Space debris generation in GEO: Space materials testing and evaluation

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\textbf{A B S T R A C T}

The aim of this work is to evaluate what happens to the spacecraft materials beyond the spacecraft End of Life. A review of spacecraft external materials and effects of space environment is presented. This paper results from a continued study on spacecraft material degradation, and space debris formation in geostationary orbit (GEO). In this paper a 20-year GEO dose profile that combines simultaneous UV, particles irradiation and thermal cycling was applied to a set of external spacecraft materials. These materials comprised MLI assemblies, Velcros fixation and spacecraft painting. The evaluation of these external spacecraft materials, exposed to simulated space environment have confirmed the criticality of degradation of MLI, Velcros fixation and spacecraft painting. The evaluation of these external spacecraft materials, exposed to simulated space environment have confirmed the criticality of degradation of MLI, Velcros fixation and spacecraft painting. The synergy of space radiation (particles, UV) and thermal cycling ages the material and induces mechanical stress, causing creation of brittle surfaces, cracks and delamination. These phenomena cause serious damage to exposed surfaces, changing the surfaces thermo-optical properties, and may induce the generation of space debris. In particular, experimental results show the delamination of internal MLI layers and the severe degradation of the Velcros.

1. Introduction

Since the first man-made satellite launched in 1957 by Soviet Union, space activities have become more and more intensive. Human space exploration activity in the near-Earth space (LEO and GEO) has left behind a tremendous number of debris that poses a potentially significant danger to active satellites, spacecraft and people working in extravehicular activity \cite{1-4}. The number of space debris in orbit, consisting of bodies of disparate sizes, has increased. Examples of such debris are decommissioned satellites, rocket bodies (e.g. rocket upper stages, adapter rings), spacecraft external materials as paint flakes, payload shrouds, explosive bolts remains, instrument covers, slag particles \cite{5,6} (see Fig. 20).

Potential colonization of Mars and the Moon puts an additional pressure on how we manage the near-Earth space, since in the future we may need this space to deploy long-living bases and transition hubs to facilitate growing traffic from the Earth to extra-terrestrial outposts (see Fig. 19).

In this paper we address the formation of space debris and the factors that cause them, in particular for multilayer insulation materials, in particular when employed in a geostationary orbit (GEO). The main contributions are: (i) A review on spacecraft materials and effects of space environment on the materials; (ii) Description of the testing implementation for a high dose profile that simulate a 20 year GEO mission; (iii) Evaluation of radiation effects on the materials (mechanical properties and TO); (iv) identification of severe degradation of MLI internal layers and MLI Velcros; (iv) Size distribution of debris for the degraded MLI layers.

The paper is structured as follows: Section 2 introduces the previous research on the system. Section 3 explains the experimental methods. Section 4 presents the experimental results. Finally, Section 5 outlines conclusions.

2. State of the art

Today, space debris has become a huge concern for orbital missions that makes remediation a critical and necessary action \cite{7}. The spatial density of space debris is often used to define the number of resident space objects (RSO) per unit volume as function of the altitude \cite{8} as

\begin{equation}
\text{Density of RSO} = \text{Density of Space Debris} \times \text{Size Distribution of Debris for the Degraded MLI Layers}.
\end{equation}
The main degradation mechanisms that can be observed are: external spacecraft materials exposed to space environment. As a summary, depending on the environment and the type of family of materials, the spatial density is shown in Fig. 1.

Fig. 1 shows that the spatial density is higher in Low Earth orbit (LEO) and geostationary orbit (GEO). It can also be observed that the spatial density is slightly higher in GEO regime than it is in LEO. Note that RSO spatial density depends also on orbit altitude and declination. GEO spacecrafts or satellites are exposed to the outer radiation belts, solar flares and cosmic rays. Spacecrafts in GEO orbits are near the magnetopause, hence they are susceptible to high energetic plasmas during geomagnetic sub storms (in the tail of magnetosphere) [9].

In previous research we have reported a study on generation of space debris in simulated GEO conditions [10,11]. Also, we have reported a first incite on a study on generation of space debris in simulated LEO conditions [12].

Spacecraft external materials play an important role in satellite protection from space environment. In the past there have been cases of satellites failures due to the satellite external material degradations and respective change in thermo-optical properties. Midori – II, an Earth observation (EO) satellite, has stopped working in 2003 possibly due to degradation of X-ETFE polymer (change in the thermo-optical properties) due to UV irradiation leading to erosion of surface materials.

With operational conditions combining these constraints, synergy will lead to enhanced degradation and potential generation of debris. In this problematic, surface (thermo-optics parameters) and mechanical properties are of prime importance.

In GEO, the synergy of space radiation, i.e. electrons, protons, UV and thermal constraints is considered as main contributor to the degradation-induced debris process with the following mechanisms:

- Temperature: mechanical stress due to thermal cycling (TC), higher scission/x-linking ratio or colour centres bleaching at temperature higher than room temperature (RT);
- UV (affect mostly polymers): bonds breaking and rearrangement (photolysis), excitation and ionization;
- Charged particles: ionization (bonds breaking, radicals and colour centres depending on material type), displacement damage (radiolysis), physical sputtering (solar wind), blistering (metals only, and for large fluency);
- Atomic Oxygen (AO) affects polymers, metals, thin coatings: oxidation leading to erosion of surface materials.

2.1. Degradation factors and experimental constrains

To evacuate the materials for space applications it is usual to perform a series of qualification tests; however, the implementation of this test is subject to experimental constrains reported in this section:

- Air vacuum testing considerations

The presence of oxygen during irradiation induces a mechanism of oxidation reactions and the formation of free radicals in the material. This mechanism impacts the scission/cross-linking rate of the materials. It can be observed an increase, of several orders of magnitude, of polymer degradations, between radiation response in air (worst case) and under vacuum.

Annealing mechanisms are also affected by oxygen during vacuum-to-air transitions with kinetics varying from minutes to hours [14,15].

Ground testing of surface materials (particles exposures) requires vacuum conditions (<10⁻⁷ mbar), so that contamination of surfaces is avoided.
Temperature

Temperature is a major parameter in the process of material degradation when acting in synergy with other components of space environment, especially radiation.

For organic materials, degradation increases with temperature due to greater chain mobility (higher scission/x-linking ratio) [15]. In inorganic materials, temperature governs annealing of coloured centres as observed in optics [16].

