxidized (peroxy) entities. Supersaturation during cooling and decompression leads to preferential segregation of reduced solutes (H₂, C and N) to dislocations, subgrain boundaries etc. The observed organic molecules are believed to derive from precursor entities formed at those segregation sites with preformed C-C, C-H and possibly C-N bonds, including aromatic and polyaromatic precursors. Oxygen is consumed at the fracture surface for forming H₂O/CO₂/O₂ through oxidative reactions which involve O⁻ states. Mg²⁺ cations which become coordinatively unsaturated are converted to Mg⁰ and lead to the observed Mg vapor emission.
Introduction

The fluid phase in the upper mantle is part of the geodynamic cycle of volatiles. It is best described as C-N-O-H system with minor additions of sulfur and halides (e.g. Burnham 1979; Eggler and Baker 1982). The major component is CO₂ with H₂O a distant second and occasional N₂, plus CO and CH₄ (Wyllie 1978; Arculus et al. 1984; Bergman and Dubessy 1984). The amount of N₂ remains uncertain and may be as low as 1 ppm as measured by fusion extraction (Becker and Clayton 1977; Sakai et al. 1984) or as high as ~20 ppm as measured by neutron activation analysis (Norris and Schaefer 1982).

Besides filling the intergranular space the fluid phase components may be consumed to form carbonates, hydrous minerals such as micas, amphiboles, humite minerals and possibly brucite. Nitride or oxynitride minerals have not yet been reported.

C-N-O-H compounds also occur in fluid inclusions which are held to represent samples of the free fluid phase trapped in microcracks during deformation and subsequent healing (Roedder 1972, 1984; Murck 1978; Bergman and Dubessy 1984).

A third possibility to store C-N-O-H in the mantle is by forming solid solutions. Evidence for solid solution with H₂O is provided by the ubiquitous occurrence of OH⁻ in olivine (Wilkins and Sabine 1973; Beran and Putnis 1983; Freund and Oberheuser 1986; Miller et al. 1987) and other nominally anhydrous minerals. The characteristic infrared (IR) absorption bands of OH⁻ lie in a spectral region where silicate minerals have little or no intrinsic absorption. Hence, they are easily detected.

CO₂-related solute species are more difficult to ascertain spectroscopically. such as carbonate anions, CO₃²⁻, are more difficult to ascertain. Different C-bearing oxyanion complexes, unknown from bulk minerals, must also be considered. Yet, the possibility of solid solution formation between CO/CO₂ and olivine has been questioned (Mathez et al. 1984; Tsong et al. 1985; Mathez 1987), while in quenched high pressure silicate glasses carbonate anions are believed to exist besides molecular CO₂ (Mysen and Virgo 1980; Fine and Stolper 1985).

Unfortunately most techniques by which carbon in minerals may be analyzed such as photoelectron spectroscopy and secondary ion mass spectroscopy are very sensitive to surface contamination (Oberheuser et al. 1983; Tsong et al. 1985; Freund 1986). Also the ¹²C(d,p)¹³C nuclear reaction, a less contamination-sensitive technique (Wengeler et al. 1982; Oberheuser et al. 1983), has been criticized (Tsong et al. 1985; Mathez 1987). Shevabreyeva et al. (1987) used a combination of two nuclear reaction, ¹²C(d,p)¹³C and ¹²C(d,n)¹⁴N, of which the latter is a bulk technique, and concluded that carbon exists in olivine at the level of several hundred ppm as reported earlier by Oberheuser et al. (1983).
In the present study we pursue our quest for methods and techniques which enable us to obtain more information about fluid phase components in olivine. The method of choice is the on-line measurement of gases and vapors released upon impact fracture in ultrahigh vacuum and during subsequent heating.

Dickinson et al. (1982, 1985) showed that fracture of brittle materials is accompanied by the emission of electrons, of ions, of excited and neutral atoms or molecules, and of electromagnetic radiation from ultraviolet to radio frequencies. Some emissions occur only during crack formation and decay rapidly. Others last longer or be delayed.

**Experimental**

Three separate experiments were performed: (i) Fracto-Emission Mass Spectrometry (FEMS), (ii) Surface Analysis by Laser Ionization (SALI), and (iii) Field Ionization Mass Spectroscopy (FIMS).

FEMS involves the measurement of the gas emission upon fracturing a single crystal as shown schematically in Fig. 1 and as described by Dickinson et al. (1986). The crystal was located 12 mm in front of the ionizer (70V) of a fast-scanning UTI 100C quadrupole mass spectrometer. The span of the three point bend fracture device was 2.5 mm. The signal output was digitized at a rate of 100 µsec per channel with a LeCroy Data Acquisition System. Fracture was induced by a stainless steel wedge (0.75 mm radius) advancing at =10 cm/sec. The moment of fracture was determined by an acoustic transducer which triggered the data acquisition system. The total pressure change was monitored by a Bayard-Alpert gauge, positioned =20 cm from the sample with a 20 msec electronic response time. In most runs the MS was operated in the scanning mode covering a preselected range of atom mass units (1-110 amu) at a rate of 80 msec/scan. While the fracture lasted typically =10 µsec, the last residual gas spectrum prior to fracture and the gas emission after fracture were recorded over =1.6 sec. Some runs were done in the single mass mode to follow the evolution of one specific gas species with higher time resolution.

The stainless steel ultrahigh vacuum (UHV) system was pumped with a magnetically suspended 400 l/sec turbomolecular pump and a He cryopump giving background pressures in the low 10^-8 torr range. Simulated runs with no sample in the fracture device were performed to test to what extent the motion of the bellow and mechanical shock led to the release of gases. The observed gas emission was weak and comprised only gases which were already seen in the residual gas spectrum.

The SALI technique has been described by Becker and Gillen (1984). It uses a high intensity (10^8-10^9 W/cm^2) ultraviolet (193 nm) pulsed (20 nsec) eximer laser beam for ionization of gas molecules (except H_2, H_2O, CO, CO_2, N_2 and a few others) by non-resonant multiphoton
processes in conjunction with a time-of-flight mass spectrometer for analysis. The base pressure in the UHV system was in the $10^{-9}$ torr range. The samples, freshly fractured fragments of 2-4 mm size, were mounted on a heatable carrier plate. The gases analyzed freely evaporated from the sample surface either at RT or upon heating. The ionizing laser pulses intersected the gas ≈1 mm above the sample surface. Depending on the signal strength the data were collected and averaged over 100-200 or more laser pulses.

FIMS is a "soft" ionization technique. It uses the steep electric field gradient at the tip of carbon whiskers generated by 5000 V to field-ionize the gases which emanate from a sample contained in a heatable fused silica sample tube. Organic compounds are ionized. The mass spectrometer, a 60° magnetic sector instrument (resolution 1300), was pumped with diffusion pumps to a vacuum in the $10^{-6}$ torr range.

While FEMS 70 V electron beam ionization and SALI laser multiphoton ionization result in strong fragmentation of the gas molecules analyzed, making the assignment of the observed mass fragments to specific molecules difficult, the "soft" field ionization in the FIMS experiments produces mostly parent molecule ions which renders the assignment relatively easy.

Samples and Sample Preparation

The olivine single crystals used in this study came from the San Carlos Indian Reservation, Arizona, USA. They were rounded, 5-20 mm, dull green and optically slightly turbid. Thin sections show a polygonized substructure of slightly misoriented subgrains separated by dislocation walls, probably formed during mechanical deformation in the mantle (Kohlstedt and Goetze 1974; Poirier 1985). Some crystals showed μm-sized veins, filled with fine-grained material, presumably of metasomatic origin (Lloyd 1987). Though some crystals may have contained fluid inclusions, our fracture experiments rarely produced pressure bursts signaling the cracking of gas-filled inclusions.

The olivine crystals fractured softly, even when visually crack-free. This softness contrasts with the usual brittleness of olivine and is probably due to its pronounced domain substructure. Scanning electron microscope examination reveals steps and terraces which appear to indicate subgrain boundaries along which the fracture would preferentially occur. Fractures may have also occasionally followed cracks containing alteration products. The observations reported below are typical of a fracture behavior which creates fresh fracture surfaces presumably cutting through intact crystal volume.

For FEMS olivine crystals of the appropriate size were cut in halves with a diamond saw, mounted with acetone-soluble resin and ground under water to 1-2 mm thickness. The unpolished slices were removed from the support, cleaned ultrasonically in acetone and $\text{H}_2\text{CCl}_2$ or $\text{H}_3\text{C}_2\text{Cl}_3$. 
degassed in vacuum (=10^-6 torr) at RT for 8-12 hrs. Thereafter they were spring-loaded in batches of 20 into the sample carousel and re-evacuated (<10^-8 torr) for several hours prior to fracturing.

FEMS was tested extensively using different materials, including glass microscope slides, high purity fused silica and synthetic MgO single crystals (Dickinson et al., 1986, 1987). Blank runs with fused silica (Hereaus-Amersil Suprasil™) produced a short-lived gas pulse during fracture and a slight increase of the mass 28 (CO/N2). No other gases were observed.

For SALI selected olivine crystals were ultrasonically cleaned with hot water, acetone and H2CCl2 and rapidly dried. The crystals were crushed in air, 2-4 mm sized pieces were hand-picked with stainless steel tweezers, mounted beneath a fine W mesh or a stainless steel washer on a heatable plate and introduced into the UHV (10^-9 torr range) within 2-4 min. The gas evolution was monitored at ambient temperature and during heating to 450°C. Procedural blanks with fused silica and pyrex glass showed no gas emission.

For FIMS selected olivine crystals were ultrasonically cleaned in water, acetone and H2CCl2, dried and gently crushed in air in an alumina mortar under a laminar flow hood. The coarse powder (40-80 mg) was loaded into the fused silica capillary sample tube, evacuated for 30 min at 50°C to the 10^-5 torr range and then heated linearly at 6°C/min to 500°C. The mass data (24-500 amu) were averaged over 4 min intervals.

Results: FEMS Experiments

Fig. 2 shows typical examples of the gas emission from olivine fracture (a) five consecutive 80 msec scans covering the first 320 msec out of the total 1.6 sec duration of the experiment and (b) six scans of which the first three are consecutive and the remaining three spaced in time. In both cases scans (1) show the residual gas spectrum just before fracture, dominated by H2, H2O, CO/N2 and CO2. Scans (2) contain the fracture near 60 amu followed by emission at 63, 65, 77, 91 and 105 amu ranging from moderate (a) to very strong (b). Scan (3) in Fig. 2a shows off-scale intensities for 18 amu (H2O), 24 amu, and 44 amu (CO2) and nearly off-scale intensity for 28 amu (CO/N2). Intensities for H2O, CO/N2, CO2 and the signal at 24 amu continue to increase in scans (4) and (5). In Fig. 2a the H2 evolution reaches a maximum in scan (4), decaying slowly thereafter. The run shown in Fig. 2b is characterized by a shorter pulse of the H2 emission and remarkably strong emission at 77, 91 and 105 amu remaining clearly detectable after >1 sec.

Other strong peaks lie at 27, 29, 39, 41, 51 amu. Together with the 63, 65, 77, 91 and 105 amu peaks and others with lesser intensities they generally fit 12, 13 or 14 amu increment sequences, suggesting that these peaks represent fragments of series of complex hydrocarbons. Fragments of chlorinated hydrocarbons were never detected, indicating that no solvent had remained after the pre-experiment evacuation procedure.
Fig. 3 shows another typical FEMS run, 3·10⁻⁸ torr full scale, with the intensity data plotted on a logarithmic scale. Fracture occurred just before 60 amu causing a initial, intensive burst of ions, atoms and molecules. After the initial burst has decayed masses 65, 77, 91 and 105 amu are seen to go off scale within <32 msec of the fracture. In the following scan, 40-112 msec after fracture, the intensity at 2 amu (H₂) is up by a factor of nearly ten, the intensities at 18 amu (H₂O), and 51 amu are off scale. Surprisingly the peaks at 28 and 44 amu (CO/N₂ and CO₂ respectively) remain of moderate intensity, indicating little CO/N₂ and CO₂. In fact the amount of CO/CO₂ released in this experiment is less than that of the complex hydrocarbons. Prominent peaks lie at 63, 65, 79, 91 and 105 amu. The remaining scans in Fig. 3 cover the time interval 120-192 msec and 200-272 msec after fracture. While all peaks decrease in intensity, including those for CO/N₂ and CO₂, the intensity of the H₂ emission increases. Also the 24 amu intensity increases, while that of the 56 amu species decreases slowly. Numerous other peaks, in particular those 63, 65, 77, 91 and 105 amu, remain relatively strong up to ~1 sec after fracture.

Occasionally intense pressure bursts were observed which shut-off of the ionization source. Such pressure bursts are caused by fluid inclusions. Most of our samples, however, yielded spectra which resembled those shown in Figs. 2/3 with sizeable but not excessively large amounts of H₂H₂O and N₂/CO/CO₂ and fragments of complex organic molecules. More often than not the emission intensities of some organic fragments, in particular at 77, 91 and 105 amu, equalled or exceeded the intensities of the common gases.

Little intensity was observed beyond 105 amu. This suggests that the cut-off occurs around this molecular weight. One experiment was performed at elevated temperature, at ~50°C, using the radiation of the ionization filament as a heat source. In this run the range of gases emitted extended to ~150 amu.

In most experiments the intensity at 24 amu was strong and increased with time, typically for 100-300 msec. This behavior resembles the 24 amu emission from MgO observed with 100 μsec resolution. The 24 amu emission from MgO increased with time up to after ~300 msec when it suddenly stopped (Dickinson et al. 1986). The 25 and 26 amu emissions followed the same kinetics. It was concluded that the 24/25/26 amu emission arises from Mg atoms, e.g. 25Mg, 25Mg and 26Mg isotopes, vaporizing from the fracture surface. Though we have not yet performed dedicated olivine fracture experiments with μsec time resolution, the 24 amu emission from olivine depicted in Figs. 2/3 appears also be due to Mg atoms. This assignment is supported by a qualitative assessment (at 80 msec time resolution) of the 25 and 26 amu emission.

Another outstanding mass peak shown in Figs. 2/3 is that at 56 amu: it often appears with medium intensity, is strongest immediately after fracture and decreases with time. Whether it is due to Fe emission requires confirmation by further experiments.

Occasionally we noticed an enhanced emission at 32 amu, suggesting O₂. In the case of MgO fracture an O₂ emission was unequivocally established. This was surprising because the MgO
crystals had been grown under extremely reducing conditions of a carbon arc (Dickinson et al. 1987). In the case of olivine the $O_2$ emission is still uncertain, mostly because fragments from many different molecules occur in the low mass range.

From the pressure rise of the Bayard-Alpert gauge and the known volume of the UHV system the total amount of non-condensable gases released at a typical FEMS fracture experiment was estimated to be of the order of $10^{15}$-$10^{16}$ molecules/cm$^2$, equivalent to 1-10 monolayers assuming a smooth fracture surface. Complex "organic" molecules account for a substantial fraction of the total of emitted gases.

**SALI Experiments**

The purpose of the SALI experiments was to test whether the same gases observed upon impact fracture in UHV can also be observed when the olivine crystal was fractured in air and then introduced into the UHV system. Since nonresonant multiphoton ionization is $\approx 10^4$ more effective than the 70 eV electron ionization used in the FEMS experiments, the high SALI sensitivity allows for the detection of low partial pressures.

Fig. 4a/b shows the gases detected by SALI from an olivine single crystal fragment prepared and introduced into the UHV system as described above. The SALI spectrum displays many of the same mass peaks which were observed in the FEMS experiments, e.g. at 105, 91, 77, 65, 63 and 51 amu.

The fact that mass peaks 105, 91, 77, 65, 63, 51 etc. are observed in the SALI experiments, e.g. after cleaving the olivine crystal in air and exposure to air for as long as $\approx 4$ min, indicates that the compounds which produce these mass peaks are not adversely affected by short exposure to air. On the other hand certain peaks observed in FEMS experiments will obviously not appear in SALI spectra, in particular those due to Mg vapor, e.g. at 24 amu for Mg$^+$. The strong peak at 24 amu in Fig. 4a is not due to Mg$^+$ but to the C$_2^+$ fragment which forms when hydrocarbons are ionized by multiphoton ionization. Note that the complete series of skeleton C$_n^+$ ions, 1$<n>6$, can be clearly seen in Fig. 4a with peaks at 12, 24, 36, 48, 60 and 72 amu.

Fig. 4b shows the mass range above 110 amu. Many peaks are detected up to $\approx 400$ amu, documenting the high sensitivity of the SALI technique. The progression is generally by 13 amu, suggesting the addition and/or removal of CH units. There also seems to be a progression by 50 amu, indicating polycondensed aromatic rings. If the peak at 78 is due to benzene, C$_6$H$_6$, then the 128 amu peak could be due to naphthalene and/or azulene, C$_{10}$H$_8$, and the 178 amu peak to anthracene and/or phenanthrene, C$_{14}$H$_{10}$.

At RT the intensities decrease slowly as a function of time. For instance, if the intensity of the 105 amu peak was 2600 relative units $\approx 1.5$ min after introduction into the UHV sample chamber (5-6
min after fracture), it dropped to 1600 and 740 after ≈5 and ≈20 min, respectively. Upon heating the intensities increase, passing through a maximum and decay. The integrated intensities of all "organic" compounds released during heating is estimated to be of the order of \(10^{15} \text{ cm}^{-2}\) corresponding to approximately one monolayer (Becker et al. 1986).

Fig. 5a compares the intensity of the 105 peak with that of the 106 peak during heating to 360°C: both masses exhibit the same "thermal profile", suggesting that they are related to the same parent molecule. Fig. 5b compares the "thermal profiles" of 91, 105 and 165 amu. The 91 amu species is a dominant fragment of toluene, \(\text{C}_8\text{H}_5\text{CH}_3\), and a fragment of xylenes, of other benzene derivatives as well as dienes. Since there is no apparent correlation between 91 and 105, 91 appears not to be a fragment of the same compound that produces 105. The thermal profile of mass 91 follows that of mass 165 indicating that 91 is a fragment of or chemically related to 165.

Figs. 5c and 5d show that 39, 63 and 65 follow the "thermal profile" of 91. By contrast 27, 50 and 51 amu behave differently. Since the peaks at 26 and 27 amu are most like due to the diagnostic N-bearing fragments \(\text{CN}^{+}\) and \(\text{HCN}^{+}\), their correlation with species at 50 and 51 amu suggests that the latter two also arise from N-bearing compounds, possibly nitriles.

Pyrex and fused silica procedural blanks did not produce any detectable gas emission. A fresh fragments of a gem-quality olivine single crystal which we included in our SALI study produced faint signals of some of the higher mass fragments as observed before. The observation indicates that the gem-quality olivine may emit minute traces of the same complex organic molecules as the optically slightly turbid crystals. At the same time the essentially "negative" result with the gem-quality olivine shows that cleavage of the crystal in air and its exposure to the laboratory environment did not lead to the contamination of the olivine surface with organic compounds from the laboratory air. It thus confirms that the strong gas emission from the other set of olivine crystals is genuine.

FIMS Experiments

Fig. 6 shows the sum of the intensities of all mass peaks from 40 to 500 amu accumulated from the 80 mg powder sample between 50-500°C. The four most intensive peaks occur at 302, 43, 106 and 128 amu.

Since FIMS produces predominantly parent molecule ions, an assignment of the observed peaks is much easier than by the other two techniques: many mass peaks can be identified as aromatics. At the same time we note that mass peaks which were strong in the FEMS and SALI experiments are often shifted by one amu to higher values. For instance instead of 105 amu one observes a strong 106 amu peak.
If 106 amu represents a parent molecule ion, it may be benzaldehyde, $\text{C}_6\text{H}_5\text{CHO}$, or any xylene, $\text{C}_6\text{H}_4(\text{CH}_3)_2$. The choice is narrowed by considering the intensities of toluene, $\text{C}_6\text{H}_5\text{CH}_3$ at 92 amu which is closely related chemically. The accumulated intensity at 92 amu is low. Therefore it is unlikely that the main contribution to 106 amu comes from xylene, $\text{C}_6\text{H}_4(\text{CH}_3)_2$. Thus the 106 amu peak is probably due to benzaldehyde.

The "thermal profile" of 107 amu is different from that of 106 amu, indicating that the parent molecule at 107 amu belongs to a different group. It may represent a pyridine derivative substituting N for CH in the aromatic ring.

Other peaks in Fig. 6 which can be assigned to likely parent molecules are: 78: benzene, 128: naphthalene or azulene, 152: biphenylene, 166: fluorene, 168: diphenylmethane, 178: phenanthrene and anthracene, 202: pyrene, 228: chrysene, 252: perylene or benzopyrene, 300: coronene, 302: benzopyrelene or dibenzopyrene, 350: benzocoronene.

Fig. 7 shows selected FINIS thermal profiles. Some thermal profiles are quite similar indicating that they belong to compounds which are chemically related such as the series 84, 98, 112 (not shown in Fig. 7) and 126. Each is separated from the next one by 14 amu, suggesting the addition of CH$_2$ units. They rise slowly and reach maximum intensity at or above 300°C. Compound 106 amu (probably benzaldehyde) appears first with its maximum intensity between 150-200°C, followed by the 302 amu compound (benzopyrelene or dibenzopyrene) which passes through a maximum around 200°C.

For a simple distillation process lower molecular weight compounds are expected to appear first during heating, e.g. at lower temperatures. Benzene at 78 amu is observed only at high temperatures, >400°C, with increasing intensity. This suggests that benzene is predominantly a pyrolysis product.

Fig. 7 provides further information about the parentage of the 106 amu compound. Its intensity reaches a maximum around 180°C but displays a second maximum at 450°C. This latter maximum coincides with a maximum for toluene (92 amu) and the increasing intensity of benzene (78 amu). It is tempting to suggest that, while the lower T maximum of 106 amu is probably due to benzaldehyde as indicated above, the higher T maximum may due to xylene pyrolyzing to toluene and benzene.

**Discussion**

The sustained emission of gas molecules during and after impact fracture of mantle-derived olivine as measured by FEMS suggests that the olivine crystals contain gaseous components. The gases include common constituents of the C-N-O-H mantle fluids such as H$_2$, H$_2$O, CO/N$_2$ and CO$_2$ but also "organic" C-N-O-H gases in the mass range up to ≈110 amu. The emission intensity of
the "organic" compounds often equals or exceeds those of the simple stable gases. Furthermore the FEMS experiments show emission at 24 amu which is probably due to Mg vapor.

According to the SALI experiments the complex C-N-O-H gases continue to be emitted for relatively long time after the crystals were fractured in air and exposed to air for a few minutes. The emission is enhanced by heating, giving rise to distinct "heating profiles". With the high sensitivity afforded by SALI the mass range could be extended to ~250 amu. There are indications that some of the complex "organic" molecules are N-bearing.

The FIMS experiments finally extend the mass range of the evolved C-N-O-H gases beyond 300 amu and provide clues as to their chemical identity. They suggest that many of the emitted gases are polyaromatic hydrocarbons.

Possible In Situ Contamination

The FEMS experiments are designed to minimize, if not exclude, in situ contamination of the fresh fracture surface from the residual gas. From the residual gas spectra we know that the UHV system did not contain any of the "organic" molecules observed after fracture, down to a level <10^-9 torr (see scan 1 in Fig. 2).

