PULMONARY TOXICITY OF SIMULATED LUNAR AND MARTIAN DUSTS IN MICE: I. HISTOPATHOLOGY 7 AND 90 DAYS AFTER INTRATRACHEAL INSTILLATION

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NASA is contemplating sending humans to Mars and to the moon for further exploration. Volcanic ashes from Arizona and Hawaii with mineral properties similar to those of lunar and Martian soils, respectively, are used to simulate lunar and Martian environments for instrument testing. Martian soil is highly oxidative; this property is not found in Earth’s volcanic ashes. NASA is concerned about the health risk from potential exposure of workers in the test facilities. Fine lunar soil simulant (LSS), Martian soil simulant (MSS), titanium dioxide, or quartz in saline was intratracheally instilled into groups of 4 mice (C57BL/6) at 0.1 mg/mouse (low dose, LD) or 1 mg/mouse (high dose, HD). Separate groups of mice were exposed to ozone (0.5 ppm for 3 h) prior to MSS instillation. Lungs were harvested for histopathological examination 7 or 90 days after the single dust treatment. The lungs of the LSS-LD groups showed no evidence of inflammation, edema, or fibrosis; clumps of particles and an increased number of macrophages were visible after 7 days but not 90 days. In the LSS-HD-7d group, the lungs showed mild to moderate alveolitis, and perivascular and peribronchiolar inflammation. The LSS-HD-90d group showed signs of mild chronic pulmonary inflammation, septal thickening, and some fibrosis. Foci of particle-laden macrophages (PLMs) were still visible. Lung lesions in the MSS-LD-7d group were similar to those observed in the LSS-HD-7d group. The MSS-HD-90d group had PLMs and scattered foci of mild fibrosis in the lungs. The MSS-HD-7d group showed large foci

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of PLMs, intra-alveolar debris, mild-to-moderate focal alveolitis, and perivascular and peribronchiolar inflammation. The MSS-HD-90d group showed focal chronic mild-to-moderate alveolitis and fibrosis. The findings in the O₃-MSS-HD-90d group included widespread intra-alveolar debris, focal moderate alveolitis, and fibrosis. Lung lesions in the MSS groups were more severe with the ozone pretreatment. The effects of O₃ and MSS coexposure appeared to be more than additive. Results for the TiO₂ and quartz controls were consistent with the known pulmonary toxicity of these compounds. The overall severity of lung injury was TiO₂ < LSS < MSS < O₃ + MSS < quartz. Except for TiO₂, the increased duration of dust presence in the lung from 7 to 90 days transformed the acute inflammatory response to a chronic inflammatory lesion. This study showed that LSS and MSS are more hazardous in the lungs than nuisance dusts.

NASA is considering returning to the Moon to establish a lunar outpost and sending humans to explore Mars (Hoffman & Kaplan, 1997). Instruments and hardware destined for these extraterrestrial bases must be tested in simulated Martian and lunar environments in NASA laboratories. NASA has used ashes obtained from the San Francisco volcano field in Arizona and from a Hawaiian volcano for environmental simulation (Glaser, 1992). These volcanic ashes have mineral properties resembling those of lunar and Martian soils, respectively (McKay et al., 1994; Allen et al., 1998). Since the inhalation toxicity of these soil simulants has not been evaluated, NASA is concerned about the health risk to workers from potential dust exposures in the Earth-based test facilities. Toxicity testing with simulants can also provide guidance for designing experiments with actual lunar dust and eventually Martian dust.

During the first Apollo lunar landing, the moon dust particles encountered were described by the crew as extremely fine (NASA, 1969). The dust stuck to their spacesuits despite attempts to brush it off and was brought into the lunar landing vehicles. After the spacecraft lifted off from the lunar surface and microgravity was reestablished, a large quantity of floating dust made breathing difficult and impaired the visual acuity of the crew (NASA, 1970). NASA conducted a few studies in the 1970s to assess the risk of exposure, but the results failed to reveal whether the dust was an inhalation hazard (Holland & Simmonds, 1972) (see Discussion for details). Russian investigators also conducted a few studies on lunar dust. They concluded that lunar soil caused fibrosis and other signs of pneumoconiosis in animals (Kustov et al., 1974, 1989). However, the studies contained considerable experimental deficiencies (see Discussion for details). The toxicity of lunar dust must be further determined before humans are sent to the Moon for long periods of habitation. In general, NASA will not release a lunar sample for study until a successful study on the simulant (JSC-1) has been conducted. Addressing this requirement was one of the purposes of the present study.