At low temperature, synergy with space radiation is not straightforward and depends on material type [17,18]: degradation mechanisms are “frozen” in materials sensitive at room temperature while no significant change is observed in more resistant materials (epoxy, polyimide); silicones are more sensitive at low temperatures.

For missions with strong temperature constraints (range and cycling), it was shown that the representative simulation of the mission profile was required to include a reliable estimate of materials degradation and prediction of End-of-Life performances [19].

Moreover, at macroscopic level, temperature cycling (alone or in synergy with space radiation) induces mechanical stress that can result in enhanced degradation either overall or at stress location [20,21].

The temperature range for ground simulation (i.e. material testing) is usually larger than operational conditions, but the number of cycles is usually lower. The higher temperature range can be considered an accelerated degradation mechanism.

- **Particle dose rate, UV flux on ground simulation (i.e. radiation exposure acceleration impact)**

Particulates and UV acceleration at ground facilities are strongly different due to different technical limitations. Acceleration may induce abnormal dose rate effects in materials. When particles irradiation takes place in air, dose rate effects have been observed (the lower the dose rate, the higher the oxidation rate).

In vacuum, some cases of dose rate effect have been observed, affecting measurements on volatile products emission, but the impact on material properties at macroscopic level is not significant [22]. White paints with the higher acceleration are the worst-case condition [23].

Dose rate effects with charged particles are very material-dependent and may be associated with parasitic mechanisms, such as over-heating of samples. Sample over heating due to large flux may lead to recovery effects that are not representative of orbit conditions (underestimate of degradation).

Regarding UV/VUV, only few papers in the literature report on acceleration effect on material response and degradation [24–26], and this over a limited range of solar constant (SC). Most of the existing solar or UV simulator provides acceleration factors in the 2–15 SC range. UV acceleration is usually limited, as the light absorption in the exposed material creates a temperature increase leading to thermal effects. That is the reason why norms and standards define a maximum UV acceleration [27–29].

Regarding VUV, high acceleration can be obtained (in the 115 nm–180 nm range) with deuterium lamps for instance. VUV only affects surface properties.

Note that, from the debris generation standpoint, UV acceleration does not seem to be much relevant (unless participating to the sample temperature local or global increase); the total Equivalent sun hours (ESH) seen by the sample (total dose) are on the other hand very relevant.

- **GEO particles dose profile**

The radiation dose on materials drops rapidly with depth. So, representativeness of dose profile is a critical parameter from the thermo-optic properties and for coating adhesion (i.e. dose at coating/substrate interface).

In orbital conditions, surface dose is deposited by VUV/UV, protons with energy lower than 0.5 MeV and electrons with energy lower than 100 keV. At ONERA, the typical GEO dose profile is simulated as disclosed in Fig. 2 (with 2 proton beams and 1 electron beam).

In Fig. 2 GEO is the total cumulative dose in a one-year GEO orbit, calculated by ONERA. The electromagnetic radiation only affects surface properties, as wavelength and attenuation length of most (opaque) materials leads to dose deposition in the first microns (or less) [30] as shown in Fig. 3.

It is usually recommended to use Xenon lamps which provide representative solar spectrum and to filter the >400 nm range to avoid overheating of samples.

Dose equivalence can be applied between Xenon, Xenon–Mercury, and Mercury lamps integrating irradiance over the 200 nm–400 nm range. This equivalence is considered valid if the threshold energy for degradation is known [29].

- **Material degradation mechanisms and debris generation**

In MLI, fragments release is assumed to be described by a three-stage process: 1) Tear initiation in regions of highest stress and enhanced by material degradation (UV, AO or particle dose); 2) Tear propagation; 3) Foil separation.

In a simulation environment, using a representative dose profile, tear initiation can be reproduced on the 1st MLI layer, while the inner layer receives a lower dose than the one expected from the dose profile (otherwise inner layers would be over tested compared to the external). The implementation of a constant dose over the whole assembly set by surface dose (very strong in GEO) is technically difficult as it requires very large electron fluency (and thus an unrealistic long irradiation).

For paints, two mechanisms of debris generation can be considered: 1) ageing processes leading to the creation of a brittle surface layer, which may crack and lead to the generation of spalling off; 2) large temperature fluctuations leading to a thermal expansion of coatings and substrates at different rates. The creation and growth of cracks, caused by local thermal stress combined with AO, UV or particle dose, leads to the delamination of the coating layer.

In a simulated environment, a representative dose profile allows the reproducing of delamination mechanisms, at interface (actual dose at coating/substrate interface) and defect locations (actual surface dose).

- **Combined/sequential testing (particles, UV)**

Combined testing is justified in presence of synergy between different constraints. Simultaneous exposure of materials to particles
Ageing test facilities are not designed for thermal vacuum testing, and, depending on capabilities (temperature range and number of cycling), TC could be partly applied during ageing optimizing test duration.

2.2. Degradation of materials and failure mechanisms

Charged particles (trapped electrons and protons, solar event protons — x-rays on the side facing sun — and solar wind) are at the origin of the ionizing dose profile absorbed by the materials. Surface materials, as MLI and paints, are not shielded and therefore are subject to a worst-case environment for a given mission with important dose surface (>109 krad). Because of the energy spectra and mode of interaction of particles with matter, protons are responsible of surface dose (first µm) and electrons of bulk dose. The dose in rad/year and for different orbits is presented in Table 1.

For orbits that cross trapped belts in GEO, the absorbed dose profile exhibits a strong variation with depth as protons are stopped at the very surface (<2 µm). The dose profile in GEO for 2 materials (1- Kapton in GEO orbit inclination 0 ± 160° Western longitude, 2- cerium glass polar orbit 800 km) is shown in Fig. 3.

In ionization process, an atom loses an electron and forms an ion (whatever the chemical link at play). At material level (organic), these excitation/ionization mechanisms will result in the formation of free radicals or ions. These reactive intermediates are capable of initiating chemical reactions which result in scission as well as in cross-linking reactions. The macroscopic effects are then linked to the accumulation with time of the ionizing dose (namely “cumulated effect”) inducing surface and volume phenomena.

The same particles are also at the origin of displacement dose (non-ionizing) leading to creation of interstitials-vacancies, single or cluster of defects. The macroscopic effect of such phenomena mostly concerns inorganic materials and large fluence.

