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Lithologic variation within bright material on Vesta revealed by linear spectral unmixing

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1	Lithologic variation within bright material on Vesta
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# 22 Abstract

<sup>23</sup> Vesta's surface is mostly composed of pyroxene-rich lithologies compatible <sup>24</sup> with howardite, eucrite and diogenite (HED) meteorites (e.g., McCord et al., <sup>25</sup> 1970; Feierberg and Drake, 1980). Data provided by the Visible and Infrared <sup>26</sup> (VIR) spectrometer, onboard the NASA Dawn spacecraft, revealed that all <sup>27</sup> Vesta reflectance spectra show absorption bands at ~0.9  $\mu$ m and ~1.9  $\mu$ m,

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which are typical of iron-bearing pyroxenes (De Sanctis et al., 2012a). Other 28 minerals may be present in spectrally significant concentrations; these in-29 clude olivine and opaque phases like those found in carbonaceous chondrites. 30 These additional components modify the dominant pyroxene absorptions. 31 We apply linear spectral unmixing on bright material (BM) units of Vesta to 32 identify HEDs and non-HED phases. We explore the limits of applicability 33 of linear spectral unmixing, testing it on laboratory mixtures. We find that 34 the linear method is applicable at the VIR pixel resolution and it is useful 35 when the surface is composed of pyroxene-rich lithologies containing moder-36 ate quantities of carbonaceous chondrite, olivine, and plagioclase. We found 37 three main groups of BM units: eucrite-rich, diogenite-rich, and olivine-rich. 38 For the non-HED spectral endmember, we choose either olivine or a fea-39 tureless component. Our work confirms that Vesta's surface contains a high 40 content of pyroxenes mixed with a lower concentration of other phases. In 41 many cases, the non-HED endmember that gives the best fit is the featureless 42 phase, which causes a reduction in the strength of both bands. The anticor-43 relation between albedo and featureless endmember indicates that this phase 44 is associated with low-albedo, CC-like opaque material. Large amounts of 45 olivine have been detected in Bellicia, Arruntia and BU14 BM units. Other 46 sites present low olivine content (< 30%) mostly with a high concentration 47 of diogenite. 48

# 49 HIGHLIGHTS

50 51 • Characterization of Vesta's bright material (BM) units by linear unmixing analysis;

- Detection of different lithologies within BM regions;
- Detection of phases other than pyroxenes from VIR data on Vesta surface;
- Calculation of mixing coefficients for the detected lithologies for each
- in BM unit;

## 57 1. Introduction

## 58 1.1. Early knowledge of Vesta

The first spectral study of Vesta dates back to Bobrovnikoff (1929). Mc-59 Cord et al. (1970) compared Vesta spectra with the meteorite Nuevo Laredo, 60 inferring a relation between Vesta and pyroxenes. Pyroxenes are character-61 ized by the presence of two crystal field absorptions, centered approximately 62 at 0.9 and 1.9  $\mu$ m (McCord et al., 1970), due to ferrous iron (Fe<sup>+2</sup>) in octa-63 hedral sites (e.g., Burns, 1993). Consolmagno and Drake (1977) suggested a 64 possible link between Vesta and eucrite meteorites, and Feierberg and Drake 65 (1980) proposed that Vesta is a mixture of howardite and eucrite. Studies by 66 Lupishko et al. (1988) show a clear inverse correlation between the polariza-67 tion and brightness of Vesta, correlated with the west-east dichotomy. Hubble 68 Space Telescope (HST) observations between 1994 and 1996 confirmed this 69 inference by highlighting the geological differences on the asteroid (Binzel 70 et al., 1997). Moreover the identification of the prominent impact basin 71 at the south pole (now named Rheasilvia) corroborates the hypothesis that Vesta is the parent body of the howardite, eucrite and diogenite (HED) me-73 teorites (Thomas et al., 1997). In addition, the disk-integrated mid-infrared 74

spectra of Vesta have shown the presence of minor constituents as olivine, 75 feldspar, and chromite (Donaldson Hanna and Sprague, 2009). Studies by Li 76 et al. (2010) revealed that the vestan surface can be divided into several geo-77 logical units (more specifically, regions of lithologic variation), in particular, 78 eucrite-rich and diogenite-rich units, as well as slightly weathered and freshly 79 exposed units. Shestopalov et al. (2010) concluded that the global diversity 80 of Vesta might be caused by variations of non-basaltic source sediments en-81 riched by dark rocks, spinel group minerals, or even chondritic-like material. 82 Moreover, they showed that certain vestan units contain olivine abundances 83 of several volume percent. 84