The composition of the lunar simulant is a good match for its extraterrestrial counterpart; a detailed comparison has been published by McKay et al. (1994). In brief, both the lunar soil and its simulant contain about 50% SiO₂; other common soil oxides (Al₂O₃, FeO, MgO, and CaO) account for
another 42% to 45%. No trace or heavy metals were found. The particle-size distribution (average about 100 µm) of the simulant fits within the range of lunar samples returned by Apollo missions. The simulant contains about 1% respirable dust (see Methods).

Martian surface soil is similar to Hawaiian volcanic ash in mineral content (Allen et al., 1998). Martian soil and the simulant (after removal of water and other volatiles) both contain about 45% SiO₂ and 15% Fe₂O₃, which imparts a brown color to both the regolith and its simulant. The other components that account for the majority of the remaining bulk are Al₂O₃, CaO, TiO₂, MgO, Na₂O, and K₂O. No trace or heavy metals have been found. Only 1% of the simulant is smaller than 5 µm. Detailed comparison of the Martian soil with its simulant can be found in a report published by Allen et al. (1998).

Martian soil is reactive; the evidence comes from experiment results obtained by the Viking Martian lander. When surface soil samples on Mars were humidified, evolution of oxygen was observed, suggesting the presence of superoxides or peroxides on the soil (Oyama & Berdahl, 1977). Reactive and oxidative species in Martian analog soil were recently produced under simulated Martian surface conditions (Yen et al., 2000; Tsapin et al., 2000). Inhalation exposures to Martian regolith would be expected to produce lesions in the lung partially resembling those caused by powerful oxidants. Since oxidatively activated Martian analog soil was not available at the time of this study, an ozone exposure of the mice followed by an intratracheal instillation with the unactivated ash was included to simulate the effects of the oxidative properties of Martian soil in the lung. Ozone was chosen because its effects in the lung have been well characterized, and it was feasible to select a dose that would produce only minimal tissue injury so the combined effects could be discerned.

**MATERIALS AND METHODS**

**Simulated Martian and Lunar Dusts**

Raw Arizona and Hawaii volcanic ashes [designated as JSC-1 (McKay et al., 1994) and JSC-Mars-1 (Allen et al., 1998)] were provided to Lovelace Respiratory Research Institute for size fractionation. According to the procedure used at Lovelace (Cheng, personal communication, 1998), the raw material was placed in a DeVilbiss dry powder generator operated with compressed air at 20 psig; the output was then delivered to a 3-stage cyclone operated at 31 L/min (Smith et al., 1979). The cutoff aerodynamic diameters were 5, 1.95, and 0.28 µm for the 3 stages. The fine particles from the second and third stages and the backup filter were collected and pooled for the present study. Thus, the mass median aerodynamic diameter (MMAD) of both soils was expected to be less than 5 µm.

Because the present study was not conducted by the inhalation route (for which particle sizes are important in determining the fractional deposi-
tion of inhaled particles into the pulmonary region), the simulants were used without further characterization of particle-size distribution before the intratracheal instillation study. Some time after the intratracheal instillation study was completed, Microtrac, Inc. (Montgomeryville, PA), offered to determine particle sizes of the simulated dusts using their newly marketed instrument. In Microtrac’s assay, approximately 50 mg of the LSS or MSS was suspended in 200 ml of distilled water and placed in the reservoir chamber of a Microtrac X-100 particle-size analyzer. This trilaser analytical unit was connected to a computer unit for data acquisition and analysis. The instrument has a particle size analytical range of 0.04 to 700 µm separated into 120 channels. Timed ultrasonication, which is controlled by the computer, can be generated in the reservoir chamber to ensure that the particles remain homogeneously suspended. The distribution profiles showed that the respirable fraction of the LSS had mass median diameters (MMDs) of 3 µm or less, and contained 5% large particles with MMD of 81 µm; upon ultrasonication, all of the large particles disappeared and the resulting suspension showed a bimodal distribution, with MMDs of 1.05 µm (44%) and 2.99 µm (56%). The MSS isolated similarly had a bimodal distribution that included a mass fraction with 70% fine dust with a mass median diameter (MMD) of 3 µm and 30% particles with MMD of 108 µm (diameters ranging from 30 to 400 µm). To our surprise, repeated ultrasonications still could not significantly change this particle-distribution profile.