Because the FEMS data are collected rapidly during and after fracture, within <1.5 sec, residual gas molecules cannot adsorb onto the fresh surface. Gas kinetic theory gives the number of gas molecules which collide with a surface per unit time at a given gas pressure. Even if we assume that each colliding molecule becomes chemisorbed, at ~10^-9 torr their number is so small that it would take ~1000 sec to form one monolayer. For any potential contaminant at a partial pressure of ~10^-9 torr to form a monolayer would require ~1000 sec. Therefore the maximum contamination from the residual gas is >10^-3 monolayer.

This is corroborated by the absence of any noticeable C-N-O-H emission during procedural blank runs and earlier work with MgO (Dickinson et al. 1986, 1987). The procedural SALI blank runs have additionally demonstrated beyond reasonable doubt that the observed C-N-O-H molecules are not contaminants from the ambient laboratory air, adsorbed after fracturing.

Next we address the question whether "organic" C-N-O-H molecules could be synthesized on the fresh fracture by surface-catalyzed Fischer-Tropsch reactions from H2, CO (and NH3), if these gases exist in the olivine crystals under study. We have noted above that 1-10 monolayers of "organic" molecules are estimated to be emitted in a typical FEMS experiment. For a surface-catalyzed reaction to produce that much gas within ~usec to msec at RT the reactants must first become adsorbed on the surface. Even if H2 and CO (and NH3) had been abundantly available in fluid inclusions (for which there is no evidence), before these gas molecules become catalytically reactive they must "stick" to the surface. Barker and Torkelson (1975) have demonstrated that
freshly broken quartz and basalt surfaces, when exposed for several minutes to gases such as CH₄, CO and CO₂ at pressures of several torr, will adsorb only a fraction of a monolayer. Hence, it appears unlikely that Fischer-Tropsch synthesis could produce the observed "organic" molecules under the FEMS conditions, e.g. on a μsec to msec time scale.

**Other Contamination Sources**

If the "organic" molecules are not products of *in situ* contamination nor synthesis, where, when and how were they introduced into the olivine crystals under study?

The three possibilities are: the organics were formed or acquired either (i) in the mantle, (ii) during ascent with the volcanic magma, or (iii) after emplacement at the surface of the earth.

(i) Fluid inclusions in mantle-derived minerals are generally believed to represent samples of mantle fluids trapped by microcrack formation and subsequent healing (e.g. Roedder 1984). Fluid inclusion studies show CO₂ to be the most abundant component in olivines, generally far in excess of H₂O and only occasionally accompanied by minor components such as CO, SO₂, COS, H₂S, H₂, CH₄ and N₂ (Roedder, 1965; Murck et al., 1978; Bergman and Dubessy, 1984). NH₃ and complex C-H, C-O-H or C-N-O-H molecules have never been reported except for traces of C₂H₄ and C₂H₆ in some mantle-derived chromites (Johan et al., 1983) and light to medium molecular weight hydrocarbons in minerals from a highly alkaline felsic intrusion (Konnerup-Madsen et al. 1979).

(ii) Next we consider the possibility that "organic" molecules were introduced into the olivine crystals during ascent. Mathez et al. (1984) and Mathez (1987) hypothesize that volcanic C-N-O-H fluids penetrate into the olivine crystals along microcracks and become catalytically reduced to carbon which could develop "organic" compounds upon further reactions.

Though no mechanisms have been proposed by which volcanic fluids can be catalytically reduced on olivine surfaces, we concede that reactions by some as yet unknown mechanism may not be entirely ruled out. If they occur, the resulting "organic" molecules would line the walls of existing microcracks.

Infrared spectroscopy (IR) is a sensitive probe for the detection of C-H in olivine which is intrinsically transparent around 3000 cm⁻¹. **Fig. 8** shows the IR spectrum between 3500 - 2000 cm⁻¹ of a 6.7 mm thick San Carlos olivine crystal from the same batch of crystals which produced the rich suite of complex "organic" molecules on which report in this paper. Except for a slight undulation of the background no IR absorption band is detected between 2800-3100 cm⁻¹, the range of the C-H stretching vibrations. This contrasts with MgO single crystals synthetically grown in a slightly moist electric carbon arc atmosphere where a nC-H band was clearly observable consisting of two broad absorption maxima (Freund and Wengeler 1982). The MgO crystals
emitted CH₄ plus several simple hydrocarbons upon fracture (Dickinson et al. 1986). However, this broad IR double band is unlike any band molecular CH₄ and other simple hydrocarbons would generate if trapped in microcracks.

The FEMS and FIMS observations provide yet another argument against organic contaminants in microcracks: when olivine crystals break which contain microcracks, the fracture will preferentially propagate along such cracks. Any "organic" compounds contained therein would become exposed and susceptible to evaporation/distillation. For homologous series such as the polyaromatic hydrocarbons the rate of evaporation will depend on the relative vapor pressures which decrease with increasing molecular weights. Therefore we should see a characteristic time dependence of the emission intensities after fracture and characteristic "thermal profiles".

Within the 80 msec time resolution achieved by FEMS in the scan mode we observe no correlation between the masses of the higher molecular weight fragments, in particular 105, 91, 77 etc. and their emission rates (see Figs. 2/3) suggesting that their respective parent molecules do not simply distill off the surface. The FIMS experiments provide additional support: the 302 amu compound, presumably dibenzopyrene, evolves below 200°C, while the 252 amu compound, presumably benzopyrene, comes between 280-380°C and pyrene (202 amu) peaks only at 380°C. Another sequence of molecules with 365-368 amu (not shown in Fig. 7) is released at still lower temperatures, around 150°C. Thus the thermal emission of the complex C-N-O-H compounds appears not to obey the simple law of distillation.

We therefore conclude that contamination by volcanic C-N-O-H fluids which penetrated microcracks and became catalytically reduced by some as yet unknown mechanism, can be reasonably ruled out as the source of the observed emission of "organic" molecules during and after fracture.

(iii) Next we consider contamination acquired at the surface of the earth, primarily biogenic contamination.

Obviously, the olivine crystals under study have been exposed for long time to environmental contaminants. Subgrain boundaries formed by dislocation arrays (Poirier 1985) are probably not wide enough for large biogenic molecules to enter. If biogenic contaminants diffused into microcracks and resided therein, their contribution to the observed gas emission can be ruled out on the basis of the same arguments as discussed above under (i) and (ii). Absence of biogenic contamination is also indicated by the paucity of the masses observed such as 302, 106/105, 92/91, 78/77 etc. Typical gas evolution and pyrolysis FIMS patterns of samples with known biogenic contaminants or kerogens tend to be much richer (Becker et al., 1986). Thus we conclude that environmental biogenic contamination is also an improbably source for the "organic" molecules emitted from the olivine crystals under study.
Fluid Inclusions as a Source for some Emitted Gases

We noted in our FEMS experiments that H\textsubscript{2}, H\textsubscript{2}O, CO/N\textsubscript{2} and CO\textsubscript{2} were generally released over relatively long time, often increasing over \approx 200 msec before decaying (see Fig. 2a). Such a behavior is contrary to what one would expect, if gases were released from gas/fluid inclusions dissected by the fracture plane. If gas-filled cavities are broken open, the gas volume becomes liberated at once, typically within \lt 10 \mu sec. The gases then freely expand and reach the ionizer which is located 12 mm away within \approx 10 \mu sec. The gas pulses will dissipate quickly, due to the large volume of the UHV system and to the powerful pumps. On the basis of experience gathered during the FEMS study of cloudy MgO crystals which contain minute cavities (Dickinson et al. 1987), pressure pulses generated by microscopic fluid inclusions will not last longer than \approx 30 msec.

The FEMS emission of H\textsubscript{2}, CO/N\textsubscript{2} and CO\textsubscript{2} for periods of \gt 100 msec point at a sustained gas supply from within. One possibility is that the gases seep out of a "leaky" reservoir of fluid inclusions connected to the fracture surface by diffusion channels. Though this is broadly in line with the fact that fluid inclusions are preferentially located on healed microcracks which may act as diffusion conduits (Roedder 1984), it also requires that the fluid inclusions lie just far enough beneath the surface to give the observed release kinetics. Our observations do not bear out the prevalence of such special conditions.

Dissolution Mechanisms of Fluid Phase Components

Next we address solid solution formation between C-N-O-H fluid phase components and olivine.

In the system MgO-SiO\textsubscript{2}-H\textsubscript{2}O to which olivine belongs by its Mg-rich end member forsterite, the occurrence of OH\textsuperscript{-} anions is not limited to regular OH\textsuperscript{-}-bearing phases such brucite, Mg(OH)\textsubscript{2}, serpentine, Mg\textsubscript{3}[(OH)\textsubscript{4}Si\textsubscript{2}O\textsubscript{5}], talc, Mg\textsubscript{2}[(OH)\textsubscript{2}Si\textsubscript{4}O\textsubscript{10}], etc. H\textsubscript{2}O also dissolves in the nominally anhydrous phases such as olivine. Together with an appropriate number of cation vacancies the OH\textsuperscript{-} anions become a part of the structure though their exact location and environment are still disputed (Beran and Putnis 1983; Freund and Oberheuser 1986; Miller et al. 1987).

CO\textsubscript{2} depresses silicate melting points and changes phase relations in multiphase systems in much the same way as H\textsubscript{2}O, though the pressures required are higher (e.g. Burnham 1979). The melting point depression is commonly explained by CO\textsubscript{2} dissolution in the liquid, either in form of CO\textsubscript{2} molecules or as carbonate anions, CO\textsubscript{3}\textsuperscript{2-} (Mysen and Virgo 1980; Bergman and Spera 1984; Fine and Stolper 1985). The idea that CO\textsubscript{2} may also form a solid solution with olivine first presented by Green and Radcliffe (1975) has been rejected by Mathez et al. (1984) and Mathez (1987).
This rejection cannot be upheld on the basis of fundamental rules of solid solution formation. Any olivine which crystallizes in the system (Mg,Fe)O- SiO2-CO2 must be non-stoichiometric and must contain at least a trace of structurally incorporated CO/CO2 component. Since carbon is a low-z element as a solute it is expected to form oxyanion complexes such as carbonate anions, CO3^2-. Other C-bearing oxyanions, however, cannot be excluded.

Nitrogen which is also a fluid phase component in the upper mantle is generally considered as chemically inert occurring only in the form of N2, unless NH4+ substitutes for K+ in micas (Hallam and Eugster 1976). Fundamentally, however, nitrogen must also be able to enter into solid solution as the ceramic system Si3N4-Si2N2O-Mg2SiO4 (Hendry et al. 1975; Lange 1979).

In other words, mantle-derived olivines may also contain dissolved nitrogen either as N2 or in the form of N-bearing oxyanions. Low levels of N have been reported in SiO2, possibly in form of NO2- oxyanions which would be isoelectronic with CO2^2- (Stathis and Kastner 1984). Knobel (1983) and Knobel et al. (1984) reported on N in synthetic MgO and natural olivine. Fusion of basaltic glass and extraction of nitrogen as N2 gave values in the range of 1 ppm (Becker and Clayton 1977). Exley et al. (1987) found 0.2 - 2.1 ppm N by fusion extraction. Total N analyses by neutron activation tend to give significantly higher values, 22 ppm (Norris and Schaeffer 1982).

Miller et al. (1987) determined the the OH^- content of olivines from different locations and found typical values of the order of 3-20 per 10^6 Si. Freund and Oberheuser (1986) have suggested that OH^- pairs in olivine may be converted to molecular H2 plus peroxy entities, O2^2- or Si OO\Si. This so-called charge transfer or CT reaction is a redox reaction which oxidizes oxygen to O^- while reducing protons to H2 (Freund 1985). It has been studied in MgO and CaO (Freund and Wengeler 1982; Freund et al. 1982) where the diagnostic H-H stretching band of molecular H2 is rather strong. Miller et al. (1987) report not to have found spectroscopic evidence for H2 and call the validity of the CT reaction in question.

The amount of CO2 component dissolved and the nature of the C-bearing species are poorly constrained. C concentrations in the range 10-500 ppm have been reported (Tingle and Green 1986; Oberheuser et al. 1983; Shilobreyeva et al. 1987). Based on studies of MgO and CaO (Freund and Wengeler 1982; Wengeler et al. 1982) it has been suggested (Oberheuser et al. 1983) that solute C in olivine occurs in the form of CO2^- anions associated with divalent cation vacancies and similar to the CO2^- and CO2^2- complexes in other matrices (Kafafi et al. 1984).

CO2 dissolution may proceed via the "orthocarbonate" anion, CO4^- which is theoretically predicted to be stable (Johnson and Wasson 1974) or by forming CO2^- and O2^2- via a CT reaction as in the case of dissolved H2O. Note that in CO2^- the carbon is reduced, while in O2^2- the oxygen is oxidized to the formal oxidation state -1.
Gas Evolution from a Supersaturated System.

The next question to be addressed concerns supersaturation. If solid solutions exist between olivine and the components of the C-N-O-H mantle fluid, each low-z element will be present at a specific concentration governed by the partial pressures ($P$) and temperature ($T$). Though the concentrations may not be known, the concentrations must decrease with decreasing $T$ and $P$, for instance during ascent of the rock to the surface of the Earth.

If equilibrium cannot be constantly adjusted, supersaturation ensues. Supersaturation means that the host mineral matrix will tend to exsolve the low-z "impurity" just like, for instance, Ca$^{2+}$ exsolves from MgO (McCune and Wynblatt 1983; Mackrodt 1987). Dislocations, subgrain and grain boundaries are always preferred impurity exsolution sites.

Naively the exsolution of C-N-O-H components may be thought of as a degassing process whereby the released gases are identical to those originally dissolved under mantle conditions, adjusted for the oxygen fugacity of the system. In other words, the gases released should consist of a mixture of H$_2$, H$_2$O, CH$_4$, CO, CO$_2$ and/or N$_2$. Indeed, under near-equilibrium conditions, for instance during a slow diapiric rise, these will be the gases exsolved.

However, when decompression and cooling are rapid such as during a volcanic ascent, we expect extreme non-equilibria. Under these conditions the C-N-O-H compounds which exsolve do not need to be the thermodynamically stable simple gases. This is particularly true, if we have reason to believe that a variety of solute species exist such as H$_2$ molecules plus C- and N-bearing oxyanions of different redox states. Then the situation may become very complex.

On a microscopic scale the exsolution will be controlled by the diffusion of the different solute entities. It suffices that one subset, for instance the reduced entities, has a higher average mobility. Then the gases or compounds which exsolve will be very different from the naively expected mixture of H$_2$, H$_2$O, CH$_4$, CO, CO$_2$ and/or N$_2$.

To further illustrate this point we note that CO$_2^{2-}$ can dissociate into CO$^{-}$ plus O$^{-}$. This dissociation has been postulated as the crucial step in the C diffusion mechanism which allows C atoms to enter interstitial sites and to become mobile. The degree of supersaturation of the solid solution can be reduced by exsolving only C and H$_2$, while retaining the oxidized entities, probably SiOO\text{Si}. Whether or not this happens is a question of relative mobilities, for instance the mobility of the C atom diffusing in conjunction with an O$^{-}$ state as a CO$^{-}$. At the exsolution sites, e.g. at dislocations, grain and subgrain boundaries, C-C bonds are predicted to form at the expense of the C-O$^{-}$ bonds. This eventually leads to the precipitation of C$_n$ (Freund 1986a/b).

$^{12}$C(d,p)$^{13}$C nuclear reaction and secondary ion mass spectrometric data have been interpreted to indicate that carbon is subject to surface segregation (Wengeler et al. 1982; Oberheuser et al. 1982).
1983; Freund 1986a), while Tsong et al. (1985) and Mathez et al. (1987) have expressed doubts whether carbon can exist in minerals in solid solution.

Tingle and Green (1986) attempted to measure the diffusion of $^{14}$C into an olivine single crystal at upper mantle pressure and temperature conditions, using $^{14}$C-doped graphite as a source for carbon. Though they observed C diffusion profiles, the carbon solubility in the bulk was less than expected, $\approx 100$ ppm, and apparently independent of P and T. Recrystallized olivine, on the other hand, had a much higher $^{14}$C concentration. The apparent difficulty to introduce $^{14}$C into an existing olivine single crystal is not in contradiction with the proposed CO$_2$ solid solution model which requires a readjustment of the overall stoichiometry (Freund 1986a/b). Such an adjustment, however, is possible when the olivine recrystallized and indeed reportedly incorporated much more $^{14}$C at the simulated upper mantle conditions (Tingle and Green 1986).

Since there is no consensus as to whether or not olivine can form a solid solution with CO/CO$_2$, the question is, whether the FEMS, SALI and FIMS experiments reported here can provide us with additional insight. If we assume that CO$_2$ dissolves by way of a CT reaction as proposed earlier (Freund 1986a), the FEMS, SALI and FIMS observations may provide a portion of the answer.

**C-N-O-H Precursors and C-N-O-H Molecules.**

The CT dissolution mechanism of CO$_2$ predicts two types of solutes, one reduced (carbon) and one oxidized (peroxy). If the C-bearing solute entities have a higher mobility, preferential C segregation occurs. If dislocations, subgrain and grain boundaries are sinks for segregating carbon, they are also the sites where polyatomic C$_n$ entities form through C-C bond formation at the expense of C-O$^-$ bonds. The same segregation arguments apply to H$_2$ and to N-bearing solute entities, meaning that the dislocations, subgrain and grain boundaries become enriched in H$_2$ and N with the ensuing possibility of C-H and C-N bond formation.

A conspicuous feature of the San Carlos olivine crystals used in this study is a dense dislocation network as part of an extended polygonized domain substructure (Kohlstedt and Goetze 1974; Poirier 1985). Assume that, deep in the mantle and prior to eruption, the olivine crystals contained carbon homogeneously dissolved at a concentration C$_o$. During decompression and cooling the solubility decreases and carbon will segregate. The segregant concentration C(x) at a distance x from such a segregation site is $C(x) = C_o \exp[-U(x)/kT]$ where U(x) is the interaction energy (Lücke and Stüwe 1971). C-C bond formation is predicted to take place at the segregation sites leading to polyatomic C$_n$ entities, plus additional C-H and C-N bonds, with n growing as segregation proceeds. A monoatomic segregation layer would correspond to a continuous carbon film with a few C-O, C-H and C-N bonds.
Another potentially important question is that the \( C_n \) entities thus formed are expected to be sterically constrained by the 2-dimensional structure of the subgrain boundary similar to an epitaxial overgrowth. \( C_6 \) rings, for instance, fit onto \( O^2^- \) anions like a crown. Several \( C_6 \) may fuse to polyaromatic structures and C-C sidechains may branch off in a specific way. Other shapes may be discriminated against by the structured environment. The result would be that the number of possible "organic" molecules synthesized by segregation from the solid state would be quite limited as compared to the total number of compounds known from liquid chemistry.

As long as the \( C_n \) entities are inside the crystal structure, even if their locus of formation is a defect such as a dislocation or subgrain boundary, they cannot be expected to be fully developed molecules. We rather should view them as precursor entities, still bonded to the mineral matrix via C-O\(^-\) bonds, possibly resembling the known cyclic \( CO_x^x\) oxocarbon anions, where \( 3 < x < 6 \) (Eggerding and West 1976; Staedeli et al. 1977). \( C_6 \) precursors on a subgrain boundary may resemble a rhodizonic anion, \( C_6O_6^2^- \), with \( n = 2 \) or \( 3 \), a benzene-like ring with six O substituting for H, formed through condensation of CO molecules on oxide surfaces with transfer of electrons from the oxide onto the adsorbate (Morris and Klabunde 1983).

If we look closely at our FEMS data, the gas emission may actually be controlled by the conversion of precursors sitting on or close to the fracture surface into mature molecules. The conversion can be done by scissoring C-O\(^-\) bonds and replacing them with C-H bonds. For instance benzaldehyde \( C_6H_5CHO \) which stands out in the FIMS experiments (as 106 amu molecular ion) and FEMS experiments (as 105 amu \( C_6H_5CO^+ \) fragment) strikes us as being a molecule which could have formed from a rhodizonite-like anion complex (with one remaining C-O\(^-\) sidechain) using a minimum of H to complete the molecular structure.

Another aspect which may be important for understanding the FEMS results relates to the extreme non-equilibrium condition at the fracture surface. This produces a strong driving force for further segregation from the near-by bulk. If C diffusion proceeds via the CO\(^-\) complex, the transport of carbon to the segregation site is coupled to the transport of a positive charge, e.g. O\(^-\). This causes the segregation site to become positively charged and to counteract further carbon segregation, even in a state of supersaturation (Freund 1986a/b). However, during fracture the charge is destroyed due to electron and ion emission during the fracture process. Diffusion of solute C from the near bulk would then be driven by the extreme gradient of the electrochemical potential bringing additional C and H to the fracture surface.

Heating further activates diffusion and, hence, enables segregation. Heating also enhances reactions between segregants and at the same time allows larger molecules to desorb.

This brings us back to the FIMS question (see Fig. 7): Why do the heavy 302 amu molecules (probably benzopyrelene or dibenzopyrene, \( C_{24}H_{14} \)) evolve at a distinctly lower temperature than a number of lighter molecules? Maybe six asymmetrically arranged condensed \( C_6 \) rings form a sterically favorable precursor entity which becomes the 302 amu compound. Other smaller entities
require additional segregation or more H relative to the number of C to convert into desorbable molecules. Obviously, at sufficiently high temperatures pyrolysis sets in with benzene $\text{C}_6\text{H}_6$ as the main pyrolysis product besides toluene $\text{C}_6\text{H}_5\text{CH}_3$. As shown in Fig. 7 benzene (78 amu) and toluene (92 amu) rise late in the heating program, suggesting that they are formed by pyrolysis.

We may also draw attention to an interesting similarity between the mass spectra as reported in this paper and some molecular ion spectra detected in ionopause of Comet Halley by the Giotto spacecraft (Mitchell et al. 1987). The ions at 63 and 65 recur with high intensities in both data sets. Outside the Halley ionopause low resolution mass spectra reveal a series of peaks centered at 75, 90 and 105 amu which Heubner (1987) proposed to be due to dissociation products of polyoxymethylenes, derivatives of aldehydes.

Maybe the complex organic molecules from Halley are formed by reactions similar to the ones discussed above, e.g. through segregation of solute C and other low-z elements from silicate grains which are constituents of the comet. The appearance of these molecular ions in the Halley ionopause suggests thermal and radiation-induced desorption from dust grains ejected from the Comet's surface, followed by UV ionization.

**The Role of the $\text{O}^-$ States.**

We need to address another puzzling aspect of the FEMS experiments: why do the olivine crystals simultaneously emit oxidized gases (CO$_2$, H$_2$O, possibly even O$_2$) and reduced gases ("organics") plus metal vapor upon fracture? FEMS experiments on MgO have shown that, following fracture, reduced "organic" molecules and Mg vapor are emitted at the same time as CO$_2$, H$_2$O and even O$_2$ (Dickinson et al. 1986). The answer appears to lie in the presence of peroxy entities in the mineral structure. Such peroxy entities are the counterparts of the reduced entities and probably of the form Si$^{\text{OOOSi}}$ (Freund and Oberheuser 1986).

Peroxy entities dissociate by breaking the O-O bond and releasing an $\text{O}^-$ state: $\text{O}_2^{2-} = \text{O}^- + \text{O}^-$. An $\text{O}^-$ is an electronic defect in the valence band, as radical state, commonly called a positive hole (Henderson and Wertz 1977; Marfunin 1979; King and Freund, 1984). $\text{O}^-$ states have several properties which are important in the context of the present study. (i) They move without mass transport through the mineral structure. (ii) They repel each other in the bulk and diffuse to the surface. (iii) They are powerful oxidizing reagents disproportionating into free oxygen: $\text{O}^- + \text{O}^- = \text{O}_2^- + 1/2 \text{O}_2$.