UV is critical for surface materials facing the sun (AM0 solar spectrum, ASTM490) but also facing earth due to the albedo effect (about 30% AM0 in terms of received energy). AM0 solar spectrum is a standard solar spectrum used for space applications, i.e. solar spectrum without atmosphere.

The UV radiation spectrum comprises wavelengths of between 200 nm and 400 nm, which correspond to energies of between 600 kJ/mol and 300 kJ/mol. These energies are in the same range as the bond energies of many organic compounds. Chemical reactions are induced when specific functional groups absorb the UV radiation. Free radicals liberated in the excitation/ionization process will trigger further reactions (bonds breaking and rearrangement: photolysis). Signs of photodegradation include embrittlement (surface cracking), discolouration and loss of transparency.

2.3. External spacecraft materials

The typical external spacecraft materials are listed in Table 2.

One of the main tasks of paints is to support spacecraft thermal control, by the reflection or absorption of electromagnetic radiation from the sun and by radiative emission to space.

Spacecraft passive thermal design allows thermal energy channels along the exterior, so that appropriate temperatures are maintained in the various elements of the spacecraft. Also, another important aspect of

Table 1

<table>
<thead>
<tr>
<th>Orbit</th>
<th>Depth</th>
<th>0.2 µm</th>
<th>2 µm</th>
<th>20 µm</th>
</tr>
</thead>
<tbody>
<tr>
<td>LEO</td>
<td></td>
<td>1 × 10^7</td>
<td>2 × 10^7</td>
<td></td>
</tr>
<tr>
<td>HEO</td>
<td></td>
<td>1.2 × 10^6</td>
<td>1.2 × 10^6</td>
<td></td>
</tr>
<tr>
<td>GEO</td>
<td>3 × 10^8</td>
<td>3 × 10^8</td>
<td>1 × 10^8</td>
<td>1 × 10^8</td>
</tr>
<tr>
<td>Lagrange L1 &amp; L2</td>
<td>3 × 10^9</td>
<td>6 × 10^9</td>
<td>6 × 10^9</td>
<td>6 × 10^9</td>
</tr>
</tbody>
</table>
the paint is its electrically conductivity, forming a protection against the formation of electrostatic charges, and the resulting discharges (ESD).

Space environment conditions that most degrade paints are UV radiation, AO, particle radiation (electrons and protons), and thermal cycles. The dose profile, number of cycles, temperature limits, strongly depend on the spacecraft attitude and orbit during the mission.

The paint degradation by space environment will have a strong impact in the function of paint (i.e. TO properties).

Regarding paints applications, one can classify the paints by the following subclasses:

### Table 2
Typical external spacecraft materials.

#### Paints

<table>
<thead>
<tr>
<th>Product type</th>
<th>Sub-categories</th>
<th>Description</th>
<th>Materials examples</th>
<th>Examples of applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thermal control</td>
<td></td>
<td>Spacecraft thermal control, by optimizing the spacecraft exterior surfaces thermo-optical specifications.</td>
<td>- NASA white paint (NASDA-1049/101-S), - NOVA astro white, - S13GP L0-1,</td>
<td>- White ceramic thermal insulating coatings for BepiColombo. - Spacecraft exterior protection (radiation, UV and AO).</td>
</tr>
<tr>
<td>paints</td>
<td>White paint</td>
<td>Applied to control the absorption and emission of energy on exposed spacecraft surfaces.</td>
<td></td>
<td>- Black paints can be used in heat shields for micro and nanosatellites to protect satellites when being transferred from orbit.</td>
</tr>
<tr>
<td></td>
<td>Black paints</td>
<td></td>
<td></td>
<td>- Thermal control coating. - Outer skin of spacecraft and launch vehicles.</td>
</tr>
<tr>
<td></td>
<td>High temperature paints</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Optical solar</td>
<td>- OSR</td>
<td>Flexible optical solar reflectors (OSRs) are used as satellite radiators. They have low absorption and high IR emissivity.</td>
<td>- FSR - multi-layered film ITO/CeO2/ polyetherimide/Ag/Ni alloy</td>
<td></td>
</tr>
<tr>
<td>reflector (OSR)</td>
<td>Rigid solar reflector (RSR)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Flexible solar reflector (FSR)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Second surface mirrors (SSM)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ATOX protection</td>
<td>NA</td>
<td>Protection from atomic oxygen. Can be applied on top of electronics as protective coating.</td>
<td>MAPATOX K, MAPATOX 41B, silicon paints, ceramer paints (part ceramic part polymer)</td>
<td>Protection from ATOX.</td>
</tr>
<tr>
<td>coatings</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thermal control</td>
<td>NA</td>
<td>Single polymeric foil with a metallic coating.</td>
<td>Aluminium coated MylarTM, TeonexTM, UPLEX, Colourless polyimide</td>
<td>Parabolic antennas, solar wings, and thermal control systems.</td>
</tr>
<tr>
<td>foil</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thermal control</td>
<td>NA</td>
<td>Its purpose is to thermal isolate the satellite, acting as radiation barrier and decreasing heat losses. Stack of thin polymeric layer (5-25 foils), separated by spacer or mesh or embossed, crinkled, external and innermost layers differ from the inner ones.</td>
<td>Aluminised-Teflon (AL-FEP) Polymide (Pi) films (e.g. Kapton), MylarTM</td>
<td>Hubble space telescope thermal shields.</td>
</tr>
<tr>
<td>blanket (i.e. MLI)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adhesives</td>
<td>NA</td>
<td>Structural gluing of satellite parts.</td>
<td>DC 93-500, RTV-5690</td>
<td>Solar cell adhesive, encapsulate, potting.</td>
</tr>
</tbody>
</table>