Titanium Dioxide and Crystalline Silica

The titanium dioxide sample, a product of Particle Information Services (Kingston, MA), had an average particle diameter of 0.45 µm. Crystalline silica (quartz) (acid-washed Min-U-Sil-5) with an average particle size of 5 µm was obtained from Pennsylvania Glass and Sand (Pittsburgh, PA). These particle-size specifications were from the respective vendors.

Animals and Animal Husbandry

Male mice (C57BL/6J, 2 mo old), free of known rodent pathogens, were obtained from Jackson Laboratory (Bar Harbor, ME). The animals were housed in groups of four in polycarbonate cages (with HEPA air filters) in the AAALAC-accredited vivarium at the Johnson Space Center (JSC). Animals were allowed to acclimate at this facility (with a 12-h light–dark cycle) for at least 1 wk before being used in the study. The mice had free access to tap water and Purina Formulab Chow number 50008 (Ralston Purina Co., St. Louis, MO). They were cared for and used humanely according to NASA Animal Care and Use Program guidelines.

Inhalation Exposure to Ozone

The animals in the MSS study that required a preexposure to ozone were placed in a 0.75-m³ Laskin-type stainless steel exposure chamber (Wahman, MD) (Kirichenko et al., 1996). Ozone was generated by passing house air through an electrical ozone generator (model V5AR-237 ozona-
tor, Ozone Research Equipment Corp, Phoenix, AZ); it was then metered and mixed with house air before entering the chamber. A small air stream was pulled (2 L/min) from the chamber through Teflon tubing into a calibrated ozone monitor (model 1003AH, Dasibi Environmental Corporation, Glendale, CA). The monitor was calibrated shortly before this study by a laboratory at the U.S. Environmental Protection Agency (Houston Regional Office). After the chamber ozone concentration had been stabilized at 0.5 ppm, 6 groups of mice (4 mice/group) were placed in the chamber for 3 h and the ozone concentration was recorded every 10 min. The exposure level was selected to produce no more than minimal tissue injury.

**Intratracheal Instillation of Dust Samples**

Each mouse was anesthetized intraperitoneally with a cocktail of 150 ml of ketamine and xylazine at a dose of about 80 and 16 mg/kg, respectively. After the animal was labeled with an ear tag and body weight was determined, it was secured on an inclined plastic platform (about 60°). The trachea was exposed by a 1-cm incision on the ventral neck skin (Leigh et al., 2000). An aliquot of 50 µl of freshly suspended and ultrasonicated dust in normal saline, containing 0.1 mg (LD) or 1 mg (HD) dust, was intratracheally instilled (using a 26-g needle) into the lung. For the saline-only and ozone-only control animals, only saline was instilled. The incision was then sutured and swabbed with Povidone iodine. The mice of each treatment group (4/group) were caged together in the vivarium and observed daily until their scheduled termination.

**Collection of Lungs from Animals for Histopathological Study**

Seven or 90 days after the dust treatment, each mouse was injected intraperitoneally with a lethal dose of pentobarbital sodium solution (Nembutal, Abbott, North Chicago, IL). Body weight was determined. An incision was made to expose the trachea for inserting a catheter; formalin (10% in a neutral phosphate buffer) was allowed to drip by gravity (from a 25-cc syringe barrel hanging 1.5 ft above the neck) through the catheter into the lung for about 10 min. The trachea was then tied and the isolated lung was placed in a glass vial containing about 10 ml of the same fixative. Each glass vial was assigned a number unknown (with respect to treatment) to the pathologists. The lungs were fixed for at least 7 days before further processing.