In some FEMS experiments the 1(H$^+$), 2 (H$_2^+$), and 17 (OH$^+$) amu intensities as a function of time indicate a negative correlation between the emission of H$_2$ and H$_2$O. A possible interpretation is that H$_2$O molecules are formed through oxidation of H$_2$ molecules by $\text{O}^-$ states, if such $\text{O}^-$ states flood the surface and scavenge the H$_2$ molecules before they desorb. With <100 µsec time
resolution, Dickinson et al. (1987) observed erratic variations of the O$_2$ emission from MgO crystals over periods of ~300 msec after fracture. They suggested that the oxidizing and reducing character of the fracture surface fluctuates due to rapid variations of the local O$^-$ density.

In the case of the olivine fracture we have not yet carried out similar experiments at higher time resolution. However, the similarity of the olivine data to the MgO results is striking. If the MgO observations also apply to the olivine fracture, areas on the olivine fracture surface with a high O$^-$ density would momentarily become oxidizing, generating H$_2$O, CO$_2$ and possibly even O$_2$, while other areas remain reducing and allow "organic" molecules to evolve.

**Metal Vapor Emission.**

If H$_2$O, CO$_2$ and other O-bearing molecules are formed through oxidation by O$^-$, the oxygen consumed in such reactions must be supplied the fracture surface. The reason is that O$^-$ states are just electronic defects in the O$_2$ sublattice traveling without mass transport via the valence band (King and Freund 1984). At the surface the O$^-$ can oxidize the solute entities according to the schemes:

\[
\begin{align*}
CO^- + O^- & = CO + O^2- \\
CO_2^{2-} + 2 O^- & = CO_2 + 2 O^2- \\
H_2 + 2 O^- & = H_2O + O^2-
\end{align*}
\]

If oxygen is lost from the surface, coordinatively unsaturated cations remain which in turn may be the source of the observed Mg vapor emission: Mg$^{2+} + 2 O^- = Mg^0 + O_2$.

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Legends

Fig. 1  Schematic view of the fracture device used in the FEMS experiments.

Fig. 2  Five consecutive 80 msec FEMS scans of a San Carlos olivine crystal. Scan (1): residual gas spectrum; scan (2)-(5): the next 320 msec including the fracture near 60 amu in scan (2). Besides H$_2$/H$_2$O, CO/CO$_2$ a number of complex H-C-N-O molecules are emitted.

Fig. 3  Similar data as in Fig. 3, plotted on a logarithmic scale.

Fig. 4  Typical SALI mass spectrum from the gases thermally released from an olivine fragment after cleavage in air and introduction into UHV.

Fig. 5a/d  "Thermal profiles" of gases released during heating in the SALI instrument.

Fig. 6  Gases emitted from crushed olivine single crystal powder sample in the FIMS experiment, accumulated over the temperature range up to 500°C.

Fig. 7  Intensities of selected mass peaks in the FIMS experiment. Note the high intensity of the 302 peak which reaches its maximum at a lower temperature than several lower molecular weight compounds.

Fig. 8  Infrared absorption spectrum of a San Carlos olivine crystal of the type used in this study. Note the near absence of any O-H stretching bands and the occurrence of a broad double band in the region of the C-H stretching absorption.
Fig. 1
SAN CARLOS OLIVINE IMPACT FRACTURE MS

H₂, H₂O, CO/N₂, CO₂, RESIDUAL GAS

INTENSITY, arb. units

MASS, amu

Fig. 2
San Carlos Olivine

Impact Fracture
Mass Spectroscopy

Fig. 3
Fig. 4
Fig. 5
Fig. 6
FIMS
(Field Ionization MS)
San Carlos Olivine

Fig. 7
Freund et al.: Gases Released from Olivine by Impact Fracture.
Electrical Conductivity Measurements under Minimum Perturbation Conditions

Part I:
Charge Distribution Analysis of Magnesium Oxide

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Abstract

A new technique, Charge Distribution Analysis, has been developed which measures electric and dielectric properties contact-free at the limiting case of 0 Hz under minimum perturbation conditions. CDA allows to determine surface charges and their associated internal electric fields. This unique feature and a high sensitivity make CDA particularly suitable to study electron and hole conductivity in otherwise highly insulating minerals. Positive charge carriers have been identified in nominally high purity MgO single crystals between 200-400°C where the electrical conductivity is in the $10^{-12}-10^{-14}$ S/m range, and between 500-800°C where it rises to $10^{-4}-10^{-5}$ S/m. The charge carriers reversibly appear and disappear during repetitive heating/cooling cycles. Because of their pronounced tendency to diffuse to the surface and cause a high surface conductivity, these charge carriers have been misinterpreted as sign of contamination. The internal electric field associated with the surface charge indicates that the charges are not due to contamination but are generated in the bulk and that their diffusion to the surface is an internally controlled process.

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Introduction

Most minerals are insulators. If they carry an electric current, they do so because ions become mobile at elevated temperatures or because they contain electronic charge carriers that can move in response to an externally applied field. Ions hop from lattice site to lattice site either interstitially or via a vacancy mechanism. Electronic charge carriers may also be visualized as hopping from site to site or delocalize via orbital overlap. The important physical information that can be gained from electrical conductivity measurements pertains to the nature of the charge carriers and to the way they conduct current by moving through the mineral structure or through a rock at given temperatures, pressures and gas or fluid environments.

Olivine, the major mineral of upper mantle rocks, has been studied extensively. Most conductivity measurements have been carried out under "simulated" upper mantle conditions [Constable and Duba, 1990; Schock and Duba, 1985; Schock et al., 1977; Schock, Duba and Shankland, 1989; Sockel, 1974], above 1000°C at ambient pressure (0.1 MPa) in H₂:CO₂ or CO:CO₂ gas mixtures for controlling the oxygen fugacity, f(O₂) [Ballhaus et al., 1990]. Measurements under high pressures have also been performed [Bradley et al. 1964; Shankland 1969; Duba 1972; Duba and Nicholls 1973; Duba et al. 1974; Omura et al., 1989] though not in the presence of the H–C–O fluids believed to exist in the upper mantle. The reason for not using high pressure H–C–O fluids is that these fluids tend to have a high electrolytic conductivity which drowns the typically much smaller solid state conductivity of the minerals to be studied. On the other hand H–C–O gases and fluids are known to be important components of the earth's interior and it may make a difference whether they are present at 0.1 MPa or at realistic upper mantle pressures. When fluid phase components dissolve in the minerals – even on a trace level – they give rise to defects and impurities which may profoundly affect the electrical conductivity. H₂O dissolves in olivine, forming OH⁻ associated with different defect sites [Beran and Putnis, 1983; Freund and Oberheuser, 1986; Miller et al., 1987; Wilkins and Sabine, 1973]. The ubiquitous occurrence of OH⁻ in olivine has led to a recent surge of interest in the possibility that hydrogen – more exactly protons – may act as charge carriers and make a significant contribution to the conductivity of the upper mantle [Hirsch, 1990; Karato, 1990].

Against the background of decades of intense study one may be inclined to think that all potentially important charge carriers and conduction mechanisms in olivine should have been identified. However, there is one peculiarity in the electric response of olivine at 0.1 MPa for which no compelling explanation has been offered so far: the enhancement of the conductivity by many orders of magnitude whenever mantle-derived olivine crystals are heated in the laboratory for the first time. The anomaly is restricted to relatively low temperatures – below about 700°C – and generally disappears while the samples are slowly heated in the reducing H₂:CO₂ or CO:CO₂ gas mixtures. The cause of this conductivity peculiarity is unknown, but since it disappears readily between 700–800°C it has not raised much concern. Most researchers have been satisfied
with the assumption that the observed conductivity anomaly of olivine is a laboratory-induced artefact, a "dirt effect" which does not need to be taken seriously.

In this study we examine the validity of this assumption. Recognizing that olivine crystals from the upper mantle are enormously complex—crystallographically and with respect to defects and impurities—we first turn to a model system with less variables. A suitable model system is MgO crystallizing in the simple face-centered cubic structure. Synthetic MgO crystals of nominally highest purity grade are available. As we shall show below they exhibit a strikingly similar electrical conductivity anomaly as mantle-derived olivine crystals. In Part I we describe a novel technique, Charge Distribution Analysis (CDA) which allows us to obtain information about the charge carriers that is unavailable through conventional conductivity measurements [Freund et al., 1989]. In Part II we show how supplementary physical techniques can help us to identify the nature of the charge carriers responsible for the conductivity anomaly. Finally, in Part III, using the knowledge gained by studying MgO we shall apply CDA to mantle-derived olivine crystals and draw conclusions about their charge carriers.

Identification of the Problem

Electrical conductivity measurements serve to detect charge carriers that can move in response to an externally applied electric field. The conductivity $\sigma$ is usually given in units of reciprocal Ohm-centimeter, $\Omega^{-1}\text{cm}^{-1}$, or Siemens per meter, S/m. It can be written as sum of contributions from charge $i,j$: $\sigma_{i,j} = \sum_i \sigma_i \exp \left[-E/kT\right]$, where $\sigma_i$, $E$, $k$ and $T$ are, respectively, pre-exponential factors, activation energies, the Boltzmann constant and the absolute temperature. Conductivity data are conveniently presented as Arrhenius plots, e.g. $\log \sigma$ vs $1/T$.

In predominantly ionic solids all charge carriers fall into two categories: ionic and electronic. Ions commonly move via a vacancy hopping mechanism or interstitially. Their diffusion involves mass transport. In refractory oxides and silicates ionic diffusion is generally restricted to high temperatures, typically $>800^\circ\text{C}$. Electronic charge carriers can be electrons or holes. Neither involves mass transport.

Electrons are negative charges. Their movement may be visualized as localized hopping or as due to orbital overlap between nearby ions with partially filled shells such as transition metal cations which results in delocalized states. Holes are defect electrons with respect to the perfect structure. They are positive charges, often symbolized by $h^\circ$. Their diffusion can take place either on the cation or the anion sublattice or on a combination of both. Holes are commonly associated with transition metal cations undergoing transient oxidation, for instance $\text{Fe}^{2+} \leftrightarrow \text{Fe}^{3+} + h^\circ$. However, holes do not need to be confined to the cation sublattice. They can also reside on and move via the $\text{O}^{2-}$ sublattice. In this case they are called "positive holes" or "oxygen associated holes" and represent $\text{O}^-$ radicals [Griscom, 1990; Henderson and Wert, 1977]. Such $\text{O}^-$ radicals have been identified in many natural minerals exposed to ionizing radiation [Marfunin, 1979].
Radiation-induced O\(^-\) irreversibly disappear upon annealing. Questions which need clarification are (i) what role do such O\(^-\) play in the electrical conductivity at higher temperatures, and (ii) where in the mineral matrix are the O\(^-\) located or stored when they are not, or not yet, mobile charge carriers?

Electrical conductivity measurements of minerals, either with direct currents (d.c.) or with alternate currents (a.c.), are commonly carried out with metal electrodes in direct contact with the samples. The electrodes are deposited by evaporation or pasted onto the sample surfaces or mechanically pressed against them to make the best possible contacts. Unfortunately metal-to-insulator contacts are also sources of concern. To detect the charge carriers which move through the sample they have to be converted into electrons at the interfaces. If the carriers are electrons or holes, their Fermi levels have to adjust across the metal-to-insulator contacts. If they are ions, the conversion to electrons needed to drive the external measuring circuit requires electrochemical reactions at the interface. In all cases the contact potentials may become perturbed by the accumulation of space charges or by chemical reactions in the near-electrode region. In extreme cases the flow of charge carriers can be totally interrupted due to electrode passivation. Another concern with conventional techniques relates to surface and grain boundary conductivity. Surfaces, grain boundaries and subgrain boundaries represent extended 2-dimensional defects where intrinsic valence and conduction bands are shifted on the energy scale, where additional defect levels are introduced and activation energies for many diffusional processes lowered. Lastly there is the possibility of surface contamination during sample handling or in the course of measurements.

Fig. 1 illustrates an Arrhenius plot, log \(\sigma\) vs 1/T [K], for olivine and olivine-rich rocks compiled from data reported by Duba and coworkers [Constable and Duba, 1990; Duba et al., 1974; Shankland and Duba, 1990]. The high temperature (HT) branch is represented by open and solid triangles and small solid squares. In the HT region the conductivity is fairly reproducible for given oxygen fugacities and is caused by cation diffusion [Buening and Busek, 1973; Morin et al., 1977a,b; Smyth and Stocker, 1975], possibly enhanced by hole diffusion via the cation sublattice, transiently trapped on Fe\(^{2+}\) sites to generate Fe\(^{3+}\) [Schock et al., 1989]. For synthetic forsterite a mixed HT mechanism has been proposed combining predominantly electronic conduction in the crystallographic a and b directions and cation diffusion via vacancy hopping in the c direction [Schock et al., 1989]. Upon cooling the HT branch never extends very far below 800°C, but changes more or less abruptly into a low temperature (LT) extension with a distinctly smaller activation energy. The conduction mechanism in the LT region is seldom discussed. Two LT extensions are depicted in Fig. 1 by the small solid and open squares.

The anomalously high conductivity which is typically observed below approximately 700°C during initial heating is delineated in Fig. 1 by large open squares. In the example selected here a fresh sample was heated in a 1:1 CO/CO\(_2\) mixture at 0.1 MPa pressure at 1°C/min [Constable and Duba, 1990]. Up to about 700°C the conductivity was many orders of magnitude higher than the downwardly extrapolated HT branch and several orders of magnitude above any of the LT
extensions measured during cooling. The anomaly ended around 700°C when the conductivity decreased precipitously, joining the HT branch above 800°C.

![Graph of electrical conductivity vs. temperature](image)

**Fig. 1** Electrical conductivity of olivine single crystal and dunite samples under simulated upper mantle conditions (P = 1 bar) compiled from data by Duba and coworkers [Constable and Duba, 1990; Duba et al., 1974; Shankland and Duba, 1990]. During initial heating an anomalously high conductivity is observed, variously assigned to “dirt” on the sample surface or to amorphous carbon precipitating out of the CO:CO₂ gas mixtures at low temperatures in the stability field of graphite.
Fig. 2 Typical electrical conductivity behavior of high purity, arc-fusion grown MgO single crystals heated in high purity Ar. The data represented by open squares are adopted from Kathrein and Freund (1983). The small maximum below 400°C (1000/T > 1.5) was confirmed by Kathrein et al. (1980). The steep section of the solid line delineates the high temperature Arrhenius branch as reported by many authors including Mitoff (1959, 1962, 1964), Lewis and Wright (1970) and Osburn and Vest (1971). The less steep section and dashed extension is adopted from graphical representations given by Lewis and Wright (1970) and from work by Kathrein and Freund (1983).

Surface contamination or structural disorder within a surface layer of unspecified thickness have long been thought to cause the conductivity anomaly in olivine [Duba, 1972; Duba et al., 1974; Duba et al., 1988; Duba and Shankland, 1982] and pyroxene [Duba et al., 1973]. Recently, Constable and Duba (1990) suggested that CO₂ undergoes disproportionation in the stability field of graphite, CO₂ = C + O₂, precipitating a layer of amorphous carbon onto the sample surface. Assuming that thermodynamic equilibrium can be achieved around 700°C the precipitated carbon
layer would react with gas phase CO₂ as soon as the CO/CO₂ mixture leaves the stability field of graphite and disappear again, revealing the intrinsic conductivity of the olivine sample under study. The fact that a 1:1 CO/CO₂ mixture at 0.1 MPa leaves the stability field of graphite around 670°C [Deines, et al., 1974] the end of the conductivity anomaly around this temperature (see Fig. 1) is quoted as supporting evidence for the gas phase CO₂ disproportionation reaction [Constable and Duba, 1990]. If this were true, the same conductive carbon layer should form on any substrate heated in a 1:1 CO/CO₂ mixture to any temperature below 670°C. Conversely, no carbon layer and, hence, no electrical conductivity anomaly, should be observed in any CO/CO₂-free gas or gas mixture.

Fig. 2 shows the d.c. conductivity of synthetic MgO single crystals measured in pure Ar at 0.1 MPa pressure, e.g. in a CO/CO₂-free atmosphere. The MgO crystals were of the nominally highest purity grades, 4N or 99.99% with <50 ppm transition metal impurities. They were grown by the carbon arc fusion technique [Butler et al., 1971]. The bold line in Fig. 2 depicts the HT branch with its high activation energy, 2.4 eV, as reported by many authors [Osburn and Vest, 1971; Sempolinski and Kingery, 1980; Sempolinski et al., 1980] and a typical LT extension, characterized by a lower activation energy, ~1 eV. As in the case of olivine, the LT branch is observed during cooling whenever the sample has undergone extensive heating and equilibration at higher temperatures. The open squares represent a conductivity anomaly that closely resembles that of olivine for which Constable and Duba [Constable and Duba, 1990] invoked carbon precipitation from the gas phase. When this anomaly was studied in greater detail [Kathrein and Freund, 1983], it became clear that it was reversible with a characteristic hysteresis behavior. It could be shown that there in fact are two distinct temperature regions where the conductivity is anomalously high, 300-400°C where the conductivity is in the 10⁻¹²-10⁻¹³ S/m range [Kathrein et al., 1980] and 500-800°C. Application of guard electrodes is not fully effective, indicating that the charge carriers may not be confined to the outermost surface [Kathrein and Freund, 1983]. Both conductivity anomalies are reversible. In the experiment depicted in Fig. 2 the heating of the MgO crystal (previously subjected to several temperature cycles up to 800°C) was interrupted by three cooling and reheating cycles at 10°C/min. Three branches were obtained with the same slope as the LT extension from the HT region. Since this measurement was conducted in Ar, the anomaly cannot be blamed on carbon precipitation from a CO₂-containing gas phase.

The similarity between the anomalous conductivity of olivine in CO/CO₂ mixtures and of MgO in pure Ar gas leads us to wonder whether this behavior can really be caused by extraneous surface contamination or by carbon that has precipitated from the gas phase. Maybe both, mantle-derived olivine and synthetic MgO crystals, contain some indigenous charge carriers which are responsible for this peculiar anomaly but which have not yet been correctly identified.

Conventional techniques use electrodes in direct contact with the samples. However, metal-to-insulator contacts often lead to serious perturbations at the interfaces. In addition, conventional techniques are notoriously inept to differentiate between surface and bulk conductivities. We have recently reported on a new non-contacting technique to measure mobile charge carriers in highly
insulating samples under minimum perturbation conditions [Freund et al., 1990, 1989]. We shall
discuss this new technique in the next chapter and illustrate its application with data of nominally
high purity MgO single crystals. Application to natural olivine single crystals and the discussion
of possible geophysical implications will be deferred to Part III.

A. Charge Distribution Analysis

In essence, conductivity measurements determine whether or not charge carriers exist in a given
sample that can diffuse in an externally applied electric field. Conventionally these measurements
are carried out with electrodes in direct contact with the samples. However, the same information
can be obtained contact–free by measuring the dielectric response [Freund, 1990].

When a dielectric is placed in an electric field $E_{\text{ext}}$ it polarizes. Its polarization $P$ is given as:

$$ P = \frac{\varepsilon_0 (\varepsilon - 1)}{4\pi} E_{\text{ext}} $$

where $\varepsilon_0$ is the permittivity of vacuum and $\varepsilon$ the dielectric constant. The polarization depends on
the frequency at which it is measured and the term “dielectric constant” has been applied to the
entire range from optical frequencies, $10^{14}–10^{16}$ Hz, to 0 Hz [Kittel, 1980]. The so–called
“static” dielectric constant is usually measured at $10^3–10^6$ Hz and extrapolated to 0 Hz.

In a homogeneous electric field the dielectric experiences no net acceleration. However, when
placed in an electric field gradient along a direction $z$, a force $F_z$ acts on the sample which draws it
towards the region of higher field density. In the case of an ideal dielectric the force is
independent of the polarity: $F_z^+ = F_z^+ (E_{\text{ext}}) = F_z^- (-E_{\text{ext}})$. Using Maxwell’s equations $F_z$ is given by:

$$ F_z = \int_V \nabla (P \cdot E_{\text{ext}}) dV $$

where the volume integral must include the sample but not the sources of the field.

In the case of a real dielectric, e.g. a material containing various defects and impurities, this force
is generally not independent of the polarity, $F_z^+ (E_{\text{ext}}) \neq F_z^- (-E_{\text{ext}})$. For the purpose of the
present paper we break the polarization down into five contributions, $P_{\text{el}} + P_{\text{ion}} + P_{\text{local}} + P_{\text{space}}$
$+ P_{\text{surface}}$. The first two contributions refer to the ideal structure [Ashcroft and Mermin, 1976].
$P_{\text{el}}$ represents the electronic polarization of the electronic shells of atoms and ions which can
follow fields that oscillate at frequencies up to $10^{14}–10^{16}$ Hz. $P_{\text{ion}}$ represents the ionic
polarization due to the field–induced displacement of atoms and ions which have typical response
times of the order of $10^{-9}–10^{-12}$ sec. The last three contributions arise from defects or impurities
and generally have longer relaxation times. $P_{\text{local}}$ contains contributions from local dipoles which
may rotate in the electric field but not diffuse over macroscopic distances. $P_{\text{space}}$, the space
charge polarization, is caused by charges which can diffuse over macroscopic distances. Space
charge polarization can also occur in the absence of externally applied electric fields. Surface charges occur because (i) the free energies of formation of cation and anion vacancies in ionic crystals in the bulk differ from those near the surface and thus create an imbalance [Kliewer and Koehler, 1965; Lehovec, 1953], (ii) the energy levels in the valence and conduction bands as well as in impurity levels are shifted [Kittel, 1980], and (iii) if charge carriers of one sign have a significantly higher mobility than the compensating charge carriers of the opposite sign, a charge layer forms at the surface together with a subsurface depletion zone [Sze, 1969]. We combine (i) to (iii) into the $P_{\text{surface}}$ term and refer to it summarily as the "surface charge".

$P_{\text{el}}$ and $P_{\text{ion}}$ weakly dependent upon temperature (T) [Thorpe et al., 1986]. $P_{\text{local}}$ may be T-dependent to the extent that local defects with a dipole moment are subject to thermally activated reorientation in the applied electric field. Their dipole moments, $\mu = d_e$, are small because $d$, the separation between the charges $e$, is of the order of interatomic distances, $10^{-8} \text{ cm}$. Hence, $P_{\text{local}}$ is not expected to make a large contribution. By contrast, $P_{\text{space}}$ can be large, if charges diffuse over large distances. At the limit $d$ may be the size of the crystal in field direction, say $10^{-1} \text{ cm}$, and dipole moments stemming from diffusing mobile charges may make a contribution $= 10^7$ times larger than that from rotating local dipoles.