### Table 3
Lists of some of the current space grade paints.

#### MLI materials

<table>
<thead>
<tr>
<th>Manufacturer</th>
<th>Product</th>
<th>Base Composition</th>
<th>Colour</th>
<th>Absorptance</th>
<th>Emittance</th>
<th>Outgassing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lord Corporation</td>
<td>Chemglaze H32</td>
<td>Polyurethane</td>
<td>Matt-black</td>
<td>0.95</td>
<td>0.85</td>
<td>TML = 1.6%, RML = 0.9%, CVCM = 0.04%</td>
</tr>
<tr>
<td>Lord Corporation</td>
<td>Chemglaze L300</td>
<td>Polyurethane</td>
<td>Matt-black</td>
<td>0.955</td>
<td>0.85</td>
<td>TML = 1.1%, RML = 0.9%, CVCM = 0.04%</td>
</tr>
<tr>
<td>Lord Corporation</td>
<td>Chemglaze Z306</td>
<td>Polyurethane</td>
<td>Black</td>
<td>0.95</td>
<td>0.9</td>
<td>TML = 1.5%, RML = 0.6%, CVCM = 0.03%</td>
</tr>
<tr>
<td>Acheson Colloïden</td>
<td>Electrodag 501</td>
<td>Fluorinated binder</td>
<td>Black</td>
<td>0.965</td>
<td>0.829</td>
<td>TML = 0.96%, RML = 0.44%, CVCM = 0.00%</td>
</tr>
<tr>
<td>Acheson Colloïden</td>
<td>Electrodag 503</td>
<td>Fluorocarbon binder</td>
<td>Silver</td>
<td>0.37</td>
<td>0.44</td>
<td>TML = 0.22%, RML = 0.21%, CVCM = 0.06%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>PBZ</td>
<td>Silicone/Metallic</td>
<td>Matt-White</td>
<td>0.2 ± 0.04</td>
<td>0.83</td>
<td>TML = 0.55%, RML = 0.52%, CVCM = 0.08%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>PCEA</td>
<td>Silicone</td>
<td>Matt-White</td>
<td>0.25 ± 0.29</td>
<td>0.88</td>
<td>TML = 1.15%, RML = 0.44%, CVCM = 0.03%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>PSB</td>
<td>Potassium Silicate</td>
<td>Matt-White</td>
<td>0.14 ± 0.02</td>
<td>0.88</td>
<td>TML = 3.04%, RML = 0.04%, CVCM = 0.00%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>PSBN</td>
<td>Silicate</td>
<td>Matt-White</td>
<td>0.15 ± 0.02</td>
<td>0.92 ± 0.02</td>
<td>RML = 0.29%, CVCM = 0.00%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>MAP – PU1</td>
<td>Polyurethane</td>
<td>Matt-Black</td>
<td>0.96 ± 0.02</td>
<td>0.89</td>
<td>RML = 0.56%, CVCM = 0.09%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>PNC</td>
<td>Silicone</td>
<td>Matt-Black</td>
<td>0.98 ± 0.02</td>
<td>0.91 ± 0.03</td>
<td>TML = 0.91%, RML = 0.52%, CVCM = 0.00%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>MAP – PUC</td>
<td>Polyurethane</td>
<td>Black</td>
<td>0.94 ± 0.02</td>
<td>0.8</td>
<td>TML = 0.83%, RML = 0.55%, CVCM = 0.02%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>MAP – SG120FD</td>
<td>Silicone</td>
<td>Matt-White</td>
<td>0.17 ± 0.04</td>
<td>0.87 ± 0.02</td>
<td>TML = 0.95%, RML = 0.54%, CVCM = 0.04%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>SG121FD</td>
<td>Silicone</td>
<td>Matt-White</td>
<td>0.17 ± 0.02</td>
<td>0.88 ± 0.03</td>
<td>TML = 1.30%, RML = 0.28%, CVCM = 0.08%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>SG122FD</td>
<td>Silicone</td>
<td>Matt-White</td>
<td>0.18 ± 0.02</td>
<td>0.90 ± 0.03</td>
<td>RML = 0.74%, CVCM = 0.07%</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>SK5</td>
<td>Silicone</td>
<td>Matt-White</td>
<td>0.29 ± 0.02</td>
<td>0.89 ± 0.04</td>
<td>–</td>
</tr>
<tr>
<td>Société MAP S.A.</td>
<td>PUK</td>
<td>Polyurethane</td>
<td>Matt-Black</td>
<td>0.97 ± 0.02</td>
<td>0.91 ± 0.03</td>
<td>RML = 0.56%, CVCM = 0.00%</td>
</tr>
<tr>
<td>Akzo Nobel Aerospace Coatings</td>
<td>P5G 120 FD</td>
<td>Silicone</td>
<td>White</td>
<td>0.19</td>
<td>0.88</td>
<td>TML = 0.56%, RML = 0.54%, CVCM = 0.03%</td>
</tr>
<tr>
<td>ALION</td>
<td>Z93P</td>
<td>Silicate</td>
<td>White</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Previous IITRI (USA)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>–</td>
</tr>
<tr>
<td>ALION</td>
<td>SG13GP6N</td>
<td>Silicone</td>
<td>White</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>
- Other paints (e.g. metal paints). These include metal plating of gold, silver or even copper, helping to protect internal electronics by reflecting UV and infrared radiation [35].

The list of some of the current space grade paints is presented in Table 3.

Multilayer insulation (MLI) material provides high performance and specifically addresses all modes of heat transfer through the basic design of the system. For ground applications MLI is often used inside of a vacuum chamber, eliminating gas convection, and minimizing gas conduction to the molecular scale. Reflective shields are used to minimize radiation heat transfer inversely proportional to the number of shields. Low conductivity spacers are used to prevent the metallic based reflective shields from touching; they also minimize the conduction through the blanket itself. Much care is taken to design the MLI blanket such that it minimizes heat transfer in every manner including edges, seams, and installation procedure.

The environmental effects that cause degradation of the MLI are primarily temperature cycling, solar radiation, particle radiation, and AO. Although VUV, from solar radiation, is abundantly present at all altitudes, AO is particularly a problem in LEO. The MLIs are made from polymer films; the AO reacts with the organic species and erodes the polymer. This phenomenon is known as AO erosion and is measured in mass loss.

The key MLI specifications and material characteristics that are correlated with the MLI materials degradation, and therefore also correlated to debris generation, are TO properties; MLI stack configuration in terms of number, thicknesses, and materials of the layers; operational temperatures.

MLI application methods

Regarding MLI fixation methods, the most common fixation methods are:

1) Stand-offs/clipwasher fixation with the stand-offs bonded or screwed to the structure. This method is considered the most reliable fixation method. Stand-offs is available in different dimensions and can be custom made to the specific application.

2) Velcro fixation: Velcro stapled to blanket, and counterpart glued to structure using pressure sensitive adhesive tape (PSA) or adhesive.