**Preparation of Lung Tissue Slides for Histopathological Examination**

The formalin-fixed mouse lungs were embedded in paraffin, thin-sectioned coronally, and mounted on glass microscope slides using standard histopathological techniques. Sections were stained with hematoxylin–eosin. For each mouse, one slide containing one lung section was examined independently by three pathologists. Van Gieson and trichrome stains were also used to stain elastin and connective tissue to provide supplementary information to the pathologists. A formatted score sheet was used for grading intraalveolar, interstitial, vascular, and bronchiolar/bronchial areas for
the presence of particles, macrophages, inflammation cells, edema, congestion, hyperplasia, fibrosis, and other parameters (when applicable). The scores from the three pathologists were averaged.

RESULTS

Effects of Instilled Titanium Dioxide

After receiving a single intratracheal instillation of TiO$_2$ (0.1 mg/mouse, low dose, LD), the lungs of the mice killed 7 days after the treatment showed focal clumps of particles; some of the particles were inside macrophages. There was no evidence of inflammation, edema, or fibrosis (Table 1). The lungs of the mice that were killed 90 days after dust treatment had no identifiable particles and were completely normal (Figure 1A). This slide, which is indistinguishable from those of the saline-treated controls, is used in this manuscript for illustrating the appearance of normal lung tissue and treatment with a negative-control dust.

### TABLE 1. Summary of lung histopathology results from intratracheally instilled mice

| Treatment          | Dust dose (mg/mouse) | Day | ↑ Macrophages | Inflammation | Edema | Fibrosis |
|--------------------|----------------------|-----|---------------|--------------|-------|----------|
| Saline             | 0                    | 7   | 0             | 0            | 0     | 0        |
| O$_2$ (0.5 ppm)    | 0                    | 7   | 0             | 0            | 0     | 0        |
| TiO$_2$            | 0.1                  | 7   | +             | 0            | 0     | 0        |
| LSS                | 0.1                  | 7   | +             | 0            | 0     | 0        |
| MSS                | 0.1                  | 7   | +             | 0            | 0     | 0        |
| MSS + O$_3$ (0.5 ppm) | 0.1               | 7   | +             | 0            | 0     | 0        |
| SiO$_2$            | 0.1                  | 7   | +             | +            | +     | 0        |
| Saline             | 0                    | 90  | 0             | 0            | 0     | 0        |
| O$_2$ (0.5 ppm)    | 0                    | 90  | 0             | +            | 0     | 0        |
| TiO$_2$            | 0.1                  | 90  | 0             | 0            | 0     | 0        |
| LSS                | 0.1                  | 90  | 0             | 0            | 0     | 0        |
| MSS                | 0.1                  | 90  | +             | 0            | 0     | +        |
| MSS + O$_3$ (0.5 ppm) | 0.1               | 90  | +             | 0            | 0     | +        |
| SiO$_2$            | 0.1                  | 90  | +             | +            | 0     | +        |
| TiO$_2$            | 1.0                  | 7   | +             | 0            | 0     | 0        |
| LSS                | 1.0                  | 7   | +             | +            | 0     | 0        |
| MSS                | 1.0                  | 7   | ++            | +            | 0     | 0        |
| MSS + O$_3$ (0.5 ppm) | 1.0               | 7   | ++            | ++           | 0     | +        |
| SiO$_2$            | 1.0                  | 7   | ++            | +++          | 0     | +        |
| TiO$_2$            | 1.0                  | 90  | ++            | 0            | 0     | 0        |
| LSS                | 1.0                  | 90  | +             | +            | 0     | +        |
| MSS                | 1.0                  | 90  | +             | +            | 0     | +        |
| MSS + O$_3$ (0.5 ppm) | 1.0               | 90  | ++            | ++           | 0     | +        |
| SiO$_2$            | 1.0                  | 90  | ++            | +++          | 0     | +        |

Note. Groups of 4 mice were intratracheally instilled with 0, 0.1, or 1 mg of a dust per mouse and killed 7 or 90 days after the single treatment. The results in the table were condensed from formatted score sheets, which recorded intraalveolar, interstitial, vascular, and bronchiolar/bronchial areas for relevant histopathological parameters.
When compared to the lungs of low-dose animals of the same treatment duration, lungs of mice in the high-dose group (1 mg/mouse) that were killed 7 days after the dust treatment had larger focal clumps of particles; the HD-90d group had increased numbers of alveolar macrophages and mild focal perivascular inflammation (Figure 1B).