$P_{\text{el}}, P_{\text{ion}}, P_{\text{local}},$ and $P_{\text{space}}$ are invariant to the reversal of the field gradient. By contrast $P_{\text{surface}}$ contributions are variant to field gradient reversal. This means that a dielectric with a surface charge will be either attracted to or repelled from the region of higher electric field density, depending on the sign of the charge and the polarity. Assuming that bulk and surface terms are independent of each other we can separate field-invariant and field-variant contributions by forming the linear combinations:

$$ F_\Sigma = \frac{1}{2} (F^- + F^+) = \frac{\varepsilon_0 (e-1)}{4 \pi} \int V [(P_{\text{el}} + P_{\text{ion}} + P_{\text{local}} + P_{\text{space}})] E_{\text{ext}} dV \quad (3) $$

$$ F_\Delta = \frac{1}{2} (F^- - F^+) = \int V (P_{\text{surface}} \cdot E_{\text{ext}}) dV \quad (4) $$

Eqs. (3) and (4) reveal an important feature of the CDA technique: its capacity to determine the sign of a surface charge and, hence, the sign of the majority carriers. This information is normally not available through conductivity measurements, but only through Seebeck coefficient (thermopotential) and Hall measurements [Ashcroft and Mermin, 1976; Kittel, 1980; Sze, 1969].

To evaluate $F_\Sigma$ and $F_\Delta$ we subtract $a_+ U^2$ from $F_\Sigma$ where $U$ is the voltage and $a_\pm$ are constants:

$$ F_{\pm} = F_{\pm 0} + a_\pm U^2 \quad (5) $$

The $a_\pm U^2$ term with $a_- = a_+$ describes the ideal dielectric, while $F_{\pm 0}$ arises from the charge at the surface which acts as an electrostatic mirror. With this approximation $F_\Sigma$ and $F_\Delta$ become:

$$ F_\Sigma = \frac{1}{2} (F^-_0 + F^+_0) + \frac{1}{2} (a_- + a_+) U^2 \quad (6) $$

$$ F_\Delta = \frac{1}{2} (F^-_0 - F^+_0) + \frac{1}{2} (a_- - a_+) U^2 \quad (7) $$
At low voltages we can set $F^0 = F_0^+ + \delta F$, and $a_- = a_+ + \delta a$, where $\delta F \ll F_0^\pm$ and $\delta a \ll a_\pm$. This corresponds to a small perturbation of the $\mathbf{P}$ surface contribution representing the electrostatic mirror. At high voltages the $F^\pm_0$ terms become negligible and $a_\pm = 1/2 (a_+ + a_-)$. We can thus define $F_\Sigma$ and $F_\Delta$ through their high and low voltage limits, respectively:

$$F_\Sigma = \frac{1}{2} (a_- + a_+) U^2 \equiv a U^2 \quad \text{for high U} \quad (8)$$

$$F_\Delta = \frac{1}{2} (F_0^- - F_0^+) \quad \text{for low U} \quad (9)$$

**Experimental**

For the CDA measurements a PERKIN-ELMER Thermogravimetric Analyzer TGS-2 was modified by the introduction of a circular Au bias electrode (3 mm diameter) at the bottom of the cup-shaped Pt furnace (10 mm internal diameter, 18 mm height), and of an annular Au counterelectrode (12 mm high) on the inside wall of the furnace reaching from its upper rim downwards. Thus the electric field lines were symmetrical to the cylinder axis, emerging vertically from the bias electrode at the bottom fanning out upwards and ending horizontally in the counterelectrode. The bias electrode and counterelectrode were connected to the ± output and ground respectively, of a KEITHLEY 240A digital voltage source (±1100V). The Pt windings of the furnace were bifilar and a.c. operated using one half period of the sine wave for power and the other for measuring the resistance and, hence, the temperature. The sample rested at four points on a Pt stirrup (0.1 wire, 6 mm diameter loop) suspended from a fused silica fiber, well insulated from ground. The Pt stirrup gave no contribution to the CDA measurements except for a constant off–set that was independent of temperature. The distance between the lower sample surface and the bias electrode was chosen at different values, generally at 150 ± 20 µm. In its most sensitive range the balance can detect forces equivalent to an apparent weight change of $<10^{-7}$ g.

The samples were nominally high purity MgO single crystals, grown from a CO/CO$_2$–saturated melt in a carbon arc fusion furnace [Butler et al., 1971] by W. & C. Spicer, Ltd., England. The crystals were rated 4N with less than 100 ppm cationic impurities, typically =10 ppm of Cr, Mn, and Ni, <30 ppm Fe, and <50 ppm (Al + Si). In addition they contain OH$^-$ in the range of 10–100 at.–ppm, due to the presence of adsorbed H$_2$O in the MgO powder starting material [Chen, et al., 1975; Freund and Wengeler, 1982]. The crystals came as large (3-10 cm$^3$) cubes or “as grown” thumb–sized forms bordered by pitted, irregular growth surfaces. The crystals were slightly yellowish–brown, due to color centers not related to transition metal impurities. Except for a clear rim of 2–4 mm thickness adjacent to the growth surfaces, the crystals were typically cloudy in the center, due to a network of curved subgrain boundaries decorated by µm–sized cavities. For the CDA experiments pieces (approximately 6 x 6 x 2 mm) were obtained by cleavage. To approximate a cylindrical shape the four edges were rounded on SiC sanding paper. The crystals were cleaned in acetone and methanol, dried in air, and subjected at least one in situ heating and cooling cycle up to 800–900°C.
All measurements were performed in high purity, dry Ar at a flow rate of about 10 ml/min with no additional control of the oxygen fugacity. Heating between temperatures (25 or 50°C increments) was at 20°C/min. After reaching the maximum temperature the crystals were cooled to 25°C or 100–150°C at 1°C/min. During heating the samples were allowed to thermally equilibrate at each step prior to the application of the bias voltages. An equilibrium time of about 5 min was generally found sufficient. The positive and negative bias was applied in 2, 5, 10, 20, 50 or 100 V increments, depending upon the experiment. After each sequence the dielectric polarization of the sample was allowed to relax under 0 V bias for several minutes while the temperature was raised to the next step. At some temperatures “creeping” occurred, e.g. the dielectric polarization increased over several minutes to an hour. The creeping was not due to slow thermal equilibration of the sample under study but to the appearance of charges which diffused in the electric field gradient. Creeping under positive bias was, as a rule, slower than creeping under negative bias at the same temperature, indicating that the diffusion of charge carriers within the sample was not symmetrical. Whenever the creeping rates differed markedly under positive and negative bias, a surface charge developed, indicating that the asymmetry in the charge carrier diffusion process is caused by the filling of surface states. Occasionally, to test the reproducibility, the sequence of positive and negative bias application was reversed. Except for the temperature intervals marked by “creeping” and except for cases where insufficient time was given for relaxation of the sample under 0 V bias, the reproducibility was typically better than ±10%, independent of whether the sample was first polarized under positive or under negative bias.
At ambient temperature, the MgO crystal responded as a nearly ideal dielectric material: immediately after application of the field gradient, a force was measured that remained constant with time. The force was proportional to the square of the applied voltage, $F^+ \propto U^2$, as shown in Fig. 3, giving a near-perfect straight line over the range 0–700 V. Its slope is proportional to the dielectric susceptibility $(\varepsilon - 1)$ where $\varepsilon$ is the dielectric constant, 9.6 for MgO [Gourley and Runciman, 1973]. At low voltages, a small deviation is noted (inset in Fig. 3), indicating an
intersect with the ordinate at about -0.75 force units. This repulsion under positive bias is due to a weak $P_{\text{surface}}$ term, suggesting a weak positive surface charge on the MgO crystal under study.

Fig. 4a  $F^+$ measured as a function of time at +20 V between 200-585°C. Note "creeping" above 300°C.

Normally one would not expect any drastic change of the dielectric polarization with temperature ($T$) because the $T$ dependence of the $P_{\text{el}}$ and $P_{\text{ion}}$ terms, characteristic of any ideal dielectric, is of the order of the volume thermal expansion coefficient. This expectation, however, is not fulfilled at $T \geq 300^\circ\text{C}$ as illustrated by Fig. 4. In (a) we show that $F^+$, measured as a function of time ($t$) at +20 V, remains nearly constant below 282°C. At 282°C and more clearly at 307°C $F^+(t)$ increases slightly during the first 200 sec. At 332°C the "creeping" extends over at least 900 sec, becoming progressively faster with increasing $T$ and reaching a steady state at 459°C after about 300 sec, about 5 times larger than $F^+$ at 200°C. At higher $T$ the kinetics further change as shown in (b). At 484°C and 510°C the creeping is slow lasting for over 1500 sec. At 534°C a steady
state is reached within $\approx 600$ sec. This time shortens to $< 90$ sec at $585^\circ$C. At $585^\circ$C, the steady state $F^+$ value is $\approx 15$ times larger than $F^+$ around $200^\circ$C. This time-dependent increase in the dielectric response suggests the appearance of mobile charge carriers in the MgO crystal under study.

![Graph showing $F^+$ measured as a function of time at $+20$ V between $200-585^\circ$C and up to $585^\circ$C.](image)

**Fig. 4a/b** $F^+$ measured as a function of time at $+20$ V (a) between $200-585^\circ$C and (b) up to $585^\circ$C. Note the different ordinate scales and the extended “creeping” above $300^\circ$C.

In order to study the surface charge contribution, $P_{\text{surface}}$, we measured $F^+$ and $F^-$ under otherwise identical conditions but at higher bias voltages of $\pm 50$ V. The higher bias voltage increases the electric field gradient and thus not only increases $F^+$ but also increases the charge carrier diffusion. In **Fig. 5** we show the result of the measurements below $425^\circ$C when the d.c. conductivity of the same purity-grade MgO crystal (**Fig. 2**) displayed a shallow maximum*. The CDA measurements were carried out at follows: The crystal was first heated in Ar to $800^\circ$C, then

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* In this and the next figures $F_\pm$, $F_\Sigma$ and/or $F_\Delta$ are plotted versus $T$ [°C], not $1/T$ [K].
cooled at 10°C/min to 150°C and annealed overnight in Ar. It was cooled further to 75°C and reheated in a stepwise fashion at 20°C/min in small 5–10°C increments to a series of temperatures between 75–410°C. After 4–5 min equilibration at each temperature −50 and +50 V were applied. $F^+(t)$ and $F^-(t)$ were measured at 2 sec resolution over about 60 sec. Between reversals from +50 V to −50 V the crystal was allowed to relax at 0 V for 2–5 min, depending upon the rate of creeping.

![Diagram](image.png)

**Fig. 5** CDA response of MgO under ±50 V below 425°C, indicating the appearance of mobile charges above 225°C and an enhancement of the positive surface charge.

Over the entire temperature interval $F^-$ was larger than $F^+$, e.g. $F_A > 0$, indicating a positive $P_{\text{surface}}$ contribution. Below 220°C both $F^-$ and $F^+$ are constant. Between 225–325°C, $F^-$ increased about three–to fourfold, reaching a small maximum around 325°C. $F^+$ decreased above 300°C and became repulsive above 325°C, indicating a positive $P_{\text{surface}}$ contribution strong...
enough to repel the sample from the bias electrode, e.g. from the region of higher electric field density.

It is instructive to compare the CDA response as shown in Fig. 5 with the d.c. conductivity of MgO as measured by conventional techniques over same temperature interval, 100–425°C [Kathrein and Freund, 1983]. As is apparent from Fig. 2 the conductivity is extremely low, $10^{-12} - 10^{-14} \text{ S/m}$, equivalent to $10^{-14} - 10^{-16} \text{ cm}^{-1}$, meaning that only $10^2 - 10^4$ charge carriers pass through $1 \text{ cm}^2$ sample cross section per volt-second. In order to measure such low conductivities special efforts have to be made to minimize electric stray fields, electrostatic charging, surface conduction and thermopotentials. In Fig. 6 we show the results of a d.c. conductivity measurement of an MgO disk of 14 mm diameter, 1 mm thickness with two circular Pt electrodes (7 mm effective diameter) and an annular guard electrode. The measurements were carried out under highest purity Ar on a fused silica support using shielded leads up to within 1 cm of the electrodes using a furnace with a low voltage SiC heater. The entire apparatus, including the d.c. circuit, was built into a fully enclosed Faraday cage, set on Teflon blocks. Batteries were used as a voltage source (500–1000 V) and the currents were measured in the $10^{-12} \text{ amps}$ range by means of a battery-operated KEITHLEY 610C electrometer during slow (2–5°C/min) heating/cooling cycles. The response was ohmic between 500–1000 V. Curve 1 represents the current measured after annealing the MgO crystal at 25°C for >12 hrs. Upon cooling the current fell into the <$10^{-13}$ amps range and eventually reversed sign, due to thermopotentials or electrode–related depolarization currents. Curves 2 and 3 were obtained upon reheating after 2–6 hrs of annealing at 100°C.
The conductivity maxima around 300–400°C and the hysteresis behavior which depended upon the annealing time were reproducible. When these measurements were reported several years ago [Kathrein et al., 1980] the nature of the charge carriers was still uncertain. On the basis of the CDA measurements depicted in Fig. 5 we can now say that the 200–400°C conductivity anomaly in the MgO crystals is not a “dirt” effect caused by the high d.c. voltages or due to an extraneous surface contamination. Furthermore, we now know that the charge carriers are positive and that they are capable of building up a strong internal field within the crystal.
Fig. 7 CDA response at the on-set of the large d.c. conductivity anomaly depicted in Fig. 2. The stepwise increase of $F^+$ above 400°C signals the appearance of mobile charges. The fact that $F^- > F^+$ indicates that the charges are positive and lead to a strong positive surface charge on the MgO crystal.

**Fig. 7** shows $F^-$ and $F^+$ between 275–550°C. At each temperature $F^-$ was measured first, followed by 2-3 min relaxation at 0 V, and $F^+$ measurement. $F^-$ remained larger than $F^+$, indicating a continuation of the positive surface charge. Above 400°C $F^-$ increased rapidly while $F^+$ decreased and eventually became repulsive around 475°C, indicating an enhancement of the positive surface charge. Creeping became prominent for $F^-$ between 400–525°C but was nearly absent from $F^+$. This was in disagreement with the pronounced creeping under positive bias shown in **Fig. 4b** probably because, after application of the negative bias, the system was given only relatively short time (2-3 min) to relax under 0 V before the positive bias was applied. Because a negative bias tends to attract positive charges to the surface, the subsequent application of the positive bias encountered a surface that was already in a saturated state.
The appearance of the strong positive surface charge ($F_\Delta$) between 450-500°C, coincident with the stepwise increase of the field-invariant dielectric polarization ($F_\Sigma$) and the concomitant sharp rise of the d.c. conductivity, does not support the notion of surface contamination [Freund, 1990; Freund et al., 1989]. Surface contamination cannot give rise to any surface charge that is maintained by an internal electric field.

**Fig. 8** Portion of the d.c. conductivity curve between 300-700°C (solid triangles) from Fig. 2 (left scale) plotted together with $F_\Sigma$ (open squares) and $F_\Delta$ (open circles) from a CDA experiment using a very slowly cooled MgO crystal.

In Fig. 8 we plot $F_\Sigma(T)$ and $F_\Delta(T)$ from a different run together with the 300-700°C section of the d.c. conductivity rise towards the anomaly from Fig. 2. $F_\Sigma(T)$, the sum of the field-invariant contributions, shows the large step-wise increase between 400-500°C that coincides
with the increase in the d.c. conductivity and an essentially flat portion above 600°C. Despite a small maximum towards positive values at 300°C $F_\Delta(T)$ was found to be weakly negative below 425°C — a behavior which seemed to be typical of very slowly cooled MgO crystals, possibly indicating an intrinsically negative surface under conditions approaching thermodynamic equilibrium, due to differences in the free energies of formation of anion and cation vacancies in the surface-near region [Kingery, 1974; Kliewer and Koehler, 1965]. Above 450°C, however, $F_\Delta$ turned strongly positive as it did in every CDA experiment conducted with these MgO crystals. In a narrow region around 500–550°C the steady state $F_\Delta$ values remained undetermined due to creeping. $F_\Delta$ stayed positive up to 950°C exhibiting a shallow maximum around 800°C.

According to eq. (9) it is best to measure $F_\Delta$ at low bias voltages because, according to eq. (5), $F^+$ and $F^-$ are proportional to the square of the bias voltage plus a linear term. Low bias voltages bring out this linear term. In Fig. 9a–c we plot $F^+$ and $F^-$ for three selected temperatures from 0 to +45 V and −35 V, measured at 2 V increments.

The “as measured” data show a repulsive response under positive bias and, less pronounced, an extra attraction under negative bias. The linear terms are given by the sections parallel to the abscissa above the critical bias voltage. Below this critical voltage ($U_{\text{crit.}}$) the absolute values $|F^0|$ and $|F^\pm|$ increase linearly with $U$ meaning that $P_{\text{surface}}$ is constant from 0 to $U_{\text{crit.}}$. Above this threshold value the surface contribution recedes as $P_{\text{surface}} \propto 1/U$. This means that the surface charge acts elastically like a spring which is pushed back into the bulk by the externally applied positive bias. Note that $U_{\text{crit.}}$ reaches values as high as +25 V. An internal electric field which pins positive charges to the MgO surface and can resist such a high externally applied field implies the presence of surface states which are energetically stable. The fact that such surface states can only be populated with charge carriers from within confirms our contention that the charge carriers in the MgO crystals under study are generated in the bulk.
MgO

550°C
Fig. 9 Deconvoluted $F^+$ and $F^-$ data at low bias voltages according to eq. (5). The solid circles are the "as measured" data. The open triangles represent the parabolic term which describes the ideal dielectric response, $F^\pm \propto U^2$. The open squares represent the linear term, $F_{0}^\pm$, arising from the presence of surface charges. Repulsion under positive bias indicates that the surface charge is positive. $U_{\text{crit}}$ is the bias voltage above which the externally applied electric field overcomes the internal electric field.

Since the MgO crystal represents a closed system and is electrically decoupled from the outside by the highly insulating fused silica hang-down fiber overall charge neutrality must be maintained. We therefore conclude that the inside of the MgO crystal contain a negative charge which is equal to the sum of all positive surface charges. The positive charges which come to the surface above 500°C diffuse relatively rapidly as is demonstrated by the kinetic CDA curves in Figs. 4a/b. Since all CDA phenomena reported here are reversible at sufficiently slow cooling rates, the charge carriers must be able to retrodiffuse into the bulk. While the charge carrier diffusion towards the surface is probably caused by mutual repulsion in the interior, the retrodiffusion may be driven by the field gradient between the positive surface and the negative interior.
Fig. 10  Variation of $F_\Delta$ as a function of temperature between 550–1000°C. The maximum observed around 750–800°C indicates that the density of positive charge carriers at the surface peaks around this temperature.

Fig. 8 already indicated that $F_\Delta$ reaches a maximum around 800°C. In Fig. 10 we focus on the temperature region 550–1000°C, plotting $F_\Delta(T)$ as measured in the 0 to ±30 V bias range in 2 V and 5 V incremental steps. The plot confirms that $F_\Delta$ is temperature dependent and reaches a maximum around 750–800°C. Prolonged exposure to temperatures above 800°C led to an irreversible loss of CDA response similar to the irreversible loss of the anomalously high d.c. conductivity that was shown for MgO in Fig. 2 and for olivine in Fig. 1. The probable cause for this irreversibility will be discussed in more detail in Part II.
Conclusion

CDA has provided us with the means to show that, contrary to widespread belief, the d.c. conductivity anomalies below 700–800°C in MgO cannot be due to surface damage or contamination either prior to or during the experiment. Instead the CDA data suggest that some peculiar charge carriers exist in the bulk of the MgO crystals under study. These charges are reversibly generated during heating and consumed during cooling. They carry a positive sign. The process is reversible upon cooling but becomes partly irreversible after the crystals have been heated for extended periods of time at temperatures exceeding 700–800°C, even in pure Ar. The strongest argument so far in favor of an indigenous source is based on the appearance of the positive surface charge in both temperature intervals of anomalously high d.c. conductivity, between 200–400°C and between 500–800°C. The unique features of CDA is its capacity to show that the surface charge is sustained by a strong internal electric field. Such an internal field implies that the charges originate in the bulk and are driven to the surface by some internal force.

This conclusion invites speculations about the nature of these charge carriers and how they were introduced into the nominally high purity MgO crystals. So far we know that these charge carriers are positive and mobile at temperatures at which ionic mobilities, for instance of \( \text{Mg}^{2+} \) or \( \text{O}^{2-} \) via vacancy hopping or by interstitial diffusion, are still extremely small in MgO [Sempolinski and Kingery, 1980; Sempolinski et al., 1980]. Only holes, e.g. defect electrons, can have sufficient mobility to explain the observations. Now, holes can exist on (i) the cation or (ii) the anion sublattice. In case (i) they would correspond to a transiently oxidized transition metal cation, for instance \( \text{Fe}^{3+} \) on a \( \text{Mg}^{2+} \) site. In case (ii) they would consist of \( \text{O}^- \) on an \( \text{O}^{2-} \) site and as such deserve the designation of “positive holes” [Griscom, 1990; Henderson and Wertz, 1977]. Given the low transition metal concentration in the MgO crystals under study it is unlikely that cation impurity–bound holes are responsible for the observed phenomena. Therefore, positive holes remain the only candidates. In Part II we shall present data obtained by other techniques which allow us to go beyond the as yet tentative assignment and unambiguously identify the charge carriers that are responsible for the electrical conductivity anomalies in MgO below 700–800°C.

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References


Electrical Conductivity Measurements under Minimum Perturbation Conditions

Part II:
O⁻ Charge Carriers in Magnesium Oxide

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Abstract

Anomalously high electrical conductivity is observed in MgO single crystals (99.99% purity grade, <50 ppm transition metal impurities plus traces of OH⁻, grown by arc fusion) below ≈800°C. This conductivity anomaly is widely believed to be a “dirt effect”, caused by surface contamination. However, this conjecture cannot be upheld. Infrared studies show that OH⁻ in the bulk undergo changes in the exactly same temperature intervals in which the conductivity shows anomalies. Between 500–800°C H₂ molecules are being generated out of OH⁻, implying an internal redox reaction: OH⁻ + OH⁻ = H₂ + O₂⁻. The appearance (disappearance) of mobile charge carriers upon heating (cooling) coincides with oxidation (reduction) of trace transition metal cations, measured by electron spin resonance spectroscopy. This is consistent with dissociation of peroxy defects, O₂⁻, into O⁻. Magnetic susceptibility measurements indicate spin-decoupling of diamagnetic O₂⁻ between 300–500°C into paramagnetic O⁻. The O⁻ concentration was estimated to be of the order of 1000 ppm. Conversion of OH⁻ into H₂ plus O₂⁻ is corroborated by a mass spectrometric study of MgO ex Mg(OH)₂ which releases molecular H₂ (2000–5000 ppm) above 400°C and atomic O above 600°C, consistent with disproportionation: O₂⁻ = O²⁻ + O. All data jointly indicate that the electrical conductivity anomaly of MgO single crystals is not a “dirt effect”, caused by surface contamination, but that it is due to the generation of O⁻ charge carriers which occurs in the bulk. The underlying reaction involves the redox conversion of lattice–bound OH⁻ into H₂ plus O₂⁻. These O₂⁻ coexist with transition metal cations in lower oxidation states. If the H₂ molecules diffuse out, the MgO crystals retain O₂⁻ defects, equivalent to interstitially bound excess oxygen, MgO₁+δ. This is a solid state reaction by which crystals which originally grew under reducing conditions can become internally oxidized.