## <sup>85</sup> 1.2. The Dawn-Era and Vesta's lithologies

Dawn entered orbit around Vesta in July 2011 (Russell and Raymond, 86 2011), and enabled many discoveries during about one year of observations. 87 For the first time, Dawn acquired data of Vesta's surface at high spatial res-88 olution, allowing for production of complete geological and lithological maps 89 (De Sanctis et al., 2012a; Williams et al., 2014). Dawn's Visible and Infrared 90 (VIR) imaging spectrometer (De Sanctis et al., 2011) provided high spectral 91 resolution data covering the majority of Vesta's surface at varying spatial 92 scales. The spatial resolution of the VIR maps ranges from  $\sim 800 \text{ m/pixel}$  in 93 the Survey phase to  $\sim 180$  m/pixel in the HAMO and HAMO-2 phases (see 94 Table 1 of Zambon et al. (2015)). 95

Vesta is the parent body of the HED meteorites. HEDs encompass a large variety of igneous rocks, similar to basalts, cumulate gabbros, orthopyroxenites
and igneous brecciated mixtures (Mittlefehldt et al., 1998). The diogenites
are coarse-grained cumulates that originated in a plutonic layer deep in the

crust (Mittlefehldt et al., 1998; Beck and McSween, 2010; McSween et al., 100 2011, 2013). The mineralogy of diogenites is dominated by orthopyroxenes 101 (from 87% to 99%); all diogenites contain < 5% chromite and some contain 102 olivine, typically at contents < 10% (McSween et al., 2011). Eucrites occur 103 as basaltic or cumulate rocks. They are dominated by Ca-poor pyroxenes 104 and plagioclase, with minor amounts of metal, troilite, chromite, ilmenite, 105 and silica (Mayne et al., 2010; McSween et al., 2011). Eucrites are believed 106 to have crystallized as lavas on the surface or within relatively shallow dikes 107 and plutons (McSween et al., 2011). Basaltic eucrites contain Fe-rich pyrox-108 enes, whereas cumulate eucrites are predominantly unbrecciated and their 109 chemistry is similar to basaltic eucrites, but richer in Mg (Mittlefehldt et al., 110 1998; McSween et al., 2011). Howardites are brecciated achondrites, prin-111 cipally mixtures of eucrite and diogenite clasts, and reflectance spectra of 112 howardites have pyroxene band center positions intermediate between those 113 of eucrites and diogenites (Mittlefehldt et al., 1998; McSween et al., 2011). 114 Impact mixing of eucrite and diogenite has produced the polymict breccias 115 and howardites. 116

Eucrite, diogenite and howardite lithologies are present on Vesta's surface, 117 as revealed by VIR data (De Sanctis et al., 2012a; Ammannito et al., 2013a) 118 augmented by Dawn Framing Camera images (Reddy et al., 2012b) and 119 Gamma Ray And Neutron Detector data (Lawrence et al., 2013; Prettyman 120 et al., 2011, 2013, 2014). Recently, many papers on the analysis of Vesta's 121 surface composition using Dawn data have been published (e.g., Ruesch et al. 122 (2014); Ammannito et al. (2013a); De Sanctis et al. (2013b); Le Corre et al. 123 (2013); Reddy et al. (2013); Thangjam et al. (2013)). The crust of Vesta is 124