**Effects of Instilled Quartz**

The lungs of the SiO$_2$-LD-7d group showed focal clumps of particles and an increased number of alveolar macrophages. Mild to moderate focal alveolar inflammation, mild focal perivascular and peribronchiolar inflammation, neutrophilic and lymphocytic infiltration, and edema also were observed. The lungs of mice in the SiO$_2$-LD group killed 90 days after dust treatment still had clumps of particles and particle-laden macrophages (PLMs) in addition to mild inflammation and mild focal fibrosis (Figure 1C).
The lungs of the HD-7d group contained larger focal clumps of particles and more macrophages compared with the lungs of the LD-7d group. In addition, the lung lesions were more severe, showing moderate to severe alveolitis, moderate focal perivascular and peribronchiolar inflammation, and edema. For the high-dose group in the 90-day study, large clumps of PLMs were still abundant in the lungs. The lungs also showed moderate alveolitis; moderate to severe focal bronchiolar, perivascular, and peribronchiolar inflammation; necrosis; edema; and focally severe fibrosis, strongly resembling bronchiolitis obliterans organizing pneumonia (Figure 1D).

**Effects of Instilled Lunar Soil Simulant (LSS)**

The lungs of mice in the LSS-LD-7d group showed the presence of focal clumps of particles and, occasionally, macrophages (Figure 2A). There was no evidence of inflammation, edema, or fibrosis. At 90 days, particles were not observed and the tissue was normal (Figure 2B).

FIGURE 2. Lung histopathology of mice intratracheally instilled with a single dose of 0.1 mg (LD) or 1 mg (HD) of lunar soil simulant and killed 7 or 90 days after the treatment. In the figures, ad = alveolar duct, b = bronchus, rb = respiratory bronchus, v = vessel; in (A), ➞ = particle-laden macrophages; (B) shows normal tissue and no particles; (C) shows mild to moderate alveolitis with lymphocyte and neutrophile infiltration, ➞ = alveolar damage, particle-laden macrophages, and inflammatory cells; (D) shows mild alveolitis, ➞ = septal thickening, and blood vessel and bronchial inflammation. Magnification: ×200.
The lungs of mice in the HD-7d group showed clumps of particles and macrophages, mild to moderate alveolitis with neutrophilic and lymphocytic infiltration, and mild perivascular and peribronchiolar inflammation (Figure 2C). When they were examined 90 days after the dust treatment, foci of PLMs, mild alveolitis, mild focal perivascular and peribronchiolar chronic inflammation, and septal thickening were visible (Figure 2D).

**Effects of Instilled Martian Soil Simulant (MSS)**

The lungs of mice in the MSS-LD-7d group showed particles and macrophages; mild focal intra-alveolar debris, infiltration of neutrophils and lymphocytes, and perivascular and peribronchiolar inflammation were evident (Figure 3A). At 90 days, PLMs and scattered foci of mild fibrosis were detected (Figure 3B).

The lungs of the HD-7d group had large foci of PLMs, mild focal alveolitis, intra-alveolar debris, and mild to moderate perivascular and peri-
bronchiolar inflammation (Figure 3C). At 90 days, PLMs, mild alveolitis, focal moderate inflammation, and fibrosis were observed (Figure 3D).

**Effects of Ozone and Instilled MSS**

A single 3-h inhalation exposure to 0.51 ± 0.02 ppm ozone (followed by saline instillation) produced no overt lung injury when lungs were examined 7 days after the exposure; when they were examined 90 days after the exposure, mild focal alveolitis and peribronchiolar inflammation were noted. When an ozone exposure was followed by an intratracheal instillation of MSS (0.1 mg or 1 mg/mouse), the lungs of the LD-7d group showed foci of particles and an increase in macrophages. Focal mild peribronchiolar inflammation and alveolitis also were visible (Figure 4A). At 90 days, the lungs of the O₃ + MSS-LD mice revealed focal mild peribronchiolar inflammation, alveolitis, and fibrosis (Figure 4B).