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Introduction

Scientists have traditionally worked very hard at cleaning and perfecting the materials they want to study. Nature, however, has worked equally hard at evading scientists, and many studies of material properties have ended ungloriously in the file labeled “dirt effects”. It has been known for some time that mantle-derived olivine crystals display a somewhat bothersome, anomalously high electrical conductivity below ≈700°C [Duba, 1972; Duba et al., 1974; Duba and Nicholls, 1973; Schock et al., 1977; Sockel, 1974]. It is reported that this anomalous behavior is poorly reproducible and disappears upon prolonged heating in reduced gases. This has led to the widespread belief that the anomaly is probably caused by contamination of some kind. At closer inspection it is found that conductivity anomalies are not unique to mantle-derived olivine crystals. They are also known from studies of other crystals, in particular synthetic MgO crystals where the conductivity enhancement covers many orders of magnitude up to ≈800°C [Lempicki, 1953; Mansfield, 1953; Lewis and Wright, 1968; Lewis and Wright, 1970; Osburn and Vest, 1971; Kathrein and Freund, 1983]. In MgO, like in olivine, the anomaly disappears upon prolonged heating to 700–800°C and has been reported to be poorly reproducible. It therefore appeared reasonable to assume that it had something to due with contamination, specifically surface contamination. Though there were enough hints to suggest that the situation might be more complicated, researchers have generally been reluctant to spend time and effort pursuing this enigmatic anomaly for fear that Nature may once more be successful in setting them up with a “dirt effect”.

Part I [Freund et al., 1991] re-examined the anomalous conductivity behavior of MgO and found it worth a detailed study applying a new technique, Charge Distribution Analysis (CDA). In the course of this study two distinct temperature regions of anomalous conductivity behavior were identified, extending from 200 to 400°C and from 500 to 800°C. In both temperature regions the effect was caused by charge carriers that reside predominantly at the surface. However, these charge carriers were not related to some kind of surface contamination. CDA gave strong indications that they arose in the bulk and that their propensity for the surface is the result of an internal driving force.

While CDA was able to ascertain that these charge carriers were positive, it could not unambiguously identify their chemical nature. To meet this objective other techniques are needed. The following presentation is based on work which has mostly been carried out in my laboratory with students and collaborators whose names are acknowledged at the end. Work by other groups will be referenced sparsely. This is despite the fact that MgO has been studied intensively over many years [e.g. Henderson and Wertz, 1977]. However, most researchers have either passed over the temperature region where the electric conductivity anomalies occur or avoided making correlations between their specific results, obtained by other techniques, and the conductivity response.
Experimental

As in the CDA experiments described in Part I arc-fusion grown, nominally high purity MgO single crystals were used, obtained from W. & C. Spicer, Ltd., Cheltenham, England. They were of the same purity (4N = 99.99 %) or a somewhat lower purity grade (3N = 99.9 %) with typically 70–100 ppm Fe, 10–20 ppm Cr and <10 ppm Mn as the crystals used in the d.c. and CDA studies [Freund et al., 1991; Kathrein and Freund, 1983]. Lesser purity grade crystals (99.5 %) were included, produced by Dynamit–Nobel AG, Troisdorf, Germany, in an industrial carbon arc production furnace. These crystals, designated below as DN–MgO, have a higher Ca, Al and Si content. The 4N–MgO crystals typically contain =10 ppm Mn, Cr, Ni and =30 ppm Fe plus Ca, Al and Si of about 10 ppm each. They are slightly yellowish–brown due to color centers [Henderson and Wertz, 1977] and often cloudy, due to submicron–sized cavities decorating dislocation lines and subgrain boundaries. All crystals, irrespective of their purity grade, contain OH⁻ impurities at concentrations in the 10–100 ppm range introduced by H₂O adsorbed to the MgO starting powder [Abraham et al., 1971; Butler et al., 1971]. Specpure grade MgO powder was obtained from Johnson Matthey Chemicals, Ltd., Batch # 3.51934, quoted to contain =3 ppm Fe, =1 ppm Ag, <1 ppm Ca and <1 ppm Cu.

For the infrared (IR) spectroscopic study a PERKIN–ELMER Model 225 grating spectrometer was used, modified to cover the range from 3000–4500 cm⁻¹. All spectra were recorded at liquid nitrogen temperature (=80 K) after heating the samples either in situ or externally to temperatures up to 1000°C. A custom–built stainless steel cryostat was available for in situ treatments up to 550°C. The cryostat was contained in an ultrahigh vacuum chamber equipped with NaCl and sapphire single crystal windows and pumped to 5 × 10⁻⁸ torr base pressure by means of a turbomolecular pump. Screwed to the bottom of the cryostat was a Cu block with a rectangular opening. A ceramic block containing a pair of 100 W heating elements could be brought in mechanical contact with the Cu block by means of a linear motion drive. Cleaved MgO crystals of 10–20 mm thickness and =10 mm width were wrapped with 0.1 mm Au foil (for thermal contact) and placed inside the Cu block. They were heated repetitively in 20-25°C increments for 30 min and then either quenched by pouring liquid nitrogen directly into the hot cryostat after decoupling the heater block or cooled slowly by allowing the system to cool dynamically to room temperature before filling the cryostat with liquid nitrogen. To reach temperatures >550°C the crystals were externally heated in a tube furnace for 30 min, air–quenched to ambient temperature and subsequently mounted in the cryostat Cu block for cooling to 80 K.

The electron paramagnetic resonance (EPR) study was conducted with a BRUKER B–ER 414 spectrometer operating at 9.7 GHz. A 2 x 2 x 2 mm MgO crystal doped with 760 ppm Cr was used as a calibration standard. Cleaved 3N–MgO single crystals (3 x 3 x 12 mm) and coarsely crushed DN–MgO crystals (100–385 μm size fraction) were sealed in thin–walled fused silica tubes under high purity He (Air Liquide), heated for 5–30 min in 50°C incremental steps up to 1000°C and for 5 min down in the same temperature steps to 400°C. Each temperature treatment
was following by a rapid quench in liquid N$_2$. The EPR spectra were recorded at 77 K or room temperature. The repositioning of the samples in the resonance cavity was reproducible with ±1% relative intensity variation.

The magnetic susceptibility study was conducted with a EG&G Princeton Applied Research Vibrating-Sample Magnetometer (VSM) Model 4500, combined with a 5.4 KGauss Varian magnet, model V-2200A. The VSM was equipped with a high temperature sample holder (25–1000°C) made of boron nitride (BN) within a double–wall vacuum chamber. The interior sample chamber was evacuated and refilled with 5 torr He for thermal contact. The MgO single crystal was ground into a cylinder of 4.8 mm diameter and 15.6 mm height. It was mounted with a small amount ZrO$_2$-based high-temperature cement onto the BN rod, preheated to 980°C and cooled slowly. The data were collected during reheating at 4°C/min. The magnetic response of the BN sample rod with cement, as determined in a separate run, was subtracted from the data.

The mass spectroscopic data were collected in an all–glass, hydrocarbon–free ultrahigh vacuum (UHV) system, bakeable to 450°C, with a base pressure in the 10–$^{10}$ torr range. The system used electromagnetically operated polished spherical stopcocks as valves and was pumped by two Hg diffusion pumps in tandem between liquid nitrogen traps. The sample was placed in a fused silica tube (4 mm i.d.), connected to the glass apparatus via a graded seal, and heated with a free–standing Pt coil in an outer glass chamber filled with Ar (to prevent diffusion of H$_2$ through the hot fused silica wall). The gases evolved from the sample were directed through selectable glass capillaries of known conductance into an Omegatron cyclotron resonance mass filter, built entirely of Pt–Ir alloy except for the ThO$_2$–covered tungsten wire for the electron source. Typically 30–70 μg specpure MgO powder samples were placed in the fused silica sample tube and outgassed for several hours at 1000°C. They were then hydroxylated and deuterioxylated to Mg(OH)$_2$ and Mg(OD)$_2$ respectively, using double–distilled H$_2$O and 99.7% D$_2$O (Merck Uvasol) at 355 torr equilibrium pressure. The hydroxides and deuteroxides thus obtained were very slowly decomposed to finely divided MgO containing a high concentration of residual OH$^-$ and OD$^-$ along with point defects. The pressure inside the UHV system was measured with a Bayart–Alpert gauge and the heating rate of the sample was controlled through an automated feed–back loop to keep the degassing rate within the dynamic range (1:10$^6$) of the Omegatron. After the main decomposition when the degassing rate decreased rapidly the Bayart–Alpert gauge was shut off to avoid decomposition at the hot wire. The purpose of this experiment was to determine small amounts of H$_2$/HD/D$_2$ and/or O$_2$/O in the presence of H$_2$O/HDO/D$_2$O vapor.
Results and Discussion

Infrared Spectroscopy of OH⁻ Impurities

OH⁻-bearing point defects in MgO single crystals were studied by IR spectroscopy over a very wide temperature range using the experimental procedures outlined above [Freund and Wengeler, 1982]. The OH⁻ impurities cause two prominent groups of bands. One group is relatively strong and consists of three, sometimes four equispaced bands at 3297, 3312, 3327, and 3342 cm⁻¹. This “quadruplet” has been assigned to a defect where a single OH⁻ sits adjacent to an Mg²⁺ vacancy, V''ₘ₉, and compensates half of its charge. Using the Kröger [1964] defect denomination* this defect is written as [OH' V''ₘ₉]. The second group is of lower intensity and consists of three bands at 3550, 3571, and 3566 cm⁻¹. This “triplet” has been assigned to a defect where two OH⁻ sit adjacent to an Mg²⁺ vacancy and fully compensate its charge, [OH'' V'ₘ₉ HO]⁺.

Fig. 1a/b shows a remarkable result. The intensity variations indicate that the number of OH⁻ at the defect sites varies considerably with temperature between 25°C and 950°C. The variations are fully reversible, indicating first that the crystals behave as a closed system and second, that a complex redistribution of OH⁻ defects takes place during heating and cooling which involves several internal reservoirs. Third, the relative intensities of the two groups are not in agreement with what one would predict from simple defect equilibria.

On the basis of minimum energy considerations, e.g. minimalization of the the local Coulomb potential, the fully compensated defects, [OH'' V''ₘ₉ HO]⁺ should be significantly more abundant that the half compensated defects, [OH' V''ₘ₉]. Instead, the triplet in (b) is 5–10 times weaker than the quadruplet in (a). In addition the intensities of both groups vary significantly not only as a function of temperature but also as a function of the cooling rate. The latter is particularly pronounced between 200–400°C: slowly cooled crystals give higher intensities than rapidly cooled crystals. Now, the 200–400°C interval is the same which stood out in the conductivity measurements as a region of anomalous electrical conductivity behavior (see Figs. 5/6 in Part I [Freund et al., 1991]). There are two possible conclusions: (i) the conductivity anomaly is directly related to OH⁻ defects, and (ii) the OH⁻ defects act as local probes for something else happening in the bulk which in turn is responsible to the conductivity anomaly.

* Superscript dot, prime, and cross indicate a positive, negative and neutral charge with respect to the ideal structure. V means vacancy. Subscripts characterize the lattice site except for O²⁻ sites which are omitted. Subscript i stands for interstitial. Square brackets delineate the point defect, generally omitting lattice points that are not essential.
Fig. 1a: Intensity variations as a function of temperature of the quadruplet group, assigned to the O–H stretching vibration, $v_{\text{OH}}$, of single OH$^-$ associated with an Mg$^{2+}$ vacancy. Each data point represents the IR intensity after the crystal had been heated to the indicated temperature and cooled. All IR measurements were done at 80 K [after Freund and Wengeler, 1982].
Fig. 1b: Intensity variations as a function of temperature of the triplet group, assigned to two OH⁻ associated with an Mg²⁺ vacancy. Each data point represents the IR intensity after the crystal had been heated to the indicated temperature and cooled. All IR measurements were done at 80 K [after Freund and Wengeler, 1982].

Turning to the T–dependent intensity variations we note that the trends are opposite: while the quadruplet intensity increases by a factor of 4 between 500–800°C, the triplet intensity decreases. This is against what one would predict on the basis of simple defect equilibria. If the number of [OH⁻ V⁰_Mg]⁻ defects increases (because a “reservoir” in the bulk releases H⁺), the number of [OH⁻ V⁰_Mg⁻ HO]²⁻ defects should increase as well. Instead they decrease.
The "reservoir" consists of $H^+$ on interstitial sites, $H^+_i$, which manifest themselves in broad background absorption that extends from 3300–3700 cm$^{-1}$ [Freund and Wengeler, 1982]. Around 500°C these $H^+_i$ diffuse to existing $Mg^{2+}$ vacancies and form additional $OH^-$-compensated cation vacancy defects:

\[
[O^x \, V^\prime_{Mg}\, ]'' + H^+_i = [OH^'' \, V^\prime_{Mg}]' \tag{1}
\]

\[
[O^x \, V^''_{Mg} \cdot O]'' + 2 \, H^+_i = [OH^'' \, V^\prime_{Mg} \cdot OH]x \tag{2}
\]

However, the fact that the intensity of the triplet assigned to the $[OH^'' \cdot V^\prime_{Mg} \cdot OH]x$ defect decreases above 500°C suggests a concurrent second reaction by which these fully $OH^-$-compensated defects convert into another type of defect that bears no $OH^-$. This second reaction correlates with the appearance of a weak band around 4150 cm$^{-1}$. On the basis of partially deuterated MgO crystals this band has been assigned to the H–H stretching vibration, $v_{H-H}$, of $H_2$ molecules occupying non–symmetrical defect sites [Freund and Wengeler, 1982].

This leads to the next question: how do $[OH^'' \cdot V^\prime_{Mg} \cdot OH]x$ defects give rise to $H_2$ molecules?

**Fig. 2** illustrates the reaction by which, due to the perturbation of the local Coulomb potential at the $Mg^{2+}$ vacancy site [King and Freund, 1984], the two $OH^-$ are destabilized with respect to the valency state of oxygen. It is often not appreciated that oxygen has two ionization states, $O^-$ and $O^{2-}$, of which only the first is thermodynamically stable, e.g. has a positive electron affinity [Edlén, 1960]. Upon addition of the first electron, $O + e^- = O^-$, the energy gained amounts to 1.4 eV. However, when the second electron is added, $O^- + e^- = O^{2-}$, the highly negative second electron affinity, $-8.9$ eV, comes to bear, making this reaction highly endothermal. At the same time, the lattice energy of a crystal like MgO, $-40$ eV/mole, primarily derives from the Coulomb interaction between the oppositely charged ions, $Mg^{2+}$ and $O^{2-}$. By containing divalent $O^{2-}$ interacting with divalent $Mg^{2+}$ the system more than recovers the energy expended on the negative second electron affinity of the oxygen anions. Hence, MgO is stable. This overall stability of the structure does not necessarily mean that $O^{2-}$ will be stable at every point in the crystal structure. A cation vacancy, for instance, represents a large perturbation of the local Coulomb energy. The possibility has to be considered that the $O^{2-}$ becomes destabilized with respect to the $O^-$ state. According to the IR data [Freund and Wengeler 1982] this destabilization indeed takes place when $OH^-$ neighbor a $Mg^{2+}$ vacancy: $OH^- = O^- + H$. Additional energy may be gained (i) by the recombination of two $H$ to $H_2$:

\[
[OH^'' \, V^\prime_{Mg} \cdot OH]x = [O^- \, (H_2)^x \cdot Mg] \tag{3}
\]

and (ii) by spin pairing of two $O^-$, leading to an $O^-$ dimer, chemically equivalent to a peroxo anion, $O_2^{2-}$:

\[
[O^- \, (H_2)^x \cdot Mg] = [O^- \cdot (H_2)^x \cdot Mg] \tag{4}
\]
Some of the energy gain is expected to be counteracted by lattice strain in the neighborhood of the defect which arises from the short (≈1.5 Å) O–O bond [Cremer, 1983] and the necessity to accommodate the local contraction.

\[
\begin{align*}
\text{MgO} & \\
\end{align*}
\]

\[
\begin{align*}
\text{\(^2-\)} & = \text{O}^{2-} \\
\text{\(\text{O}^-\)} & = \text{O}^\cdot
\end{align*}
\]

**Fig. 2** Conversion of two OH\(^-\) associated with an Mg\(^{2+}\) vacancy defect into O\(^{2-}\) (peroxy) plus H\(_2\).

In addition H\(_2\) molecules can leave the cation vacancy site and diffuse away. If the crystals are small and/or the time is long, the H\(_2\) molecules diffuse out, leaving behind peroxo-compensated Mg\(^{2+}\) vacancies:

\[
[O\text{O}^{-}(\text{H}_2)_{\text{Mg}}]^x = [O^{-}V_{\text{Mg}}]^x + (\text{H}_2)^\uparrow \tag{5}
\]
Identification of O\textsuperscript{-} by Magnetic Susceptibility Measurements

MgO is intrinsically diamagnetic, $\chi_{\text{dia}} = -10.8 \times 10^{-6}$ emu/mole and its diamagnetism is independent of temperature [Nagel et al., 1976]. Protons will not change the magnetic susceptibility of MgO\textsuperscript{**}. The O\textsubscript{2}\textsuperscript{-} anion, if it forms according to eq. [5] and Fig. 2, has 14 valence electrons and is diamagnetic. O\textsuperscript{-} states, however, have 7 electrons and an unpaired electron spin.

Fig. 3 Anomalous paramagnetic contribution observed during heating at 4°C/min of a nominally high purity MgO single crystals (99.99%, <50 wt.-ppm transition metal impurities). The error bar indicates the probable uncertainty of the measurements (after [Batlló et al., 1991]).

\*\* The nuclear spin of the proton is orders of magnitude too small to be of interest here.
Fig. 3 shows the magnetic susceptibility $\chi$ of a 4N-MgO crystal between 180–980°C. The crystal was first heated to 900°C, cooled at 10°C and reheated to 980°C at 4°C/min while measurements were taken [Batllo et al., 1991]. The $\chi(T)$ curve was adjusted at 200°C to the intrinsic diamagnetism of MgO, corrected for the small transition metal impurities, <50 at.-ppm.

Fig. 3 shows a paramagnetic anomaly that starts above 300°C and reaches a maximum around 500°C, approximately +1.5 $10^{-6}$ emu units above the base value. This rise in paramagnetism indicates the generation of unpaired spins in the MgO matrix. The few 3d transition metal impurities cannot explain this anomalous rise. Mg$^{2+}$ cannot turn paramagnetic, for instance by auto-reduction to Mg$^+$ or auto-oxidation to Mg$^{3+}$ because the +1 and +3 valency states of Mg are highly unstable. By default we are left with O$^-$ as the only alternative paramagnetic species that may form thermally in the MgO matrix, for instance through dissociation of existing peroxy defects, O$_2^-$.

Assuming that the Landé factor for O$^-$ in the O$_2^-$ matrix of MgO is the same as for free O$^-$, the magnitude of the paramagnetic contribution $\chi_{\text{para}}$ at 500°C gives an O$^-$ concentration of the order of 1000 at.-ppm [Batllo et al., 1991]. This value is significantly higher than the typical concentration of residual OH$^-$ in arc-fusion grown MgO [Abraham et al., 1971; Freund and Wengeler, 1982], suggesting that a major fraction of OH$^-$ originally contained in these MgO crystals has converted to O$_2^-$ defects as described by eqs. [3-5].

Next we have to address the question whether the observed paramagnetic anomaly supports the contention that O$^-$ are the mobile charge carriers that are also responsible for the electric conductivity anomaly and positive surface charge determined by CDA [Freund et al., 1991]. For this we need to understand why the paramagnetism starts around 300°C and reaches its maximum around 500°C, whereas the large electric conductivity anomaly begins around 500°C and reaches its maximum near 800°C.

The paramagnetic anomaly is consistent with the prediction that two unpaired O$^-$ are unstable at low temperatures because they undergo self trapping to form the diamagnetic peroxy defect, t'O$_2^-$ [Batllo et al., 1991; King and Freund, 1984]. By virtue of its short (=1.5 Å) O–O bond [Cremer, 1983] the peroxy anion represents a small polaron, e.g. it is associated with a strong and highly localized lattice strain. The dissociation of t'O$_2^-$ into two paramagnetic O$^-$ occurs in two sequential steps described by eqs. [6] and [7]: (i) decoupling the spins; (ii) dissociating into an O$^-$ bound to the Mg$^{2+}$ site plus an unbound O$^-$ which delocalizes through O$^{2-}$/O$^-$ valency fluctuations. This highly mobile, unbound O$^-$ represents a large polaron, e.g. it is associated with lattice strain spread over a relatively large volume [Freund, 1990]:

$$[t'O_{\text{V}_\text{Mg}}] \times = [t'O_{\text{V}_\text{Mg}} O^+] \times$$

[6]

$$[t'O_{\text{V}_\text{Mg}} O^+] \times = [t'O_{\text{V}_\text{Mg}}] + O^-$$

[7]
The defects with two O$^-$ and one O$^-$ at the Mg$^{2+}$ vacancy site are also known as V$_0$ and V$^-$ centers, respectively [Henderson and Wertz, 1977]. The magnetic susceptibility data presented in Fig. 3 are consistent with a 2-step dissociation by which V$_0$ centers start to form around 300°C according to eq. [6]. The maximum of the paramagnetic anomaly is reached when all O$_2^-$ have become V$_0$ centers. Above 500°C the V$_0$ centers dissociate into V$^-$ centers plus unbound O$^-$ according to eq. [7] without further increase in the number of spins. Since $\chi_{\text{para}}$ is proportional to $\frac{1}{T}$, it is expected that $\chi(T)$ decreases above 500°C as shown in Fig. 3.

**Evidence for O$^-$ from Electron Paramagnetic Resonance Studies**

If peroxy defects exist in the MgO crystals and dissociate according to eq. [7], releasing O$^-$ as mobile charge carriers in the O$^{2-}$ sublattice, EPR spectroscopy can be used in two ways: (i) to measure the concentration of quenched V$^-$ centers or (ii) to measure the oxidation of transition metal cations according to:

\[
\begin{align*}
\text{Fe}^{2+} + \text{O}^- &= \text{Fe}^{3+} + \text{O}^{2-} & [8a] \\
\text{Mn}^{2+} + \text{O}^- &= \text{Mn}^{3+} + \text{O}^{2-} & [8b] \\
\text{Cr}^{3+} + \text{O}^- &= \text{Cr}^{4+} + \text{O}^{2-} & [8c]
\end{align*}
\]

Fe$^{2+}$, Mn$^{2+}$ and Cr$^{3+}$ represent the oxidation states in which these cation impurities are normally found in the MgO crystals. The oxidation states, Mn$^{3+}$ and Cr$^{4+}$, are unstable with respect to disproportionation [Henderson and Wertz, 1977]. The presence of V$^-$ centers in rapidly quenched CaO has been reported early [Tomlinson and Henderson, 1969] but no systematic study seem to have followed this lead. We used EPR to measure the appearance of the Fe$^{3+}$ signal and disappearance of Mn$^{2+}$ and Cr$^{3+}$ signals as a result of oxidation [Kathrein et al., 1984].
Fe$^{3+}$ in DN-MgO
(b) 

Relative EPR Intensities

\[ \text{Mn}^{2+} \text{ in DN-MgO} \]

Temperature [°C]
Fig. 4 Relative intensities of the EPR signals for three transition metal impurities (a) Fe$^{3+}$, (b) Mn$^{2+}$, and (c) Cr$^{3+}$ in DN-MgO heated (solid lines) and cooled (dashed line) in steps under He, measured at 77 K (after Kathrein et al., 1984). The reversible increase of Fe$^{3+}$ intensity and decrease of Mn$^{2+}$ and Cr$^{3+}$ intensities between 500–800°C indicates reversible redox reaction inside the MgO matrix.