3) Bonding of the blanket directly to structure by means of pressure-sensitive adhesive tape (PSA).

An example of MLI application is shown in Fig. 4.

In Fig. 4 (A) are the staples; (B) is the cover tape; (C) are the staples; (D) the blanket #2; (E) is the Velcro type 2; (F) is the adhesive; (G) is the Velcro type 1; (H) is the blanket #1.

3. Experimental methods

A simulation of a 20-year of GEO dose profile was performed at ONERA, DESP, Department of Space Environment, France. The simulation combines particles and UV doses in synergy with thermal cycling to investigate potential generation of debris from the ageing of surface materials.

3.1. Tested materials and mounting on sample plate

- MLI test samples

Two MLI layups strips of 44 mm × 120 mm were prepared for the GEO test campaign: one MLI layup with an outer layer of black Kapton (used in satellites with optical instruments), and another layup with an outer layer of Kapton. The MLI layups materials are depicted in in Fig. 5.

In Fig. 5 (a): (A) is 1 × 1 mm of Kapton; (B) is VDA coating; (C) is 5 × Dacron spacer plates; (D) is 5 × 0.3 mm Kapton perf.; (E) is 5 × Polyester Fleece Spacer PV-8g; (F) is 5 × 0.25 mm Mylar perf.; (G) is 1 × Polyester Fleece Spacer PV-8g; (H) is 1 × 1 mm Mylar foils.

In Fig. 5 (b): (I) is 1 × 1 mm of black Kapton XC; (B) is VDA coating; (C) is 5 × Dacron spacer plate; (D) is 5 × 0.3 mm Kapton perf.; (E) is 5 × Polyester Fleece Spacer PV-8g; (F) is 5 × 0.25 mm Mylar perf.; (G) is 1 × Polyester Fleece Spacer PV-8g; (H) is 1 × 1 mm Mylar foils.

Two different fixation methods were implemented for each MLI strip. The MLI stirs were grounded during the test. The fixations were: Velcros (ref SJ3571 Nylon6.6, loops part only) attached with staples; and Stand-off with clip/washers in Vestel.

The implementation of the MLIs in the sample holder is depicted in Fig. 6.

- Paints test samples

The selected paints are used in satellites’ exteriors; different substrates and base compositions were considered. The tested paints (procured at MAP), with dimensions 19.8 mm × 19.8 mm, are presented in Table 4.

SG121FD is white silicone paint and is a frequently used paint in satellites. PSB is white silicate paint; it was applied on two different substrates (AUG4 and CFRP). PNC is black silicone paint and is used in satellites that have optical instruments.

- Sample holder

The MLI and painting samples as mounted on sample plate are shown in Fig. 6.

In Fig. 6 (A) are the PT100 wires; (B) are the clip/washers; (C) are the Velcros/staples; (D) is the MLI2; (E) is the MLI1; (F) is the PSB/CFRP; (G) is the PNC/Au4G; (H) is the PSB/Au4G; (I) is the SG121FD/Au4G; (J) is the ground connection. In Fig. 6 MLI layups strips size is 44 mm × 120 mm. Paint samples size is 20 mm × 20 mm. MLI are attached loosely on the adaptor plate with the clip/washers (hanging) and maintained in contact with it with Kapton tape at the bottom edge. The lateral edges of the MLI have been left open. The two ends are closed by staples and Kapton tape (when mounted on sample holder). The MLI strips are grounded to avoid charging effects.

Painting samples have their substrate (back side) in good thermal contact with the sample plate and pressure applied with fastener bars on front side. For avoiding mechanical stress to the coating, masking during painting allowed to leave fixation zones free of coating.

The sample holder is heated through a resistor and cooled down with LN2.
3.2. Experimental procedures on test preparation

- **Contamination**

Precautions (overall cleanliness of facilities, bake-out of sample plate) were taken to avoid and monitor contamination (intrinsic contamination or from porous samples and degradation products). A crystal quartz microbalance was also used during the test campaign. Only 20 Hz of difference were measured in situ, which is within the noise-signal margin and thus shows the absence of contamination.

- **Samples handling and storage**

The presence of dust on sample surface and damage due to physical contact with the sample is very critical here. Therefore, special precaution has been taken, namely: handling of samples with gloves and adequate tools to avoid damaging the samples (handling being limited to the minimum); storage at room temperature under nitrogen; use of polyethylene or polypropylene bags for protection.

- **Temperature calibration prior to test campaign**

Calibration of sample/holder temperature prior to the test campaign was performed. The objective was: to validate parameters set for heating/cooling rates and dwell times; to measure MLI external foil temperature under UV exposure (during electrons/protons irradiations, the power brought by the exposure is typically few mW/cm² whereas UV brought up to 100 mW/cm²).

Spare samples used for this calibration step were fitted with temperature sensors, to monitor the sample plate and MLI foils temperatures. Painting temperature is equal to sample plate temperature due to the good thermal contact ensured by the mounting principle. The temperatures measured are presented in Table 5.

Samples are irradiated with 6 suns UV resulting in a heating of the outer layer to approximately 100 °C (measured temperature). Temperature was measured with a PT100 (uncertainty of temperature of this sensor typically in the 0.1 °C to 1 °C range).

The test configuration of layers considered for the calculations is shown in Fig. 7.

![Fig. 7](image)

In Fig. 7 (A) is 1 × 1 mm of Kapton; (B) is VDA coating; (C) is 5 × Dacron spacer plate; (D) is 5 × 0.3 mm Kapton perf; (E) is 5 × Poly-ester Fleece Spacer PV-8g; (F) is 5 × 0.25 mm Mylar perf.; (G) is 1 × Polyester Fleece Spacer PV-8g; (H) is 1 × 1 mm Mylar foils Δt (t > 70) = 36 °C; (I) is stand-off/clip-washer; (J) is Kapton tape; (K) is −85 °C to +100 °C TC with 4 h periods; (L) is Velcros Δt (t > 50) = 54 °C; (M) is ≈100 °C at T0 (alpha = 0.35).