Lungs of O₃ + MSS-HD-7d group showed large foci of PLMs, focal moderate peribronchiolar inflammation, alveolitis, intraalveolar debris, and early focal moderate to severe fibrosis (Figure 4C). The HD-90d group had widespread intraalveolar debris, alveolitis, and focal moderate inflammation and fibrosis (Figure 4D).

**Body Weight Changes after Instillation of the Dusts**

There were no statistically significant differences in body weight gains between the exposed groups and control groups killed either on 7 or 90 days after instillation of the dusts. After 7 days, the group average body weight gains ranged from a low of −0.3 g (0.1 mg MSS) to +2.2 g (ozone control group). After 90 days, the weight gains ranged from 4.0 g to 6.4 g. The groups with the lowest average body weight gains were the O₃ + MSS-HD (4.0 ± 1.6 g), SiO₂-LD (4.6 ± 1.3 g), and SiO₂-HD (4.9 ± 0.7 g) groups. These were among the groups that showed more severe histopathological effects in the lungs.

**DISCUSSION**

The present histopathology study and a lavage study (see companion paper, Lam et al., 2002) were conducted to assess the pulmonary toxicity of simulated Martian and lunar dusts. Another goal was to establish experiment protocols that could be applied to the study of lunar soil and eventually Martian soil. We anticipate that the quantity of extraterrestrial dusts in the respirable range that NASA could make available for toxicology studies will be very small. Inevitably, mice would be the animal species of choice and intratracheal instillation would be the route of administration to conserve the scarce dusts. Thus, this animal model and route of exposure were selected for the present study, with TiO₂ and crystalline silica serving as negative and positive controls.

The doses for intratracheal instillation in this study were chosen on the basis of information from Henderson et al. (1995). Henderson reported that
in rats exposed by inhalation to 10 mg/m$^3$ of quartz or TiO$_2$ for 1 wk, the lung contained 0.76 mg quartz or 0.44 mg TiO$_2$. Information compiled by Boggs (1992) and Lai (1992) shows that the minute respiratory volume (MRV) of a 300-g rat, a 35-g mouse, and a 20-g mouse was 0.210, 0.040, and 0.025 L/min, respectively. The MRV of the mice (average 25 g) in our study would be about 0.030 L/min. If it is assumed that dust deposition in the lung is roughly proportional to the MRV of the exposed animals, then if Henderson et al. (1995) had included mice (~25 g) in their study, the dust burden of the mouse lung would have been about 0.11 mg quartz or 0.06 mg TiO$_2$. In the present study, a single dose of 0.1 mg per mouse was chosen as a low dose and 1 mg as a high dose.

Bermudez et al. (2000) reported that the lung burdens (determined at the end of the last exposure) of mice exposed to TiO$_2$ at 10, 50, or 250 mg/
m³ for 13 wk (6 h/day, 5 days/wk) were 5.2, 53.5, and 170.2 mg/g of dry lung tissue respectively (Bermudez et al., 2000). The corresponding values for lung burden of TiO₂ per mouse calculated by these authors were approximately 0.2, 2, and 13 mg (Bermudez, 2000, personal communication). It is noteworthy that the low dose (0.1 mg/mouse) used in our study fell between the values for lung TiO₂ burden of mice exposed to 10 mg/m³ for 1 wk (0.06 mg/mouse, extrapolated above using Henderson’s data) and those exposed to 10 mg/ m³ for 13 wk (0.2 mg/mouse; Bermudez, 2000, personal communication). The high dose in our study (1 mg/mouse) was half the value of the body burden of mice exposed to 50 mg/m³ for 13 wk in Bermudez’s study. This information is useful for quantitative comparisons of dust exposures by different route of pulmonary administrations, as long as the bolus nature of the intratracheal instillation is recognized (Driscolll et al., 2000). It also should be noted that the residual lung burden determined at the end of the last inhalation exposure is the dynamic result of cumulative intermittent deliveries and continuous elimination.