Fig. 4a–c shows that the number of Fe$^{3+}$ increases between 500–850°C and decreases between 850–1000°C. Upon cooling to 400°C with 5 min holding time at each temperature step the oxidation reverses, e.g. auto-reduction occurs. Likewise, the intensity of the Mn$^{2+}$ and Cr$^{3+}$ signals decreases upon heating to 1000°C and recovers upon stepwise cooling, also suggesting auto-reduction. The reversibility and the speed with which the reversibility is achieved is inconsistent with the proposition that gaseous O$_2$ may be involved in these redox reactions. Instead, the experimental data strongly suggest an internal process driven by O$^-$ states that are thermally generated upon heating above 500°C and disappear upon cooling according to eqs. [8a–c]. The results are also in agreement with the magnetic susceptibility data presented above,
indicating that the internal redox reactions of the transition metal cations are coupled to the dissociation of the peroxy defects:

$$2 \text{Me}^{n+} + O_2^{2-} \leftrightarrow 2 \text{Me}^{(n+1)+} + 2 \text{O}_2^{-} \quad [9]$$

At low temperatures when the $O^-$ are self-trapped and, hence, localized (see eq. [6]) transition metal cations in low oxidation states coexist with $O_2^{-}\,$ in the same MgO matrix. An Fe$^{2+}$ impurity on a regular Mg$^{2+}$ site represents neutral defect, [Fe$^{2+}_\text{Mg}$]$^x$. Likewise, the V$_0$ center and the peroxy anion adjacent to a Mg$^{2+}$ vacancy are neutral defects, [O$^{\text{v"O"}}_{\text{Mg}}$]$^x$ and [O$^{\text{O"}}_{\text{Mg}}$]$^x$, respectively. Between 500–800°C, when the V$_0$ centers or peroxy entities dissociate according to eq. [7] and release unbound O$^-$, some O$^-$ are captured by the transition metal cations leading to their oxidation. An Fe$^{3+}$ on a regular Mg$^{2+}$ site represents a positive charge, [Fe$^{3+}_\text{Mg}$]$^x$. The V$^-$ center is the remnant of the peroxy defect after releasing an unbound O$^-$ (see eq. [7]). It represents a negative charge, [O$^{\text{v"O"}}_{\text{Mg}}$]$^x$. Therefore, the capture of an O$^-$ state by any transition metal cation leaves two oppositely charged defects. Their long-range Coulomb interaction provides the force which, upon cooling, drives the auto-reduction of the transition metal cations, e.g. causes eq. [9] to proceed from right to left.

**Fig. 4a** showed that, after the nearly two-fold increase between 500–800°C, the Fe$^{3+}$ intensity decreases above 800°C, suggesting that there is yet another redox process at high temperatures. This is reminiscent of the decrease of the anomalous electrical conductivity above 800°C as measured by d.c. techniques and of the positive surface charge as measured by CDA and expressed through $F_\Delta$ [Freund et al., 1991]. To illustrate the correlation between the EPR data for the $O^-$ driven Fe$^{2+}$/Fe$^{3+}$ redox reaction and the d.c. conductivity data **Fig. 5** replots the d.c. conductivity of a 4N–MgO single crystal (in log Siemens units versus temperature in °C) and one of the EPR Fe$^{3+}$ intensity curves from **Fig. 4a**.

**Emission of H$_2$ Molecules and O Atoms**

It would be certainly be of value to be able to provide an independent proof that OH$^-$ defects in MgO are responsible for the *in situ* formation of H$_2$ plus O$_2^{-}\,$. To address this question I turn to the last technique to be discussed here: the mass spectroscopic measurement of gas evolution from finely divided, ultrahigh purity MgO obtained by thermally decomposing Mg(OH)$_2$.

According to textbook knowledge the decomposition of Mg(OH)$_2$ should be a simple, straightforward reaction: Mg(OH)$_2$ = MgO + H$_2$O. However, the finely divided MgO first formed still contains residual OH$^-$ balanced by Mg$^{2+}$ vacancies. Its composition may be written as $\text{Mg}_{1-\delta}\text{O}_{1-2\delta}(\text{OH})_{2\delta}$, with $\delta$~1. At sufficiently high temperatures, however, the OH$^-$ should be expelled as H$_2$O leaving stoichiometric MgO.
Fig. 5 Correlation between the electrical conductivity anomaly between 500–800°C as measured by d.c. techniques and the Fe$^{3+}$ intensity variation as measured by EPR (after [Kathrein and Freund, 1983; Kathrein et al., 1984]) providing support that the anomalous rise of the d.c. conductivity and the internal oxidation of Fe$^{2+}$ to Fe$^{3+}$ have a common cause, viz. the generation of unbound O$^-$ states.

The redox reaction given in eq. [5] indicates that H$_2$ molecules may form at OH$^-$–compensated Mg$^{2+}$ vacancy sites. As long as the H$_2$ molecules remain in the MgO matrix the bulk composition remains unchanged, but the non-stoichiometric MgO should be written Mg$_{1-\delta}$O$_{1-2\delta}$(O$_2$)$_{\delta}$(H$_2$)$_{8\delta}$ = Mg$_{1-\delta}$O(H$_2$)$_{8\delta}$. The situation changes as the H$_2$ molecules diffuse out of the MgO crystals:

$$\text{Mg}_{1-\delta}\text{O}(\text{H}_2)_{8\delta} = \text{Mg}_{1-\delta}\text{O} + \delta \text{H}_2$$

[10]

The cation–deficient MgO on the right hand side contains excess oxygen in the form of peroxo defects, O$_2^{2-}$. With $\delta<1$ it may also be written as MgO$_{1+\delta}$. Under thermodynamic equilibrium conditions such MgO with excess oxygen can only be expected to form at a high O$_2$ pressure, never under reducing conditions and never in a vacuum [Osburn and Vest, 1971; Wriedt, 1987]. Obviously the MgO$_{1+\delta}$ will not be stable. Upon further heating it is expected to decompose by...
releasing its excess oxygen just like any inorganic peroxide decomposes into oxide plus $O_2$ [Cotton and Wilkinson, 1980]. However, microscopically this decomposition reaction is a rather complicated process. It proceeds by way of $O^-$ diffusing to the surface and combining to $O_2^{2-}$ which in turn release atomic $O$ through disproportionation:

$$O^- + O^- = (O_2^{2-})_{surface} = O^{2-} + O \uparrow$$  \[11\]
Fig. 6a/b shows the H$_2$O, H$_2$ and O evolution as measured mass spectroscopically starting from ultrapure Mg(OH)$_2$ and leading to finely divided MgO [Martens et al., 1976]. The H$_2$O evolution begins above 200°C and leads to a maximum near 300°C. Above 300°C cubic MgO forms, e.g. Mg$_{1-8}$O$_{1.28}$(OH)$_{28}$, with $\delta<1$. At the same time H$_2$ molecules start to evolve. Above 600°C the H$_2$ evolution is joined by O evolution which reaches a maximum around 750°C and then decays. The H$_2$ evolution above 900°C comes from the hot fused silica sample tube. The H$_2$ evolution between 300–900°C as well as the O evolution above 600°C comes from the OH$^-$–containing MgO.

Fig. 6b dramatically illustrates the sequence of reactions described by eqs. [10] and [11]. The number of H$_2$ molecules evolved was found to vary between 2000–5000 ppm with respect to the number of OH$^-$ contained in the original Mg(OH)$_2$, the higher value applying to deuterated samples [Freund et al., 1982]. This implies an equal number of peroxy defects. Using an independent approach, a wet-chemical titration technique, Tench and coworkers [Praliaud et al.,
found highly oxidizing properties in MgO produced in the same way, through dehydration of high purity Mg(OH)$_2$ and determined the concentration of oxidizing centers to be $3.5 \times 10^{19}$ g$^{-1}$ MgO, equivalent to 2250 at.-ppm. This proves that dehydration can lead to oxidation, even in the absence of transition metal cations. The only way to balance the redox reaction is to let O$^2-$ be oxidized to O$^-$ while reducing protons to H$_2$. The 2250 at.-ppm value determined by titration is in good agreement with the range of peroxy concentrations obtained from the H$_2$ evolution of MgO shown in Fig. 6b. It corroborates the 1000 ppm O$^-$ concentration value derived from the magnetic susceptibility data shown in Fig. 3.

According to eq. [11] the O emission from the MgO surface irreversibly consumes O$^-$. This process is expected to affect the electrical conductivity and the CDA response, in particular $F_\Delta$ which is an indicator of the O$^-$ surface density. In the preceding paper [Freund et al., 1991] we reported that the anomalously high electrical conductivity of MgO ended around 700–800°C when the log $\sigma$ vs 1/T [K] curve joined the high temperature branch [Sempolinski and Kingery, 1980]. $F_\Delta$ was at a maximum around 750°C.

Fig. 7a combines the evolution of O atoms as measured from the MgO powder by mass spectrometry with $F_\Delta(T)$ as measured with an MgO single crystal by CDA. Though the two experiments were carried out under very different conditions, the results closely match. The onset of the O emission coincides with the build-up of the high positive surface charge. The maximum O emission rates lies within 50°C of the $F_\Delta$ maximum. Fig. 7b combines the O evolution curve with the rising portion of the d.c. conductivity between 500–800°C (plotted as log $\sigma$ versus T [°C]). Both plots support the arguments leading up to eq. [11], e.g. the conductivity anomaly of MgO below =800°C is caused by O$^2-$ defects which generate mobile O$^-$. The increase of $F_\Delta$ indicates an increase of O$^-$ density at the surface, but O emission destroys O$^-$ charge carriers and, hence, decreases $F_\Delta$. Likewise the annihilation of O$^-$ must also lead to a decrease of d.c. conductivity.
Fig. 7a Comparison between $F_\Delta$ above 500°C of a 4N-grade MgO single crystal with the atomic O evolution from ultrahigh purity MgO powder suggesting the decomposition of surface O⁻ into O²⁻ plus O (after [Freund et al., 1991; Kathrein and Freund, 1983; Martens et al., 1976]).

In vacuum, and in the absence of transition metal cations, the only way to irreversibly annihilate O⁻ charge carriers is by O atom emission according to eq. [11]. Obviously, the O emission can only take place from the MgO surface. Therefore, reduced gases which are brought in contact with the surface and are oxidized by O⁻ will have the same effect: they irreversibly annihilate O⁻ charge carriers. The gases of interest are H₂ and CO, and their “cleansing action” can be described in simple chemical terms as:

$$O^- + O^- + H_2 = O^{2-} + H_2O \uparrow$$  \[12a\]

$$O^- + O^- + CO = O^{2-} + CO_2 \uparrow$$  \[12b\]
Fig. 7b Comparison between the d.c. conductivity anomaly above 500°C of a 4N-grade MgO single crystal with the atomic O evolution from ultrahigh purity MgO powder suggesting the decomposition of surface $O^-$ into $O^2-$ plus O (after [Freund et al., 1991; Kathrein and Freund, 1983; Martens et al., 1976]).

Conclusion

There are three important conclusions to be drawn from the data presented here and in Part I [Freund et al., 1991]. The first conclusion pertains to the cause of the anomalous electrical conductivity of MgO crystals below $\sim$800°C. The cause is not extraneous as has been widely assumed until now but indigenous, pointing at defects or impurities that exist in the bulk of the crystals under study. The mobile charge carriers have been recognized as positive holes, equivalent to defect electrons on the $O^2-$ sublattice, chemically $O^-$ states. At low temperatures, below $\sim$300°C, they are localized, spin-paired and diamagnetic, probably consisting of $O^2_2$ defects at cation vacancy sites. Above 500°C the $O^2_2$ defects dissociate and release delocalized, highly mobile $O^-$ into the $O^2_2$ sublattice. The fact that these $O^-$ have a strong propensity to come to the surface must not be mistaken for a sign of surface contamination. This raises the possibility
that a similar anomalous conductivity behavior observed in structurally and compositionally more complex crystals such as mantle-derived olivine may also have an indigenous cause.

The second conclusion pertains to the origin of the \( \text{O}^- \) charge carriers. The IR data presented here strongly support the contention that \( \text{OH}^- \)-compensated \( \text{Mg}^{2+} \) vacancies are sites where \( \text{OH}^- \) pairs undergo an internal redox reaction and become converted into molecular \( \text{H}_2 \) plus \( \text{O}_2^- \). This reaction consumes lattice-bound \( \text{OH}^- \) which are obvious signs of dissolved \( \text{H}_2\text{O} \) component. Therefore, the \( \text{OH}^- \) concentration that can be measured by IR spectroscopy is not representative of the total \( \text{H}_2\text{O} \) content. If \( \text{H}_2 \) molecules diffuse out, the only "memory" of the earlier \( \text{H}_2\text{O} \) content consists of \( \text{O}_2^- \) defects. In the case of the \( \text{MgO} \) crystals studied here the total amount of \( \text{H}_2\text{O} \) incorporated during crystal growth appears to have been significantly larger (by a factor 5-10) than the current \( \text{OH}^- \) content suggests*. This suggests that the conjecture \( \text{OH}^- \) content equal dissolved \( \text{H}_2\text{O} \) content, hitherto taken for granted in all discussions concerning dissolved "water" in minerals, may need revision.

The third conclusion has potentially even more far-reaching implications. It pertains to the overall redox state of the \( \text{MgO} \) crystals under study. The single crystals were grown in the chemically reducing environment of a carbon arc furnace where the atmosphere is dominated by \( \text{CO}/\text{CO}_2 \) with traces of \( \text{H}_2\text{O} \) [Abraham et al., 1971; Butler et al., 1971]. The powder was obtained by dehydrating \( \text{Mg(OH)}_2 \) in the intrinsically reducing environment of very high vacuum. In both cases the defect stoichiometry of nominally high purity, transition metal free crystals should be characterized by oxygen vacancies, equivalent to an excess of \( \text{Mg} \) [Yamaka and Sawamoto, 1954; Mitoff, 1959; Osburn and Vest, 1971]. Instead our experimental data indicate that the \( \text{MgO} \) contains excess \( \text{O} \) in form of peroxy defects and have the approximate composition \( \text{MgO}_{1.58} \). Being equivalent to interstitial \( \text{O} \) atoms peroxy defects are the epitome of highly oxidizing conditions [Wriedt, 1987]. The amount of excess \( \text{O} \) appears to be substantial, of the order of 1000 ppm in the arc fusion grown single crystals and 2000 ppm in \( \text{MgO} \) powder obtained from \( \text{Mg(OH)}_2 \).

Fig. 8 shows an isobaric section through the three-component phase diagram \( \text{Mg-O}_2-\text{H}_2 \). The diagram is divided into a reduced region delineated by \( \text{Mg}-\text{H}_2-\text{H}_2\text{O}-\text{MgO} \) and an oxidized region delineated by \( \text{MgO}-\text{H}_2\text{O}-\text{O}_2 \). Also included are compounds \( \text{MgH}_2, \text{H}_2\text{O}_2 \) and \( \text{MgO}_2 \). \( \text{MgO} \) straddles the boundary between the reduced and oxidized phase regions. As shown schematically in the enlarged portion below \( \text{MgO} \) is oxygen-deficient, if heated under low \( \text{O}_2 \) partial pressures [Mitoff, 1959; Osburn and Vest, 1971] or in the presence of \( \text{Mg} \) vapor [Yamaka and Sawamoto, 1954]. It is oxygen-rich, if heated in \( \text{O}_2 \), equivalent to an \( \text{MgO-MgO}_2 \) solid solution [Wriedt, 1987]. The phase region extends towards \( \text{Mg(OH)}_2 \), due to the known solubility of \( \text{H}_2\text{O} \) component. Though the extent of isobaric solid solution formation along the three axes is unknown the shaded area outlines a probable scenario. The upper open circle is representative for non-stoichiometric \( \text{MgO} \) as expected to crystallize under reducing conditions (excess \( \text{H}_2 \) or \( \text{Mg} \)) in the presence of \( \text{H}_2\text{O} \). The line joining \( \text{MgO-Mg(OH)}_2 \)

* Assuming that \( \text{H}_2\text{O} \) is the only gas the dissolution of which leads to peroxy defects.
represents compositions $\text{Mg}_{1-\delta}\text{O}_{1-\varepsilon}(\text{OH})_{2\theta}$. The bold straight arrow pointing downwards delineates the path which the system $\text{Mg}_{1-\delta}\text{O}_{1-\varepsilon}(\text{OH})_{2\theta}$ would take after internal conversion of $\text{OH}^-$ into $\text{H}_2$ plus $\text{O}_2^2$ by loosing $\text{H}_2$. The endpoint lies on the abscissa and represents the "water–free" composition $\text{Mg}_{1-\delta}\text{O}_{1-\varepsilon}(\text{O}_2)_{\theta} = \text{MgO}_{1+\delta}$.

Fig. 8 Mg–O2–H2 phase diagram with the non–stoichiometry region for MgO redrawn and enlarged in the lower half (schematic). The shaded area outlines the compositional variability of MgO crystallizing in the presence of H2O under reduced and oxidized conditions. The open circle in the shaded area represents $\text{Mg}_{1-\delta}\text{O}_{1-\varepsilon}(\text{OH})_{2\theta} \leftrightarrow \text{Mg}_{1-\delta}\text{O}_{1-\varepsilon}(\text{O}_2)_{\theta}(\text{H}_2)_{\delta}$ obtained under reducing conditions with conversion of $\text{OH}^-$ to $\text{H}_2$ plus $\text{O}_2^2$. The open circle on the abscissa represents the composition $\text{Mg}_{1-\delta}\text{O} = \text{MgO}_{1+\delta}$ which would be achieved by complete conversion of $\text{OH}^-$ and loss of $\text{H}_2$.

Fig. 8 illustrates that, while the representative point for the initial non–stoichiometric MgO lies in the "reduced" region, it can cross the dividing line between "reduced" and "oxidized" and settle in the "oxidized" region. It needs to be stressed is that such a transition would be achieved without exposing MgO to oxidizing conditions but solely by way of the internal conversion reaction discussed above, $2 \text{OH}^- \leftrightarrow \text{H}_2 + \text{O}_2^2$, followed by out–diffusion of $\text{H}_2$. If the isobaric solubility for H2O is larger than the solubility for O2, the excess O in the MgO structure in form of peroxo...
defects can exceed the solubility limit for O₂ at the same pressure. This possibility is illustrated in Fig. 8 by placing the representative point for MgO₁⁺₆ outside the shaded region.

So far MgO has been used as a model substance to study this internal reaction sequence. Chemically and structurally more complex minerals such as olivine do not lend themselves to some of the techniques applied here. However, given the fundamental nature of the reaction steps discussed above, it cannot be ruled out that traces of dissolved H₂O component in olivine and other minerals will undergo a similar internal conversion, followed by out-diffusion of H₂. The petrological consequences of such a process could reach far beyond the concern about electrical conductivity behavior emphasized here. Until now it has been common practice to judge the redox state of minerals from their transition metal equilibria as reflected, for instance, by the Fe²⁺/Fe³⁺ ratio measured at 20°C. This Fe²⁺/Fe³⁺ ratio is then used to derive the conditions under which these minerals have formed. The results presented here show that the Fe²⁺/Fe³⁺ ratio is not a unique, not even a reliable indicator of the internal redox state. Crystals that originally formed under highly reducing conditions may acquire the sigma of an oxidized condition without concomitant adjustment of the ratio of Fe²⁺/Fe³⁺ and other transition metal cations.

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References


Electrical Conductivity Measurements under Minimum Perturbation Conditions

Part III:
Peroxy in Olivine

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Abstract

Mantle-derived olivine crystals exhibit an anomalously high electric conductivity up to \( \approx 700^\circ \text{C} \) when heated for the first time. This anomaly is commonly believed to be a "dirt effect", due to extraneous surface contamination, to polishing damage of the surface or to vapor-deposited amorphous carbon in the stability field of graphite. Using a new technique, Charge Distribution Analysis or CDA which measures contact-free the dielectric polarization in an electric field gradient at 0 Hz, the conductivity anomaly can be shown to be correlate with the appearance of highly mobile, positive charge carriers that are reversibly generated in the bulk between 450-550°C. They give rise to a surface charge, supported by an internal electric field which disappears upon cooling. The positive sign of the charge carriers suggests that they consist of holes. However, they are not holes on the cation sublattice, e.g. transient Fe\(^{3+}\) states due to oxidation, but holes on the \( \text{O}^{2-} \) sublattice, e.g. \( \text{O}^- \) states. Their reversible appearance during heating/cooling cycles suggests that the \( \text{O}^- \) states are generated from indigenous peroxy defects, probably peroxy links, \( \text{O}_3\text{Si}/\text{O}^0\text{SiO}_3 \), which coexist with Fe\(^{2+}\) in the olivine structure. The peroxy defects dissociate thermally between 450-550°C, causing the observed conductivity anomaly. The \( \text{O}^- \) charge carriers diffuse to the surface, causing the positive surface charge. At the same time they are susceptible to oxidation by reduced gases at \( \geq 700^\circ \text{C} \), e.g. \( \text{CO} + 2 \text{O}^- = \text{CO}_2 + \text{O}^2^- \). At the same time the \( \text{O}^- \) can internally oxidize Fe\(^{2+}\) to Fe\(^{3+}\) as if the olivine became oxidized by traces of \( \text{O}_2 \) in the ambient atmosphere. By analogy to MgO where cation vacancy bound OH\(^-\) pairs undergo an internal redox reaction to \( \text{H}_2 \) plus \( \text{O}_2^2^- \), it is proposed that the peroxy defects in olivine derive from traces of dissolved fluid phase components, notably \( \text{H}_2\text{O} \), and that the \( \text{O}^- \) charge carriers are an integral part of the defect structure which the olivine crystals acquire in the upper mantle.

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Introduction

Olivine is by far the major mineral of upper mantle rocks and therefore continues to occupy a central place in geophysical research. Understanding its defect structure and the role of impurities is essential to understanding the behavior of rocks under upper mantle conditions. Given the tremendous amount of effort that has been devoted to the experimental study of olivine from a variety of standpoints it appears unlikely that anything fundamentally new may remain to be discovered. Yet, in this paper we shall present evidence that olivine crystals from upper mantle rocks contain some intriguing charge carriers, highly mobile and evasive, which have resisted recognition until now.

The electrical conductivity of olivine has been measured with great care, both at ambient pressure (0.1 MPa) and at high pressures, under a wide range of oxygen fugacities [Constable and Duba, 1990, Duba et al., 1974, Duba and Nicholls, 1973, Duba et al., 1988, Schock et al., 1977, Schock et al., 1989, Shankland, 1981, Sockel, 1974]. As a result of these studies various charge carriers and conductivity mechanisms have been identified which influence or control the behavior of olivine in the high temperature (HT) region, above approximately 800°C. At the same time these studies have also laid open a striking anomaly in the conductivity behavior in the lower temperature (LT) region, below approximately 800°C. Fig. 1 in Part I [Freund et al., 1991] contained typical electrical conductivity data of olivine and olivine–rich rocks at oxygen fugacities of the upper mantle, all obtained at 0.1 MPa pressure. In the HT region, above =800°C, Arrhenius plots with constant slopes suggest a simple relation between conductivity $\sigma$ and absolute temperature $T$: $\sigma_{ij} = \sum_{ij} \sigma_0 \exp[-E/kT]$, where $\sigma_0$ are pre-exponential factors and $E$ activation energies for the charge carriers $i,j$, while $k$ is the Boltzmann constant. In the LT region, below =800°C, the conductivity behavior was more complex. Each time an olivine sample is heated for the first time there appears to be a peculiar enhancement of the conductivity, often by several orders of magnitude. This anomalous behavior must have annoyed many researchers. It is reportedly irreproducible in the sense that the high conductivity wanes during extended treatment in reducing CO/CO$_2$ gas atmospheres at temperatures around 700°C and disappears above 800°C. The waning and eventual disappearance has led to the belief that the high conductivity is a “dirt effect”, probably caused by surface contamination during sample handling, by structural disorder introduced into the surface layer through the sample preparation, for instance during polishing, or by the deposition of a thin layer of amorphous carbon, vapor–deposited from the CO/CO$_2$ atmosphere by way of the disproportionation reaction $CO_2 = C + O_2$ in the stability field of graphite [Constable and Duba, 1990].