### Table 4

<table>
<thead>
<tr>
<th>Samples</th>
<th>Paint Name</th>
<th>Substrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>SG121FD (silicone white)</td>
<td>Au4G</td>
</tr>
<tr>
<td>2</td>
<td>PSB (silicate white)</td>
<td>Au4G</td>
</tr>
<tr>
<td>3</td>
<td>PSB (silicate white)</td>
<td>CFRP</td>
</tr>
<tr>
<td>4</td>
<td>PNC (silicone black)</td>
<td>Au4G</td>
</tr>
</tbody>
</table>

### Table 5

<table>
<thead>
<tr>
<th>Irradiation</th>
<th>MLI 1 (Kapton) (°C)</th>
<th>MLI 2 (black Kapton) (°C)</th>
<th>Plate (°C) Set to</th>
</tr>
</thead>
<tbody>
<tr>
<td>UV (6 suns)</td>
<td>98</td>
<td>92</td>
<td>25</td>
</tr>
<tr>
<td></td>
<td>102</td>
<td>94</td>
<td>100</td>
</tr>
</tbody>
</table>
3.3. GEO test specifications

Considering a 20-year GEO mission, beams fluences are $2 \times 10^{16}$ with 400 keV electrons/cm$^2$, $4 \times 10^{15}$ with 240 keV protons/cm$^2$, and $4 \times 10^{16}$ with 45 keV protons/cm$^2$, leading to an overall 3 week–4 week exposure with ionizing particles. The execution of the irradiations and the test sequences with details of beam currents, particle fluences and homogeneity factors (standard deviation/mean fluence) are shown in Fig. 8.

During the testing, 2 failures happened:

1) Failure of the Minco heater at the early stage of the testing changed for a thermo-coax device considered as more reliable. The mending step required a venting/pumping step that could not affect the final results as failure happened very early in the test campaign (520 ESH, i.e. equivalent sun hours).

2) Temperature regulation failure: the electronics in charge of thermal regulation (TC program with defined parameters) failed probably due to dose effect (electron irradiation produces secondary X-rays in the room). The failed board was changed and the automatic system shielding against secondary radiation improved.

During the failure, the samples’ temperature returned to positive level (~60 °C in absence of UV) during about 72 h (failure at week-end). This failure too could not affect final results (no overheating of samples, short duration compared to total duration of testing).

3.4. GEO samples test flow

The GEO MLI/paint samples test flow is shown in Fig. 9.

<table>
<thead>
<tr>
<th>Transition</th>
<th>5 years</th>
<th>10 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air</td>
<td>Vacuum</td>
<td>replacement with thermo-coax</td>
</tr>
<tr>
<td>26-oct</td>
<td>03-nov</td>
<td>13-nov</td>
</tr>
<tr>
<td>TC (cycles)</td>
<td>heater failure</td>
<td></td>
</tr>
<tr>
<td>020</td>
<td>60</td>
<td>100</td>
</tr>
<tr>
<td>UV (ESH)</td>
<td>19</td>
<td>19</td>
</tr>
<tr>
<td>240keV</td>
<td>15-19</td>
<td>15-19</td>
</tr>
<tr>
<td>4E15</td>
<td>1E15</td>
<td>1E15</td>
</tr>
<tr>
<td>Protons</td>
<td>21.5</td>
<td>24E15</td>
</tr>
<tr>
<td>240keV</td>
<td>10,4%</td>
<td>21%</td>
</tr>
<tr>
<td>4E15</td>
<td>21%</td>
<td>21%</td>
</tr>
<tr>
<td>Protons</td>
<td>23.4</td>
<td>27.2</td>
</tr>
<tr>
<td>45keV</td>
<td>1E16</td>
<td>1E16</td>
</tr>
<tr>
<td>4E16</td>
<td>15%</td>
<td>10%</td>
</tr>
<tr>
<td>visual inspection</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td></td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

Fig. 7. Test configuration of layers considered for the calibration.

Fig. 8. Test sequence with each beam current (defines flux: $1 \text{nA/cm}^2 = 6.4 \times 10^9 \text{part.}/\text{cm}^2$.s), deposited fluence and homogeneity (for protons; with electrons homogeneity is ±10%).
In Fig. 9 the text box “In situ visual inspection” addresses the initial paintings and MLI inspection and evolution of thermo-optical properties, which is included inspection of any self-flaking, discolouration of white paints and photos; the text box “Ex situ visual inspection” addresses ex situ visual and mechanical properties inspection, which includes visual and mechanical properties inspection after dismounting from sample plate, provides the same outputs as in situ ones; the text box “Mass measurement” addresses the measuring the mass loss of paintings and MLI; the text box “Peeling” addresses the peeling test to evaluate the mass loss of paintings and MLI. Final sample characterization is the final inspection of the material sample for evaluation by photos and mass determination.

4. Experimental results

4.1. In situ visual inspection and evolution of thermo-optical properties

In situ observation of painting did not reveal any self-flaking, but discolouration of white paints mostly. Fig. 10 discloses pictures showing evolution of yellowing/browning of the white paints and the very slight whitening of the black one (20-year picture). This discolouration effect is due to the shift of the reflectance cut-off in the UV part of the spectrum (modification of chemical structure induced by radiation: scission/cross-linking of chains in binder, coloured centres in pigments) leading to increased solar absorptance. The Evolution of solar absorptance in the range from 250 nm to 2500 nm is presented in Table 6.

<table>
<thead>
<tr>
<th>Sub-system</th>
<th>Initial</th>
<th>20 year Geo</th>
<th>Delta</th>
<th>Delta (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SG121FD/Au4G</td>
<td>0.233</td>
<td>0.626</td>
<td>0.393</td>
<td>169</td>
</tr>
<tr>
<td>PSB/CFRP</td>
<td>0.142</td>
<td>0.659</td>
<td>0.517</td>
<td>365</td>
</tr>
<tr>
<td>PSB</td>
<td>0.131</td>
<td>0.623</td>
<td>0.491</td>
<td>374</td>
</tr>
<tr>
<td>PNC</td>
<td>0.973</td>
<td>0.907</td>
<td>−0.006</td>
<td>−1</td>
</tr>
<tr>
<td>Black MLI</td>
<td>0.930</td>
<td>0.880</td>
<td>−0.050</td>
<td>−5</td>
</tr>
<tr>
<td>MLI</td>
<td>0.350</td>
<td>0.680</td>
<td>0.330</td>
<td>94</td>
</tr>
</tbody>
</table>

Table 6
Evolution of solar absorptance in the range values 250 nm–2500 nm.