When the mice were each intratracheally instilled with 0.1 mg of TiO₂ particles, an increased number of macrophages, compared with that in the lung of saline controls, was seen 7 days after treatment. This increase was not visible 90 days after treatment and the lung tissue was apparently normal. When treated with a high TiO₂ dose (1 mg/mouse) and killed 90 days after dust instillation, the mice had particles persisting in their lungs, and numerous macrophages were present. Tissue histopathological manifestations were minimal. These findings were consistent with the observations reported by Bermudez et al. (2000) and colleagues (Everitt et al., 2000). In their studies, the lungs of the mice exposed for 90 days to 50 mg/m³ (TiO₂ lung burden of ~2 mg) had increased numbers of macrophages and neutrophils and “elevated soluble indices of inflammation.” These parameters remained elevated when lungs were examined at 26 wk after the last exposure. The increases were not observed in mice exposed to 10 mg/m³ (lung burden ~0.2 mg).

For those animals instilled with 0.1 mg/mouse and killed 90 days after treatment, MSS and SiO₂ particles, but not LSS and TiO₂, were still visible in the lungs. This observation indicated that TiO₂ and LSS were easier to clear from the lungs than MSS and SiO₂. However, when the animals were given 1 mg/mouse, all 4 dusts were visible in the lung regardless of whether the animals were killed 7 or 90 days after dust treatment. The high doses apparently overwhelmed the pulmonary clearance mechanisms. As mentioned earlier, particle size characterization after the instillation studies showed that the respirable fraction of LSS ultrasonicated in water had a bimodal distribution with mass median diameters (MMDs) of 1 μm (44%) and 3 μm (56%). The MSS prepared similarly had a bimodal distribution that included a mass fraction of 70% fine dust with an MMD of 3 μm and 30% particles with an MMD of 108 μm (diameters ranging from 30 to 400 μm), which is not in the respirable range. Repeated ultrasonications
did not significantly diminish the population of these large particles. Thus, about 30% of the mass of the MSS given to the mice consisted of particles in the nonrespirable range. In general, per unit weight, toxic insults to the lung from particles decrease with an increase in particle size because a fine dust sample contains a much greater number of particles than a coarse sample of equal weight. The presence of large particles in MSS would quantitatively, but not qualitatively, influence the dose response. However, large particles, which are more difficult for macrophages to phagocytize, would be more persistent in the lung, and this fact may partially explain why MSS was more persistent than LSS in the lung.

LSS was more toxicologically reactive than TiO$_2$ in the mice; the difference was more evident at the high dose. Like TiO$_2$, LSS at the low dose produced very little tissue response. At the high dose, acute mild to moderate inflammation of the lung tissue was observed among the LSS-treated mice. With the continuous irritation of the high-dose LSS dust in the lung, acute inflammation observed in the 7-day group was transformed into chronic inflammation, which was observed 90 days after dust treatment. However, the inflammatory manifestations were relatively mild with this dust. MSS elicited a more severe reaction than the LSS (compare Figures 2 and 3). Inflammatory responses, not seen in animals in the LSS-LD groups, were detected in both the MSS-LD-7d and MSS-LD-90d groups. Inflammatory reactions, which were mild in the MSS-LD-7d group, were mild to moderate in the lungs of the MSS-HD-7d group. Fibrosis was observed in the MSS-HD-90d group.

Ozone pretreatment followed by MSS instillation was used to emulate the oxidative and reactive nature of actual Martian soil. As expected, mice receiving such treatment showed a greater pulmonary toxicity response than those receiving either compound alone; the toxic manifestations were especially discernible in ozone-pretreated animals instilled with the high dose of MSS (compare Figures 3 and 4). The O$_3$ + MSS-HD-7d group exhibited moderate pulmonary reactions with early focal moderate to severe fibrosis. The animals in the 90-day study had widespread intraalveolar debris, moderate alveolitis, and fibrosis. The effects of ozone and MSS coexposure appeared to be more than additive. Nevertheless, these effects were less severe than those seen in animals treated with high doses of crystalline silica.