As illustrated in Fig. 2 of Part I the same electrical conductivity anomaly occurs when cleaved, synthetic MgO single crystals are heated in a pure Ar [Freund et al., 1991]. Therefore, disproportionation of CO$_2$ can be dismissed as the culprit. By judiciously selecting the right experimental conditions, the anomaly can be shown to be quite reproducible. The remainder of Part I was dedicated to describing a new technique Charge Distribution Analysis or CDA which has made it possible to show that, far from being erratic, the on–set of the electrical conductivity
anomaly in MgO and the stepwise increase of the static dielectric constant correlates with the appearance of a positive surface charge, supported by an internal electric field. These surface charges and their internal support field cannot be explained by mere surface contamination. Instead they require indigenous charge carriers that originate in the bulk – charge carriers which appear to have "fallen through the cracks" of prior investigations.

This brash conclusion was underscored in Part II by presenting results from several studies of MgO obtained with independent techniques, selected to illuminate different aspects of the enigmatic charge carriers which appear to be associated with the anomalous conductivity [Freund, 1991]. At the end of Part II a strong case could be made that the charge carriers in question are electronic defects on the O\textsuperscript{2−} sublattice, commonly referred to as positive holes or O\textsuperscript{−} states.

From a solid state physics viewpoint O\textsuperscript{−} are defect electrons in wide band gap insulating oxides. They occupy energy levels near the O 2p-dominated valence band. If their wave functions spread out over a large lattice volume, causing large polarons, the O\textsuperscript{−} may become highly mobile charges of positive sign [Freund et al., 1989]. They carry electric current without mass transport. If the O\textsuperscript{−} are localized, forming small polarons, they are electrically inactive though they may still be detected because of their unpaired spin. From a solid state chemistry viewpoint O\textsuperscript{−} states are radicals, paramagnetic and intrinsically unstable [Henderson and Wertz, 1977, Marfunin, 1979]. They are oxidants which may oxidize transition metal cations from low to higher oxidation states. If two O\textsuperscript{−} undergo self-trapping and pair their spins, they become invisible to all known physical techniques. They form dimers, O\textsubscript{2}\textsuperscript{2−}, conveniently called peroxy entities, they become diamagnetic and non-reactive. Formally an O\textsubscript{2} corresponds to an interstitial oxygen atom, O\textsubscript{2} = O\textsuperscript{2−} + O.

In Part III we return to olivine – a mineral which is more complicated than MgO, both structurally and chemically. The striking similarity between olivine from the upper mantle and synthetic MgO crystals in their respective anomalous conductivity behavior below 700–800°C raises an interesting question. After having corroborated by independent techniques that the conductivity anomaly in MgO is not a “dirt effect” but caused by indigenous O\textsuperscript{−}, we must be open to the possibility that the same enigmatic charge carriers exist in olivine. Unfortunately, because of their high Fe\textsuperscript{2+} content, olivine crystals from the upper mantle do not lend themselves to some of the powerful techniques presented in Part II through which we could unambiguously identify the charge carriers in MgO as O\textsuperscript{−} states [Freund, 1991]. For olivine Charge Distribution Analysis (CDA) remains an option. With its unique capability to study electrical and dielectric properties under minimum perturbation conditions we can learn something about the charge carriers of olivine which control the anomalous conductivity behavior.
Experimental

The CDA experiments were performed in an asymmetrical capacitor arrangement using two similar PERKIN–ELMER Thermogravimetric Analyzers TSG–1 and TGS–2 with a CAHN and a PERKIN–ELMER microbalance respectively. The TGS–1 had a small furnace, Pt wire wound on a cup–shaped alumina ceramic body, 6 mm internal diameter and 12 mm high, depicted in Fig. 1, while the furnace of the TGS–2 measured 10 mm internal diameter and 18 mm high. The Pt winding was bifilar and the furnace was a.c. operated using one half period of the sine wave for power and the other for measuring the resistance and, hence, the temperature. The furnace was surrounded by a 0.1 mm Au foil serving to equalize temperature gradients and as the counterelectrode. The sample was placed on a loop made of 0.1 mm diameter fused silica, suspended from the microbalance via a fused silica wire. The bias electrode was approximately circular, 4 mm diameter of Ni or Au, and introduced laterally above the sample. The critical sample–bias electrode distance could not be accurately controlled but was adjusted to 0.2–0.3 mm. Bias electrode and counterelectrode were connected to the ± output and ground respectively, of a KEITHLEY 240A digital voltage source (±1100V). The contribution from the fused silica support loop to the force measurement under CDA conditions was considered negligible.

All CDA measurements were performed at 0.1 MPa pressure in high purity 20–40 ml/min high purity N₂ (99.99 %, <5 vppm O₂, <10 vppm H₂O) during stepwise heating and cooling programs to be detailed below. Heating and cooling between incremental and decremental steps were usually fast, 20 or 40°C/min as noted in each case, in order to minimize exposure to the potentially detrimental effects of the unbuffered N₂ atmosphere. For some test runs slower heating and cooling rates were used, 1-10°C/min. The samples were allowed to thermally equilibrate at each temperature step prior to the application of the bias voltages. An equilibrium time of about 5 min was generally found sufficient. The ± bias voltages were applied in preselected increments, usually in 10–20 V.
It should be emphasized that, from the outset, the CDA runs with olivine were not intended to simulate the temperature, pressure and oxygen fugacity conditions appropriate for the upper mantle but to duplicate our previous experiments with MgO single crystals as described in Part I. The purpose of the study was to collect as much information about olivine as possible in the temperature interval where the electrical conductivity anomaly is most pronounced, e.g. below 800°C. We exerted no control of the oxygen partial pressure but instead chose the same high purity N₂ gas as in the MgO runs. Though the stated <5 vppm O₂ content of N₂ atmosphere lies outside the thermodynamic stability range of olivine, even if oxidation would take place, it will be kinetically hindered at the relatively low temperatures. The heating times were kept to a minimum, typically 30–45 min, and heating never exceeded 800°C. Tests were performed as described below to assess the reversibility of the observed CDA response. As long as the CDA response was reversible, it was assumed that the olivine crystals represented a closed system with no detectable oxidation by the traces of O₂ in the unbuffered N₂ atmosphere. The chance of oxidation was also kept to a minimum by the fact that the single crystals used in this study provided a minimum surface area across which oxidation could occur. We noted that crystals heated for 2 hrs or more to 750–800°C tended to acquire a brownish surface hue, indicative of incipient oxidation [Iishi, 1989].
Olivine single crystals from San Carlos, Arizona, USA, were available as large rounded pebbles, optically slightly turbid. Thin sections revealed a polygonized substructure of misoriented subgrains separated by dislocation walls, indicative of a high degree of disorder and/or mechanical deformation in the mantle. Occasionally small veinlets were filled with finely divided matter, probably due to metasomatic alternation. Microscopic inspection of two thin sections of random orientation revealed no visible fluid inclusions though the possibility remains that occasional fluid inclusions were missed during the survey [Pasteris and Wanamaker, 1988, Roedder, 1984]. No infrared (IR) signals characteristic for fluid or gaseous CO or CO₂ were observed. A large single crystal measuring 5.5 x 8.6 x 9.5 mm was cut and polished on all sides. The IR absorption spectrum recorded through its 8.6 mm thickness showed no detectable VOH absorption band in the 3700-3300 cm⁻¹ range. This indicates a vanishingly small concentration of residual OH⁻ impurities – a result that is not inconsistent with the wide OH⁻ concentration range down to nil as reported by Miller et al. [Miller et al., 1987] for many olivine crystals from different locations and definitely less than the ~20 at.ppm OH⁻ determined by Mackwell and Kohlstedt [Mackwell and Kohlstedt, 1990] in a hydrothermally treated olivine single crystal from the same San Carlos locality.

The 5.5 x 8.6 x 9.5 mm single crystal was cut into eight pieces, each measuring 4.5 x 4 x 2.5 mm, using a thin–blade, water–cooled, slowing rotating diamond saw. The edges were rounded off on SiC sanding paper to approximate a somewhat cylindrical shape. The pieces were cleaned in methanol or chlorinated hydrocarbon solvents and in distilled water. No traces of solvent residues could be detected by IR after drying. The crystals were placed on the fused silica loop with their polished sides facing the bias electrode. The effect of reversing the samples and placing them on the fused silica loop with the rough sides towards the bias electrode was not specifically studied, but it is known both from theory (see Part I) and prior experience with irregularly shaped samples [Freund et al., 1990] that the force measured is a function of the volume integral over the region containing the sample. Hence, the measurements are position–sensitive only to the extent that the volume integral changes.

Results

Our first task was to carry out a careful reversibility test up to 600°C. During experiment #j4 the following heating/cooling protocol was adhered to: (i) a fresh olivine crystal was loaded on the fused silica loop and dried in situ at 120°C for 1 hr; (ii) the sample was cooled to ambient temperature (24°C) and then reheated at a rate of 20°C/min to 100°C and on to 600°C in 100°C increments with approximately 30 min hold time at each temperature step during which plus and minus bias voltages were applied sequentially in 20 V increments up to ±100 V; (ii) the sample was cooled in one sweep to 400°C at a cooling rate of 20°C/min and the ± bias voltages were applied as before; (iii) the sample was reheated to 600°C at 20°C/min; (iv) the sample was cooled at 20°C/min from 600°C in one sweep to 300°C; (v) the sample was reheated a third time to 600°C.
at 20°C/min; (vi) the sample was cooled to 200°C and then to 100°C, interrupted by application of the ± bias voltages; (vii) the sample was heated a fourth time to 600°C at 20°C/min; and finally, (viii) the sample was cooled to 100°C at 20°C/min. Total duration of run #j4 was >10 hrs with ~2 hrs accumulative exposure time to 600°C.

Fig. 2: F⁺ and F⁻ as a function of voltage, measured during four consecutive heating/cooling cycles up to 600°C to demonstrate the reproducibility of the CDA measurements. 1st cycle: circles and dashed lines; 2nd cycle: squares; 3rd cycle: crosses; 4th cycle: triangles. The labels refer to the sequence of the ±100 V data.

In Fig. 2 all F⁺ and F⁻ values obtained during exp. #j4 are compiled in a force versus voltage plot. The ordinate is in arbitrary units. As expected from the CDA theory presented in Part I the response to increasing bias voltages gives rise to a series of approximately parabolic curves. If
the olivine crystal behaved as an ideal dielectric with a static dielectric constant $\varepsilon_{\text{static}}$ that is essentially temperature-independent [Cygan and Lasaga, 1986], the response would be exactly parabolic, $F^\pm \propto U^2$, and all curves would coincide. The variable steepness of the parabolas reflects a variations in $\varepsilon_{\text{static}}(T)$. Deviations from the exact parabolic shape reflect field-variant contributions, $P_{\text{surface}}$, due to surface charges. The surface charges are very sensitive to the cooling rate and cause some significant changes in the $F^+$ and $F^-$ values between 100–500°C. The excellent reproducibility (interpreted as absence of irreversible oxidation) is demonstrated by the fact that even after four heating/cooling cycles the $F^\pm$ values at 600°C agreed within $\pm 5\%$.

In Fig. 3 the same $F^\pm$ values are used to calculate $F_\Sigma$, e.g. the field-invariant part of the dielectric response which is proportional to $\varepsilon_{\text{static}}$. These $F_\Sigma$ values are plotted as a function of $T$ for the four consecutive heating/cooling cycles*. At all temperatures except at 400°C where the reproducibility lies within $\pm 10\%$ the reproducibility is better than $\pm 5\%$, indicating that the surface contributions were self-cancelling for $F^+$ and $F^-$. At 600°C the $F_\Sigma$ values coincide nearly perfectly. This is a strong indication that the olivine underwent no noticeable chemical changes during the heating and cooling cycles due to oxidation. This lends credence to the presumption that the olivine crystals under study can be safely heated repeatedly to at least 600°C without suffering oxidation, even though the ambient N$_2$ atmosphere was unbuffered. Other experiments similar to #j4 but not shown in detail here were extended to temperatures above 600°C. They indicated a progressive deterioration of the reproducibility above 700°C, suggesting the beginning of oxidation.

* Due to an internal normalization routine the ordinate force units are different than those of Fig. 2.
Fig. 3: $F_\Sigma$ for the four consecutive heating/cooling cycles plotted as a function of temperature. The $F_\Sigma$ values are reproducible within better than ±5% at 600°C, indicating that the sample did not suffer irreversible changes due to oxidation.

According to Fig. 3, during the first heating run, $F_\Sigma$ was slightly high at 24, 100 and 200°C, lower between 300–500°C and increased again between 400–600°C. Upon cooling the shape of the $F_\Sigma(T)$ curve was reproduced, including the enhanced values at 200 and 100°C**. Most noteworthy is the stepwise $F_\Sigma$ increase between 450-600°C which was fully reversible within ±5%, even at cooling rates as fast as 20°C/min. Since $\varepsilon_{\text{static}}$ of an ideal dielectric must be essentially temperature-independent [Ashcroft and Mermin, 1976] this stepwise $F_\Sigma$ increase

** In other experiments with slower cooling rates no enhancement of the $F_\Sigma$ values upon cooling below 300–400°C were observed, indicating that the lower temperature portion of the $F_\Sigma(T)$ curves shown in Fig. 3 is affected by the rapid cooling. The reason is that some mobile charge carriers generated at 600°C are easily frozen upon rapid cooling, causing residual polarization.
signals the appearance of mobile charge carriers which increase the dielectric polarizability of the olivine through their $P_{\text{space}}$ contribution. The reproducibility suggests (i) that during reheating approximately the same number of mobile charge carriers is generated and (ii) that the charge carriers disappear upon cooling or become immobilized through a trapping process. The $F_\Sigma(T)$ reproducibility remained within ±10% limits upon heating to 700°C and deteriorated slightly after exposing the olivine crystals to temperatures above 700°C for extended periods of time, typically 2–4 hrs.

**Fig. 4a:** $F^+$ as a function of temperature during stepwise heating to 800°C.
After having gained confidence that heating in the unbuffered N₂ atmosphere was benign up to 600°C, acceptable to 700°C and even higher, provided the time of exposure was kept short, we performed stepwise heating experiments up to 800°C without intermittent cooling. The experiments were repeated in nearly the identical way, using a fresh sample. **Fig. 4a/b** shows the result of the most complete run of the series, experiment #h4/5, featuring 15 temperature steps. The sample was first dried *in situ* at 120°C, cooled to 24°C and heated in 100°C steps up to 400°C, in 50°C steps from 400-500°C, in 25°C steps from 500-600°C and to 800°C in 50°C steps, the heating rate between steps always being 40°C/min. After holding the temperature for 5 min the bias voltages were applied in 5 V increments up to ±30 V and in 10 V increments to ±100 V.
The $F^\pm$ values were measured continuously but the values used for Fig. 4a/c represent those 2 min after application of the bias voltages. Since each temperature step required ~60 min to complete the sequence of $F^+$ and $F^-$ measurements, the total duration of run #h4/5 was >18 hrs with >6 hrs exposure to temperatures >600°C. After completing the heating cycle the sample was cooled fast (40°C/min) from 800–600°C and slowly (2°C/min) from 600–24°C while maintaining the flow of N$_2$. Thereafter the $F^\pm$ values were remeasured and were found to agree with those of the first heating run to within approximately ±20%, except below 150°C where the agreement was less than 50%.

![Graph](https://example.com/graph.png)

**Fig. 4c:** $F_\Sigma$ as a function of temperature during stepwise heating to 800°C.

As shown in Fig. 4a/b $F^+$ and $F^-$ were nearly constant from 24–450°C but rose sharply between 450–600°C. $F^-$ continued to rise between 600–800°C, while $F^+$ fell off slightly. $F_\Sigma(T)$, depicted in Fig. 4c, shows a stepwise increase between 450–600°C, similar to the $F_\Sigma$ increase in MgO in the same temperature interval (see Part I). Such a stepwise increase can only be caused by the
appearance of mobile charge carriers that are capable of diffusing in the externally applied electric field gradient. Above 600°C $F_\Sigma$ tended to level out, possibly indicating saturation of charge carriers. This tendency of $F_\Sigma$ to reach a plateau was repeatedly observed in similar experiments, sometimes even a trend towards a decrease, but upon prolonged heating $F_\Sigma(T)$ continued to rise moderately from 700–800°C as shown in Fig. 4c.

\[ F_\Sigma(T) \text{ is depicted in Fig. 4d. We recall that } F_\Delta \text{ reflects the field-variant contribution, due to } P_{\text{surface}} \text{, and that its sign gives the sign of the surface charge layer. Below 450°C the surface charge of fresh, previously unheated olivine crystals was weakly but consistently negative or near-zero. Between 450-575°C } F_\Delta \text{ always changed sharply to positive and remained positive up to 800°C. The minimum in the } F_\Delta(T) \text{ curve in Fig. 4d between 600–650°C may be a artefact of the data acquisition routine, caused by different response times of the } F^+ \text{ and } F^- \text{ after application of the bias voltage. If } F^+ \text{ and } F^- \text{ values recorded 5 or 10 min after application of the bias voltage were used to calculate } F_\Delta, \text{ the amplitude of the minimum would diminish. We did not} \]
observe a decrease of $F_\Delta$ above 750°C similar to that reported for MgO (see Part I). Through repetitive heating/cooling cycles it was ascertained that $F_\Delta$ always tended to be slightly negative or zero between 100–450°C and always became strongly positive above 450–550°C. Upon cooling at <10°C/min the positive $F_\Delta$ values reverted to near zero or even became negative again. Upon quenching at fast cooling rates, >20°C/min, the $F_\Delta$ values remained positive as mentioned above. If heating was carried out beyond 700°C, the reversibility was lost and $F_\Delta$ remained positive.

Next we conducted an experiment to demonstrate the internal electric field associated with the positive surface charge above 500°C. In Figs. 5a and 5b $F^+$ values obtained during run #H3 are plotted as a function of $U$ and $U^2$ respectively. (For the sake of clarity not all data contained in Fig. 5a are replotted in Fig. 5b.) The sample was pretreated in a similar way as for run #h4/5, e.g. dried in situ for a few hours around 100°C. After this there was essentially zero surface charge. The sample was then heated in 50°C or 25°C increments to 800°C and positive bias voltages were applied in 2 V or 4 V increments to +20 V, ending with +25 V. All $F^+$ used for these plots were obtained ≤2 min after application of the bias voltages. The $F^+(U)$ and $F^+(U^2)$ curves for the temperature interval 100–500°C are represented by the dotted bold lines. They are nearly perfect parabolas, passing through the origin. All $F^+$ values within this range were attractive, indicating an essentially zero surface charge and confirming the observations of run #h4/5. The $F^+(U)$ curves obtained between 550–650°C were no longer simple parabolas passing through the origin. Their linearly extrapolated $F^+$ vs $U^2$ graphs shown in Fig. 5b intersect the ordinate at negative values. Between 675–800°C the $F^+$ became clearly repulsive at low bias voltages, indicated by negative force values. Their linearly extrapolated $F^+$ vs $U^2$ graphs intersect the ordinate at ever larger negative values.
**Fig. 5a:** $F^+$ measured under low positive bias during stepwise heating plotted as a function of $U$. The dotted lines represent data up to 500°C, near-perfect parabolas passing through the origin.
Fig. 5b: $F^+$ measured under low positive bias during stepwise heating plotted as a function of $U^2$. The dotted lines are the extrapolated parabolic components measured above 550°C. Some data from (a) are omitted in (b) for the sake of clarity.

We deconvoluted the $F^+$ values of Fig. 5a/b according to $F^+ = F_0^+ + aU^2$ (see eq. [5] in Part I) where $aU^2$ describes the ideal dielectric and $F_0^+$ arises from the charge at the sample surface. The deconvoluted curves are shown in Fig. 6. Up to 500°C $F_0^+ = 0$, indicating essentially zero surface charge as already mentioned. Between 550–800°C the positive surface charge gives rise to a repulsive term (marked in the lower half of Fig. 6 by bold lines). $U_{\text{crit}}$ as marked by the short vertical ticks indicates the critical voltage up to which the surface charge withstands the externally applied bias voltage. In other words it is the externally applied electric field up to which the surface charge acts as an electrostatic mirror. Above $U_{\text{crit}}$ the surface charge is pushed back into the bulk. When the external bias is removed, the positive charge returns to the surface within a very short time, typically $<$10 sec at 600–650°C. The to-and–from migration of charges between surface and bulk is a fully reversible process as long as no oxidation takes place. In
keeping with the qualitative nature of the measurements presented here the force values in Fig. 6 are given in arbitrary units. Theoretically it can be shown that \( |F_0^+| \) is proportional to the surface charge density \[Freund et al., 1989\]. In MgO single crystals, assuming a surface layer thickness of 15 nm, the charge carrier density was estimated to be of the order of \(10^{21} \text{ cm}^{-3}\). No attempt was made with the present CDA set-up to quantify the surface charge density data for the olivine crystals.

![Graph showing San Carlos Olivine: \( F^+ \)](image)

**Fig. 6:** Same data as shown in Fig. 5a, deconvoluted according to \( F^+ = F_0^+ + aU^2 \).

In closing the results section we note that the data presented in Figs. 2–6 comprise only a fraction of the experiments which we carried out with olivine to ascertain the reproducibility of the phenomena to a degree of certainty comparable to that of the CDA study MgO as described in Part I \[Freund et al., 1991\]. We also carried out a preliminary experiment during which we added about 5 vol.-% \( H_2 \) for 10-15 min to the \( N_2 \) stream close to \( 700^\circ C \). We observed an immediate decrease of the \( F^+ \) and \( F^- \) response. The rapidity of the changes indicated that a surface reaction had taken place. After the \( H_2 \) pulse \( F^+ \) and \( F^- \) did not fully recover, indicating an at least in part
irreversible reactions. A more complete study of the effects of H₂ and other reduced gases on the
CDA response of olivine will be presented in a separate report.

Discussion

The main advantage of the CDA technique is that, by measuring contact-free under minimum
perturbation conditions, it provides insights into charge generation and mobilization processes
which cannot be achieved by any other single method. By way of the dielectric polarization at 0
Hz CDA detects mobile charge carriers without external carrier injection or extraction. In
particular, CDA has the capability (i) of detecting the appearance and disappearance of charges in
the bulk, (ii) of detecting the appearance and disappearance of surface charges, and (iii) of
determining the sign and strength of the internal fields which support the surface charges.