End-of-Life alpha are consistent with existing data (orders of magnitude) for lower but equivalent dose levels [36,37].

The first consequence of solar absorptance increase is the potential overheating of MLI external foil during the test campaign (evolution of thermal conditions of the assembly).

Table 7
Evolution of emissivity in the range values 3 μm–21 μm.

<table>
<thead>
<tr>
<th>Sub-system</th>
<th>Initial</th>
<th>20 year Geo</th>
<th>Delta</th>
<th>Delta (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SG121FD/Au4G</td>
<td>0.895</td>
<td>0.868</td>
<td>−0.027</td>
<td>−3.0</td>
</tr>
<tr>
<td>PSB/CFRP</td>
<td>0.886</td>
<td>0.897</td>
<td>0.011</td>
<td>1.2</td>
</tr>
<tr>
<td>PSB/Au4G</td>
<td>0.895</td>
<td>0.902</td>
<td>0.007</td>
<td>0.8</td>
</tr>
<tr>
<td>PNC</td>
<td>0.906</td>
<td>0.901</td>
<td>−0.005</td>
<td>−0.6</td>
</tr>
<tr>
<td>Black MLI</td>
<td>0.839</td>
<td>0.798</td>
<td>0.041</td>
<td>4.9</td>
</tr>
<tr>
<td>MLI</td>
<td>0.660</td>
<td>0.677</td>
<td>−0.017</td>
<td>−2.6</td>
</tr>
</tbody>
</table>

Table 7
Evolution of emissivity in the range values 3 μm–21 μm.

Fig. 10. Visual inspection (i.e. photos during irradiation).
The emissivity, known to be less sensitive, has also been measured, and is presented in Table 7.

Paint photos during irradiation is shown in Fig. 10.

The reflectance curves of the paintings (% vs wavelength) is shown in Fig. 11.

4.2. Ex situ visual inspection and mechanical properties

The sequence for testing is the following:

1) Visual inspection of samples (pre-peeling, after TO measurements);
2) Mass measurement;
3) Peeling: 3.1) Compress tape onto sample with load of nominal 5 kg for 60 s; 3.2) pull apart tape and sample at a nominal 0.2 cm/min until separation happens;
4) Mass measurement;
5) Visual inspection of samples and tapes (post peeling).

Ex situ visual inspection (after dismounting from sample plate) provides the same outputs as in situ ones. The ex situ pictures of the paintings before (right), after 20-year GEO (left) are shown in Fig. 12. (exposed to 20 Years) (paints before test).

Peeling test has been performed to investigate adherence of coatings and check for degradation of peel and pull-off strength. The test was performed with PSA (pressure sensitive adhesive) tape.

There was no delamination on the SG121 samples and PNC samples. The PSB paints sample and tapes before/after peeling are shown in Fig. 13 and Fig. 14.

For the PSB paint on aluminium, there was no sample fracture, but detachment of particles (pristine and aged) was observed. A slight enhancement of particle detachment in the central part of the aged sample was observed.

Fig. 14 shows the PSB on CFRP sample and tapes before/after peeling.

For the PSB/CFRP coating, substrate fracture in the central part (more strength) on both pristine and aged samples was observed. A larger fracture size is observed on pristine sample due to better adherence of coating surface (compared to aged sample where degradation occurred). Fracture comes from detachment of upper layers of CFRP substrate due to the bending of the sample at pulling step (“flexible” 0.5 mm substrate attached on opposite edges). Also, detachment of small parts was observed due to handling for dismounting from sample plate or TO measurements, illustrating brittleness of surface.

For the PNC samples, it was observed that peeling and pull-off strength remains good, and no flaking is observed, only particles detachment. However, the surface of painted samples turns “glassy” and

\[\text{Fig. 11. Reflectance curves of the paintings (% vs wavelength). (a) and (b) exhibit the same feature (same paint, PSB, different substrates); in (d) evolution of black silicone paint PNC is very limited (see Y scale).}\]
more brittle (Proton/UV irradiation induced surface degradation).

4.3. MLI test results

In situ visual inspection did not reveal any cracks or tear initiation of external foil, but strong discolouration of Kapton (strip1), clips/washers and curling of assemblies at edges (not close). There was no visible damage of Kapton tape used to close upper/lower ends. Fig. 15.

The curled edges of the MLI (not closed) due to thermal/radiation effects are shown in Fig. 16.

From ex situ visual inspection, it appeared that Mylar (internal layers) foils of MLI assemblies were so embrittled that they fell into pieces at handling. It was therefore decided to cut one edge of the strip to allow for flipping the assembly and take pictures foil by foil.

Then, each Mylar foil has been removed, piece by piece, and “reassembled” for pictures. Procedure for dismounting MLI for adaptor plate is shown in Fig. 17.

Figs. 18–22 shows all foils from strip 1 (see Fig. 21). It results that:

- Mylar foils are brittle (especially around Velcros);
- Polyester fleece is yellowed but not brittle;
- Kapton inner foils are not brittle;
- Dacron spacers are yellowed but not brittle;
- Kapton external foil is discoloured and not brittle;
- Velcros are bended, brittle and discoloured;
- Clip/washers are discoloured (darkening);
- Kapton tape used to close top/bottom strip ends is discoloured but not brittle.

The ex situ visual inspection of the Velcros revealed bending features, darkening and brittleness.

Bending can be due to combined temperature/dose effects. The vitreous transition temperature $T_v$ of Nylon is quite low (in the 47 °C-60 °C range). When temperature is greater than $T_v$ (probably in the duration of
the cycle above +35 °C, see section 2.4), polymer chains are more mobile and sensitive to dose-induced scission, then cross-linking leading to hardened and more brittle material. This effect also occurs at loops level.

Bending also induces local mechanical stress of attached foils, and then as Mylar is also strongly degraded a “cutting at edges” mechanism takes place (at handling).

Regarding thermo-optical properties of external foils, the shift of the cut-off of the reflectance curve for strip 1 is characteristic of darkening of Kapton (more absorption in the 500 nm–1000 nm range is shown in Fig. 23 b).

Regarding the black MLI, the whole spectrum is affected with higher reflectance which is characteristics of whitening of the black Kapton is shown in Fig. 23a.