Toxicity of lunar soil samples was investigated in the 1970s by NASA, but the data obtained were of limited usefulness for risk assessment. The studies involved intraperitoneal and subcutaneous injections of the lunar soil into mice and intratracheal instillation of the material into guinea pigs. The results showed acute inflammatory responses throughout the peritoneum or surrounding the subcutaneous injection site; low-grade inflammatory reactions, but no fibroblastic lesions, were detected 20 mo after the injection. In a study consisting of 3 groups of guinea pigs (4/group), 3 animals per group were each given 2 ml saline containing 2% soil; the remaining animal was
given saline containing no soil (Holland & Simmonds, 1972). Two of the
groups were killed either 2 or 4 days after the injection. No information was
given on exactly when the third group, which was held for long-term ob-
servation, was killed. Without specifying the group(s), the authors reported
that diffuse alveolar-cell hypertrophy, septal edema, mononuclear infiltr-
ation, and proliferation of alveolar macrophages were observed. The authors
stated, “The tissue response seemed out of proportion to the amount of par-
ticle material present.” Unable to discern the treatment effects because of a
significant degree of spontaneous pulmonary pathology in both the control
and dust-treated guinea pigs, the authors concluded that “the potentially
harmful or innocuous nature of the dust remains to be investigated.”

Toxicity data collected by Russian investigators are also of limited value.
In one study mice were exposed for 4 days (4 h/day) to air (0.5 L/min) that
was “passed through a layer” consisting of a preweighed sample of lunar
surface material (Kustov et al., 1974); no pathomorphological effects were
detected and no information on weight loss after the air passed through the
test material was reported. Another Russian study (Kustov et al., 1989) was
conducted with Wistar rats, each injected intratracheally with a single dose
of 50 mg (in 1% starch solution) of lunar dust or terrestrial analog (a basalt
from Kamchatka, Russia). The compositions of the Russian lunar soil and ter-
restrial analog were comparable to U.S. counterparts. Six months after the
dust treatment, the lung tissue revealed many fibrous cellular nodes that
contained dust particles, macrophages, lymphocytes, plasmocytes, fibro-
blasts, and giant multinuclear cells. The terrestrial analog, like the extrater-
restrial counterpart, also produced pneumoconiosis characterized by the
development of fibrous cellular nodes, pronounced interstitial fibrosis, and
interlobular septa.

These results are to be expected because of the massive dose of dust
used by the Russian investigators. For example, Bermudez et al. (2000)
reported that 13-wk exposures (6 h/day, 5 days/wk) of rats to 250 mg/m³
TiO₂ (a dust that at low concentrations produces very little effect in rats)
resulted in hypertrophy and hyperplasia of alveolar epithelium, and fibrotic
changes; these effects were not seen in rats exposed to 50 mg/m³ TiO₂. The
lung burden of dust in rats exposed to 250 or 50 mg/m³ TiO₂ determined at
the end of the exposures was 40 or 8 mg/rat, respectively. The bolus dose of
50 mg/rat in the Russian study apparently seriously overloaded the rats’
lungs. It is well known that greatly overloading the lungs of rats with dust
impairs cellular responses that clear and sequester particles, leading to a
spectrum of histopathology that could include cancer (Trochimowicz, 1988;
Muhle et al., 1990; Morrow, 1992; Hext, 1994). Thus, a conclusion could
not be drawn from the results of Kustov et al. (1989) on whether lunar soil
is fibrogenic at reasonable human exposure levels.

The overall results of the present study showed that LSS and MSS are
not simply nuisance dusts. It is reasonable to predict that human inhala-
tion exposures to LSS or MSS will produce more pulmonary responses than exposures to TiO$_2$, but less severe responses than crystalline SiO$_2$. Thus, workers using these dusts in test environments must use respiratory protection. The toxicity information and safety recommendation have been provided to the NASA Astromaterials Curator to generate material safety data sheets on the simulants. Even though the chemical compositions of these dust simulants are similar to lunar or Martian soil, the results of this study can give no more than a qualitative indication of the hazards of the actual extraterrestrial soils. The results of the coexposures to ozone and MSS suggest that Martian soil, a dust with a reactive surface and probably persistent in the lung like MSS, could produce more severe injury than that caused by the simulant alone. These findings support NASA’s intention to evaluate the oxidative and reactive Martian surface soil before sending humans to the hostile Martian environment (Sullivan, 2000, personal communication). In the present study, the choice of doses, mice as the animal model, positive and negative controls, and intratracheal instillation as the route of exposure have enabled us to toxicologically evaluate these simulated extraterrestrial soils using very small amounts of materials. Thus, the design and methodology of the present study could be adopted for pulmonary toxicity studies on lunar dust and Martian dust.

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