The CDA data of olivine presented in this first report are still qualitative. We have not yet
developed the mathematical tools to calculate the concentration of mobile charge carriers by finite
element analysis. However, the reality of the observed physical phenomenon cannot be denied.
The measurement of the dielectric polarization, contact-free in an electric field gradient of
reversible polarity at 0 Hz, is a sufficiently simple technique that a number of conclusions can be
drawn. The measurements clearly indicate that charge carriers are generated in the olivine crystals
upon heating to 450–550°C, that these charge carriers are highly mobile and positive, and that they
persist to higher temperatures, at least 700°C, possibly higher. They reversibly disappear upon
cooling. These charge carriers are not the product of extraneous contamination. The most cogent
argument against surface contamination stems from the fact that a surface charge develops which
is supported by a strong internal electric field and that this surface charge vanishes when the
mobile bulk carriers disappear. There is no physical process by which an internal electric field
could support an surface charge layer, unless the charges which appear at the surface originate in
the bulk of the crystal. The internal electric field signals that something inside the crystal “pushes”
the charge carriers to the surface and maintains them there, even against an externally applied
counterfield of considerable strength. Neither a layer of “dirt”, deposited onto the surface through
amorphous carbon vapor deposition, nor defects introduced into the surface through polishing and
other structurally damaging treatments can explain the presence of such an internal field.

The existence of surface charge layers on predominantly ionic crystals has long been predicted on
thermodynamic grounds [Kliewer and Koehler, 1965, Lehovec, 1953]. They are a consequence
of the fact that the free energies of formation for different families of point defects differ slightly at
the surface and in the bulk. As a result, intrinsic defects such as cation and anion vacancies are
present in different stoichiometric proportions near the surface than in the bulk. This creates
charge imbalance even under thermodynamic equilibrium conditions. The surface charges can be
exacerbated by the extrinsic defects and impurities, in particular by the segregation of aliovalent
impurities. Such a segregation will always occur when the impurity concentration in the bulk
exceeds the solubility limit [Kingery, 1974a/b]. However, the appearance and disappearance of
the positive surface charge in olivine between 450–550°C does not fall into this category of
aliovalent impurity segregation. First, the temperature is too low and the reversible changes too
rapid for atomic or ionic diffusion as we know it. The observations rather suggest electronic
charges moving to and from the surface without attendant mass transport. Second, the surface
charge is supported by an internal field, even against an externally applied counterfield. Such an
internal field can only be generated, if the charges originate in the bulk and internal forces cause
them to diffuse to the surface.

Positive charge carriers have already been observed in olivine at 1300°C through thermoelectric
effect measurements in the stability region of olivine [Schock et al., 1989]. It was suggested that
these charge carriers consist of holes that are reversibly generated by the oxidation of Fe\(^{2+}\) to Fe\(^{3+}\)
via gas phase O\(_2\) according to the reaction:

\[
8 \text{Fe}_{\text{Mg}}^{2+} + 2 (O_2)_{\text{gas}} \leftrightarrow 8 \text{Fe}_{\text{Mg}}^{3+} + 4 \text{O}_2^{-} + 2 \text{V}_{\text{Mg}}^{\text{vac}} + \text{V}_{\text{Si}}^{\text{vac}} \tag{1}
\]

In keeping with the non-stoichiometry of olivine as a function of oxygen fugacity [Duba et al.,
1974, Duba and Nicholls, 1973], eq. [1] is written as a reversible reaction. It postulates that,
on oxidation, the concentration of Fe\(^{3+}\) increases together with the concentration of Mg\(^{2+}\), Fe\(^{2+}\)
and of Si\(^{4+}\) vacancies [Nakamura and Schmalzried, 1983, Sockel, 1974]. Eq. [1] has been
formulated for temperatures well above the temperature region which we have covered in our
CDA experiments. At 1300°C diffusional processes involving cation vacancies and oxygen
incorporation are most certainly activated, allowing the olivine structure to adjust to a changing
oxygen fugacity within short time by (Mg,Fe)\(^{2+}\) and Si\(^{4+}\) vacancy formation. It is quite possible
that the charge carriers responsible for the sign of the thermoelectric effect at 1300°C do indeed
consist of defect electrons on the cation sublattice as postulated by Schock et al. [1989], e.g. of
holes that form Fe\(^{3+}\) hopping from Fe\(^{2+}\) to Fe\(^{2+}\) site.

At 450-550°C, however, it is very unlikely that holes could be generated by the same processes as
described in eq. [1], e.g. by incorporating gas phase O\(_2\) into the bulk of mm-sized olivine single
crystals and concomitant formation of (Mg,Fe)\(^{2+}\) and Si\(^{4+}\) vacancies. Even if we assume for a
moment that Fe\(^{2+}\) would become oxidized despite kinetically unfavorable factors by O\(_2\) in the
unbuffered N\(_2\) atmosphere, and that (Mg,Fe)\(^{2+}\) and Si\(^{4+}\) vacancies would form in accordance to
eq. [1], we still have to explain the reversibility observed by CDA. Since the CDA experiments
were carried out in N\(_2\) containing >5 vppm O\(_2\), the ambient P\(_{O_2}\) was many orders of magnitude
above the oxygen fugacity prevailing in the upper mantle [Nitsan, 1974]. As a result any
oxidation of the olivine at 450–550°C, if it occurs through reaction with the gas phase O\(_2\), can
only be unidirectional. This means that each successive heating of the olivine crystals to 600°C
should have led to progressively more oxidation. Instead, as depicted in Fig. 2, the CDA data
show a nearly perfect reproducibility of the field–invariant part of the dielectric polarization,
\(F_{\Sigma}(T)\). Since \(F_{\Sigma}\) responds to the charge carrier concentration in the bulk, we conclude that the
number of holes generated within the crystal during repetitive cycling up to at least 600°C was
constant and that the holes cannot be due to Fe$^{2+}$ oxidation through traces of O$_2$ in the N$_2$ atmosphere.

If the appearance of hole-type charge carriers in olivine cannot be blamed on extraneous O$_2$, we are left with only one alternative: In the *as received* state, e.g. as delivered by the upper mantle and retrieved in the field, the olivine crystals must contain a hidden reservoir of hole-type charge carriers. One of the characteristic features of this reservoir is that, upon thermal activation through repetitive heating and cooling cycles, charge carriers are reversibly released and consumed. We therefore now direct our attention to solid state processes by which hole-type charge carriers may be reversibly generated. Since there exist no known structural or magnetic phase transitions in olivine between 450-600°C and no mixing/unmixing of the forsterite-fayalite solid solution [Brown, 1980], and since the (Mg,Fe)$^{2+}$ sublattice cannot be the source on the basis of the arguments just presented, the only alternative is that the hidden reservoir lies in the O$^{2-}$ sublattice. The hole-type charge carriers which become mobile are then, by definition [Henderson and Wertz, 1977, Marfunin, 1979], positive holes, e.g. O$^-$ states. They acquire their mobility not by hopping from cation to cation sites but by entering acceptor levels near the O$^{2-}$-dominated valence band.

During an earlier single crystal IR study of San Carlos olivine [Freund and Oberheuser, 1986] two observations were made which are relevant here. First, two groups of IR absorption bands due to OH$^-$ impurities were observed. With respect to their relative position and intensities they resembled the OH$^-$ impurity bands in MgO which had been assigned to single OH$^-$ and to OH$^-$ pairs at cation vacancy sites, [OH$^-$ V$_{Mg}^-$] and [OH$^-$ V$_{Mg}^-HO$]$^x$. As in MgO the bands assigned to OH$^-$ pairs had a lower intensity than the bands assigned to single OH$^-$ defects. This was in conflict with the expectation that OH$^-$ pair defects, associated with divalent cation vacancies, should be more abundant. Second, a weak, but distinct IR band was observed in the 4150 cm$^{-1}$ region which indicated the presence of H$_2$ molecules in the olivine structure. By analogy to MgO it was therefore suggested that the OH$^-$ pair defects in olivine had been used up by a redox reaction to form H$_2$ molecules plus peroxy entitles as OH$^-$ pair defects in MgO (see eqs. [3/4] in Part II).

While there is only one substitutional mode for H$_2$O into MgO, in the olivine structure H$_2$O may substitute for either (Mg,Fe)O or SiO$_2$ component. The substitution for SiO$_2$ is arguably improbable, given the fact that olivine crystals in upper mantle rocks is generally associated with SiO$_2$-richer phases such as pyroxenes. Therefore, they are more likely (Mg,Fe)O-deficient than SiO$_2$-deficient as had been suggested by Beran and Putnis [1983]. If dissolution of H$_2$O in olivine occurs by substituting for (Mg,Fe)O, the structural formula becomes:

$$\text{(Mg,Fe)$_2$SiO$_4$} + \delta \text{H}_2\text{O} \leftrightarrow \text{(Mg,Fe)$_2$SiO$_4$-$8\text{SiO}_2$} + \delta \text{(Mg,Fe)O} \quad [2]$$

If OH$^-$ pairs in olivine, like their equivalent in MgO, underwent a redox conversion to H$_2$ plus peroxy and if these H$_2$ molecules subsequently diffused out of the olivine crystals, we have:
(Mg,Fe)$_{2-\delta}$SiO$_4$/(H$_2$)$_{\delta} \Rightarrow$ (Mg,Fe)$_{2-\delta}$SiO$_4$ + $\delta$H$_2$

where the slash on the left hand side indicates that the H$_2$ molecules at first still reside in the olivine structure. After they diffused out the remaining, peroxy-bearing olivine would have the non-stoichiometry shown on the right hand side. If $\delta<1$, this simplifies to (Mg,Fe)$_2$SiO$_4$+$\delta$ representing an olivine with excess oxygen contained in the structure in form of peroxy entities and balanced by a certain concentration of cation vacancies, primarily among Mg$^{2+}$ and Fe$^{2+}$.

It should be noted that the actual oxygen content of natural minerals is never determined analytically. The reason is that oxygen is always assumed to be in the $-2$ oxidation state. Therefore, the contention that mantle-derived olivine may be slightly oxygen–rich, due to the sequence of redox and diffusional steps described by eqs. [2] and [3], cannot be supported by quoting chemical analyses from the literature. If $\delta<1$, it is questionable whether the precision of current analytical techniques suffices to ascertain a small deviation from (Mg,Fe)$_2$SiO$_4$ towards (Mg,Fe)$_2$SiO$_4$+$\delta$. The task is made even more difficult by the fact that any small increase of the Fe$^{3+}$ concentration would also be reflected in a deviation of the bulk stoichiometry towards (Mg,Fe$_{2+}$,Fe$^{3+}$)$_2$SiO$_4$+$\delta$. In order to chemically analyze for the alleged peroxy content one would have to determine both the oxygen to cation ratio and the Fe$^{2+}$/Fe$^{3+}$ ratio. The latter may be possible by means of Mössbauer spectroscopy. It has been noted that the Fe$^{2+}$/Fe$^{3+}$ ratios as determined from Mössbauer spectra are often inconsistent with the Fe$^{2+}$/Fe$^{3+}$ ratios as determined by conventional wet-chemical techniques or as calculated from microprobe data [Dyar et al., 1989]. During wet-chemical analysis any peroxy entities which exist in a mineral will be acid-hydrolyzed and used up to oxidize Fe$^{2+}$, thus resulting in a high apparent Fe$^{3+}$ content. Mössbauer spectroscopy, by contrast, would measure the Fe$^{2+}$/Fe$^{3+}$ ratio irrespective of whether or not peroxy coexist in the mineral matrix. A possible way to assess the peroxy content would be to heat a given olivine crystal in a small fused silica tube, sealed under vacuum, up to a temperature at which all peroxy would be consumed by the oxidation of Fe$^{2+}$ to Fe$^{3+}$, followed by an accurate Mössbauer determination of the Fe$^{3+}$ content retained in the olivine structure or used to form other phases:

$$Fe_{Mg}^x + O_O^x \Rightarrow Fe_{Mg}^x + O_O^x$$

A major uncertainty of such a procedure rests with the assumption that the olivine crystals do not contain any other oxidizable components such as traces of amorphous carbon lining the walls of fluid inclusion cavities and/or dislocations [Green and Guegen, 1983] which would also potentially react with O$^-$ to give CO and CO$_2$.

While MgO is highly ionic, the bonding within the SiO$_4$ tetrahedra in olivine is strongly covalent and every oxygen belongs to an SiO$_4$ tetrahedron. If there is formation of peroxy entities in olivine, they will probably be quite different from the peroxy anion, $O_2^2$, believed to exist in the MgO structure. The exact structure of the peroxy entities in olivine is unknown, but by analogy to other cases general features may be outlined. For the substitutional mode described by eq. [2] the
two most likely OH−-compensated cation vacancy defects contain one or two OH−. Since all O
are bound to Si, they are of the form O₃Si−OH V" Mg and O₃Si−OH V" Mg HO−SiO₃, where V" Mg
stands for a divalent cation vacancy*. A likely peroxy configuration is one that can be described
as peroxy link:

\[
\text{O}_3\text{Si−OH HO−SiO}_3 \Leftrightarrow \text{O}_3\text{Si}^{\text{OO}^-}\text{SiO}_3 + \text{H}_2
\]  

Peroxy links between SiO₄ tetrahedra, O₃Si/OO⁻SiO₃, have been discussed as a possible defect in
vitreous SiO₂, beside the alternative configuration O₃Si/O/SiO₃. They are believed to be the
source defects for the peroxy radicals, O₃Si/OO⁻, which are produced by low temperature (77 K)
γ- or x-irradiation [Friebele et al., 1979]. Shelby [1980] reported on a large increase of Si−OH
when molecular H₂ was allowed to diffuse into initially OH−-poor fused silica and ascribed it to a
reaction similar to eq. [5] reversing from right to left.

Miller et al. [Miller et al., 1987] who sampled olivine crystals from a worldwide distribution of
localities failed to observe the very weak IR band of H₂ molecules. On the basis of this negative
evidence they dismissed the possibility that OH− pairs could undergo a redox conversion into H₂
plus peroxy entities. Unfortunately, the absence of H₂ molecules is not a strong argument against
eq. [3] because H₂ molecules are not only intrinsically IR-inactive but can also diffuse out of the
olivine crystals under study, removing the evidence of former OH− and leaving peroxy entities as
the only memory. We recall that the olivine crystals studied here show no IR band attributable to
OH− impurities. Hence, according to conventional wisdom, these olivine crystals would have to
be classified as OH−-free, suggesting that they come from an anhydrous portion of the upper
mantle. If we admit on the basis of the evidence presented above that the same crystals contain
peroxy entities and that these elusive defects may have formed according to eq. [5], our outlook
changes.

As in the case of O₂⁻ in MgO, peroxy defects in olivine become detectable only after they
dissociate into O−. Peroxy links are expected to dissociate as follows:

\[
\text{O}_3\text{Si}^{\text{OO}^-}\text{SiO}_3 \Leftrightarrow \text{O}_3\text{Si}^{\text{OO}^-}\text{SiO}_3 + \text{H}_2
\]  

The two radicals on the right hand side may also be written as [SiO₄]⁺ or [SiO₄]²⁻ with the hole
delocalizing over the [SiO₄]⁴⁻ tetrahedra [Marfunin, 1979] and possibly beyond. Propagation of
the positive hole can then occur in a similar manner as for O− states in MgO, e.g. by delocalizing
on energy levels near the O 2p-dominated valence band [Freund et al., 1989]. In the case of
MgO, because of its intrinsic diamagnetism, we could follow the spin decoupling which precedes

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* Footnote: Here we do not use the strict Kröger defect nomenclature because the result would be rather
unyielding, given the complexity of the structural unit considered.
the dissociation $\text{O}_2^2^- = O^- + O^-$ by magnetic measurements [Batllo et al., 1990] and compare them to the CDA data. In the case of the olivine crystals the high concentration of Fe$^{2+}$ and other transition metal cations prevents magnetic measurements from being a useful diagnostic tool and we have to rely on CDA for obtaining the necessary information.

The fact that highly mobile charge carriers appear in olivine upon heating to 450°C, generate a positive surface charge, reach saturation between 550–600°C, and disappear upon cooling is strikingly similar to MgO. Since we can be sure that the charge carriers in MgO are positive holes, e.g. O$^-$ states propagating through the O$^{2-}$ sublattice via some sort of O$^{2-}$/O$^{-}$ valency fluctuations on energy levels near the O 2p-dominated valence band, we suggest that the positive charge carriers in olivine are of the same nature. We conclude that the mantle–derived olivine crystals under study contain the equivalent of the peroxy defects in MgO, probably peroxy links as described by eqs. [5/6] or some other similar configuration of self–trapped O$^-$ dimers. These peroxy entities probably derive from solute H$_2$O (and possibly other fluid phase components [Freund, 1987]) that became incorporated into the olivine structure during crystallization in the high pressure–high temperature environment of the upper mantle.

There is a distinct difference to MgO, however, with respect to $F_\Delta$ and its temperature dependence. In the case of MgO we noted (in Part I) that $F_\Delta$ reached a maximum around 750°C and then decreased upon further heating. In Part II we presented evidence that the $F_\Delta$ decrease correlates with the emission of atomic O from the MgO surface. If O is emitted from the top surface layer by O$^-$ disproportionating according to reaction $O^- + O^- \rightarrow O^{2-} + O_2$, two positive charges which had been transported by the O$^-$ from the bulk to the surface become permanently fixed in the form of an anion vacancy, $V_{O^\text{a}}^-$. In order to balance this fixed charge and reduce $F_\Delta$ an aliquot number of Mg$^{2+}$ vacancies, $V_{Mg}^{\text{a}}$, have to diffuse outwards from the bulk to the surface. In the case of MgO it is known that cations become diffusively mobile around 750–800°C by a vacancy hopping mechanism [Sempolinski and Kingery, 1980]. Consequently, annealing of $V_{Mg}^{\text{a}}$ is the most likely process that accounts for the reduction of $F_\Delta$ above 750°C in MgO. In the case of olivine, as shown in Fig. 4d, we do not observe a clear maximum in the $F_\Delta(T)$ curve but only a transient feature followed by a further rise beyond 750°C. Concurrently the $F_\Delta(T)$ reversibility is lost. In some experiments we noted a brownish hue on the olivine surface, indicating the onset of oxidation. Whether or not this oxidation ≥750°C is caused by Fe$^{2+}$ reacting with extraneous gas phase O$_2$ in the unbuffered N$_2$ atmosphere or by Fe$^{2+}$ reacting with indigenous O$^-$ is yet unclear. In both cases $F_\Delta$ is expected to become even more positive through the formation of Fe$^{3+}$ on (Mg,Fe$^{2+}$) lattice sites.

At the beginning, in Part I, we pointed out that the electrical conductivity of olivine widely believed to be relevant to the upper mantle is the conductivity experimentally determined after the anomaly ≤700°C has been removed. The removal is achieved by heating the mantle–derived olivine crystals in CO/CO$_2$ or H$_2$/CO$_2$ gas mixtures until their electrical conductivity values come down to the low level characteristic for 0.1 MPa conditions. This procedure is justified by the conjecture that the conductivity anomaly is nothing but a “dirt effect”, caused by extraneous
contamination. Now we know that the anomaly is due to indigenous O⁻ charge carriers. To the best of our current understanding these O⁻ charge carriers and their parent defects, the peroxy entities, represent a “memory” of a solute H₂O content, e.g. of an earlier OH⁻ content. IR spectroscopy in inapplicable because H₂ molecules and peroxy are intrinsically IR-inactive. Any attempt to use Raman spectroscopy would probably fail because of the relatively low H₂ and peroxy concentration. Electron paramagnetic resonance spectroscopy cannot used either because mantle-derived olivine crystals typically contain a much too high Fe²⁺ content.

By heating mantle-derived olivine to 700°C in CO/CO₂ or H₂/CO₂ at 0.1 MPa and thus removing their conductivity anomaly, the crystals loose their indigenous O⁻ type charge carriers. Thus, their natural state is altered. The experimentally measured conductivity may no longer reflect the “true” conductivity which the same crystals would display, if they still had their full complement of O⁻. One may argue that 700°C is too low a temperature to be of significance because, in the upper mantle, temperatures are likely to exceed 1000°C. However, this argument is fallacious because it is based on the assumption that the O⁻ charge carriers must disappear around 700°C. In reality, the annihilation of O⁻ around 700°C is probably an artefact of the experimental set-up used in any conductivity measurement: the O⁻ charge carriers are only consumed because the samples possess a free surface across which the O⁻ react with the CO/CO₂ or H₂/CO₂ gases. In the laboratory set-up the system is not closed. In the upper mantle, however, the olivine crystals will not possess a free surface (unless CO/CO₂/H₂O fluids freely percolate around each grain). Therefore, to the extent that the upper mantle can be considered a closed system, the possibility has to be taken into consideration that O⁻ charge carriers persist beyond the 700°C threshold annihilation temperature.

Very little is known about the effect of pressure on the stability of O⁻ and on the stability of their parents defects, the peroxy entities. Given the fact that the O⁻—O⁻ peroxy bond is short, typically ≈1.5 Å [Cremer, 1983] compared to 2.8–3.0 Å for most O²⁻—O²⁻ distances in silicates, pressure is expected to delay peroxy dissociation and, hence, the generation of O⁻ charge carriers to higher temperatures [Freund, 1987]. Unless there are competitive mechanisms by which O⁻ can be consumed, for instance by oxidizing Fe²⁺ to Fe³⁺ and incorporating these Fe³⁺ into phases such as spinel or orthopyroxenes which accommodate or require Fe³⁺ in their respective crystal structures, the range over which O⁻ charge carriers might exist under upper mantle pressure conditions may extend into the 1000–1200°C region.

So far we only studied the O⁻ charge carrier generation at 0.1 MPa pressure during heating. From the available evidence we conclude that the O⁻ derive from peroxy parents. We recognize that the olivine crystals under study have a complex history in the pressure–temperature–time (P–T–t) space. At which point in the P–T–t space the peroxy parents have been introduced is unknown. They may have formed in the mantle, e.g. before the olivine crystals were brought up to the surface of the earth by volcanic activity. They may have formed during ascent while the olivine crystals underwent decompression and cooling. They may have formed after cooling. Each scenario would lead to different consequences with respect to the conductivity behavior of olivine
in the upper mantle. As long as the P-T-t pathway of the peroxy formation reaction is unknown, the field remains open to speculation.

One question, however, can be answered with some degree of confidence: The long-standing controversy concerning the sensitivity of olivine to oxidation in unbuffered environments. Many researchers have observed that olivine crystals readily show signs of oxidation when heated in any gas environment where the oxygen fugacity lies outside the thermodynamic stability range of the olivine phase [Nitsan, 1974]. As tell-tale signs for beginning oxidation color changes are reported and the precipitation of minute crystals of iron oxides at the surface and along dislocation lines [Duba and Nicholls, 1973, Durham and Goetze, 1977]. If one is unaware of the possible presence of peroxy entities in olivine, the conclusion appears unavoidable that traces of O$_2$ in the gas or traces of H$_2$O in microcracks are the oxidants and that only a prolonged exposure to reducing CO/CO$_2$ or H$_2$/CO$_2$ gases will correct the situation. However, if mantle-derived olivine crystals contain peroxy entities in their structure, due to solute H$_2$O component, followed by the redox reaction and loss of H$_2$ by diffusion as discussed above, the outlook changes: Crystals containing peroxy defects are susceptible to an internal oxidation according to eq. [4]. Such an internal oxidation occurs in the absence of any trace of O$_2$ in the gas phase. Even if one were able to produce an Ar or N$_2$ atmosphere with only a few O$_2$ molecules per liter, equivalent to P$_{O_2}$<10$^{-20}$ as required by the linear extrapolation of the thermodynamic data pertinent to the high temperature stability of olivine, the olivine crystals under study would still oxidize from within.

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References


VI List of Publications Describing Further Work Performed