### 4.4. MLIs: debris distributions

ImageJ, a public-domain image processing program, was used to extract debris area distribution from the set of pictures provided in section 4.3.

Fig. 24 discloses all distributions by layer for Kapton and Black Kapton MLI assemblies with “bins start” in mm² (debris area) and with Layer 1 the “back layer” (in contact to the sample holder).

As a reminder, this debris has been generated from handling and not from self-delamination or self-flaking (even if layer 1 from strip 1 was almost self-flaking). An uncertainty of 10%–20% is estimated on area measurements, as the debris could not be flattened between transparent plates due to high electrostatic conditions. The debris distribution by
layer is disclosed in Fig. 24.

5. Conclusions

Satellite material testing is usually done for the satellite expected lifetime. However defunct satellites in GEO are exposed to very long periods, beyond satellite lifetime. In this paper we study the effects of synergistic effects of space environment (electrons, VUV, protons, TC) on materials for long periods. The aim was to evaluate the creation of debris by degradation of the materials.

The materials studied were representative spacecraft GEO materials, namely: two MLI blanket samples (GEO-1 MLI layup and GEO-2 MLI) fixed to the sample holder with Nylon6.6 Velcros and Vestel standoffs and four paints SG121FD (silicone white paint) on Au4G, PSB (silicate white paint) on CFRP, PNC (silicone black) on Au4G.

A simulation of 20-year of GEO dose profile was performed combining particles and UV doses in synergy with thermal cycling. Beams fluences were: $2 \times 10^{16}$ with 400 keV electrons/cm$^2$, $4 \times 10^{15}$ with 240 keV protons/cm$^2$, and $4 \times 10^{16}$ with 45 keV protons/cm$^2$. The TC was 300 cycles, from $-85$ °C to $+100$ °C. UV simulation was provided with Xenon 6500 lamp whose spectrum is close to the solar spectrum, the sun acceleration was measured periodically and adjusted between 5.2 and 6.6 SC (variation due to lamp ageing) a total of 8211 ESH was done.

As to major results, in situ observation of painting did not reveal any self-flaking, but discolouration of white paints was observed. The discolouration of white paints showed an evolution to yellowing/browning. The evolution of solar absorptance in the paints and MLI external foil was measured. The major degradation was observed in SG121FD on Au4G (absorptance changed from 0.233 to 0.626), PSB on CFRP (absorptance changed from 0.142 to 0.659), PSB (absorptance changed from 0.131 to 0.623) and MLI (absorptance changed from 0.350 to 0.680). The first consequence of solar absorptance increase is the potential overheating of MLI external foil during the test campaign (evolution of thermal conditions of the assembly). Emissivity was also measured but changes were less relevant.

Adhesion test on paints have not indicate major differences in paint adhesion of the aged samples compared with the reference samples.

Regarding debris generation by material aging due to space environment, the more relevant results were obtained from MLI degradation. The external layers of the MLI layups were curled and TO properties changed significantly (i.e. burned up appearance). After disassembling the MLI layups it was found that the Mylar inner foils were brittle and white paint) on Au4G, PSB (silicate white paint) on CFRP, PNC (silicone black) on Au4G.

Fig. 15. Clip/washer pictures before/after GEO test campaign (in and ex situ): (a) Initial feature; (b) 20-year GEO Shadowing of MLI ext. layer underneath; (c) greater degradation at surface (p+).

Fig. 16. Curled edges of the MLI (not closed) due to thermal/radiation effects.

layer is disclosed in Fig. 24.

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Satellite material testing is usually done for the satellite expected lifetime. However defunct satellites in GEO are exposed to very long periods, beyond satellite lifetime. In this paper we study the effects of synergistic effects of space environment (electrons, VUV, protons, TC) on materials for long periods. The aim was to evaluate the creation of debris by degradation of the materials.

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As to major results, in situ observation of painting did not reveal any self-flaking, but discolouration of white paints was observed. The discolouration of white paints showed an evolution to yellowing/browning. The evolution of solar absorptance in the paints and MLI external foil was measured. The major degradation was observed in SG121FD on Au4G (absorptance changed from 0.233 to 0.626), PSB on CFRP (absorptance changed from 0.142 to 0.659), PSB (absorptance changed from 0.131 to 0.623) and MLI (absorptance changed from 0.350 to 0.680). The first consequence of solar absorptance increase is the potential overheating of MLI external foil during the test campaign (evolution of thermal conditions of the assembly). Emissivity was also measured but changes were less relevant.

Adhesion test on paints have not indicate major differences in paint adhesion of the aged samples compared with the reference samples.

Regarding debris generation by material aging due to space environment, the more relevant results were obtained from MLI degradation. The external layers of the MLI layups were curled and TO properties changed significantly (i.e. burned up appearance). After disassembling the MLI layups it was found that the Mylar inner foils were brittle and white paint) on Au4G, PSB (silicate white paint) on CFRP, PNC (silicone black) on Au4G.
Fig. 18. Mylar foils and polyester fleeces from strip1 (numbering from the back foil).

Fig. 19. Flipping the Kapton/dacron foils of strip1 (layer 7 to 12 numbering from the back foil).
Fig. 20. Mylar foils and polyester fleeces from strip2 (numbering from the back foil).

Fig. 21. Flipping the Kapton/dacron foils of strip2 (layer 7 to 12 numbering from the back foil).
Fig. 22. Velcros degradation; (a) degradation; (b) Darkening of Velcros (loops); (c) and (d) embrittled Velcros (broken parts at handling).

Fig. 23. Reflectance curves of the MLI external layers (R% vs wavelength): (a) black Kapton (strip2), (d) Kapton (strip1).
fall into pieces at handling. Extreme care was taken handling the brittle MLI layers, the Debris distributions for the layer was measured.

Regarding the MLI fixation methods the Velcros degraded extremely, becoming yellow and brittle. It may be possible that severe degradation of the Velcro leads to the separation of the full MLI blanket (if the MLI is only attached with Velcros) however to prove this hypothesis further study is needed.

These results are expected to contribute to a better practice of choice of materials when building satellites, to avoid the significant problems caused by debris.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Fig. 24. Debris distributions for all layers; bins start is in mm² (debris area): (a) Kapton layup, K1 is the first layer in contact with sample holder; (b) Black Kapton layup, BK1 is the first layer in contact with sample holder; Bins start is in mm² (debris area).

References