dominated by howardite enriched in eucrite (De Sanctis et al., 2012a, 2013b). 125 A few outcrops of diogenite are present in localized areas in the south po-126 lar region, corresponding to the rim of the large impact basin Rheasilvia. 127 Large areas of diogenite-enriched howardite have also been identified in the 128 northern hemisphere (longitude  $0^{\circ} - 90^{\circ}E$ ) and are interpreted to consist of 129 Rheasilvia ejecta (De Sanctis et al., 2012a; Ammannito et al., 2013a), 130 Ammannito et al. (2013a) suggested a high content of olivine (50-80 vol.%) 131 in the area of Bellicia (lat 40°N, lon 40°E) and Arruntia craters (lat 40°N, lon 132 70°E), highlighting the presence of lithologies different from that of typical 133 HEDs. Others eleven sites containing much smaller amounts of olivine (lo-134 cated up to 39°S latitude) have also been identified by Ruesch et al. (2014), 135 and six other new olivine-rich regions were proposed by Palomba et al. (2015), 136 almost all located at latitudes below 28°N. Dark material units on Vesta 137 were discussed by Jaumann et al. (2012); McCord et al. (2012); Reddy et al. 138 (2012a), and Palomba et al. (2014). Hydrated mineral phases were shown to 139 be correlated with this low-reflectance material (De Sanctis et al., 2012b), 140 confirming that the dark material can be attributed to the presence of a 141 carbonaceous chondrite (CC)-like component delivered by impacts. 142

## 143 1.3. Bright material units: A slice of fresh material

The spectral analysis of the bright material units reveals that they are generally characterized by greater band depths than their surroundings. Most bright units have a howardite rich-eucrite, composition, with some exceptions such as the bright unit called "BU15" in Zambon et al. (2014).

BM units could represent pristine material that has been recently exposed
on the surface, and thus contains less carbonaceous-chondrite-like contami-

nation introduced by impact mixing than other areas of Vesta (see Zambon 150 et al. (2014) for more detail). To be consistent with Zambon et al. (2014), we 151 follow the same classification of the BM units, as described by Mittlefehldt 152 et al. (2012). The abbreviation CWM stands for crater wall material, SM 153 means slope material, and RM refers to radial material. The spectral anal-154 ysis employed in Zambon et al. (2014) to derive Vesta's lithologies is based 155 on selected spectral parameters, namely band center (BC), band depth (BD) 156 and Band Area Ratio (BAR). 157

The goal of our work is to use linear spectral unmixing to automatically 158 identify the main lithologies of specific regions and determine their semi-159 quantitative mixing coefficients, including identification and mapping of min-160 eralogical phases other than the spectrally dominant pyroxenes. The mixing 161 coefficients depend on various effects, such as, the grain size, the abundance 162 of the mineralogical phases present in the scene, and the scale at which the 163 components are mixed (i.e., macroscopic vs. intimate; Combe et al. (2008)). 164 Since the grain size on Vesta, at the VIR spatial scale, is quite homogeneous 165  $(< 25 - 45 \ \mu m)$  (Hiroi et al., 1994; Palomba et al., 2014; Zambon et al., 166 2014), we can assume that the grain size does not substantially affect our 167 derived mixing coefficients. Thus the unmixing results principally depend on 168 the abundance of the mineralogical phases and on the type of mixing. 169

Unmixing methods are useful for understanding the composition of a surface
(e.g. Pieters and Englert, 1993; Keshava and Mustard, 2002; Bioucas-Dias
et al., 2012). Mixing models can be either linear or non-linear. In the linear
case, the spectrum of a region can be considered the area-weighted average of
the endmembers present (Singer and McCord, 1979; Hapke, 1993; Bioucas-

Dias et al., 2012). The endmembers can be extracted from the scene, or can be reference spectra of plausible analogue phases. The endmembers are generally assumed to represent the different components, which are the fundamental constituents of the scene, so that each pixel in the scene can be modeled as a linear combination of the endmembers (Bioucas-Dias et al., 2012).

The case of non-linear mixing arises when light is scattered by multiple mate-181 rials in the scene. Scattered light interacts with more than one endmember, 182 and the resulting spectrum is a non-linear combination of the components. 183 Models for non-linear mixing require comprehensive information on the ex-184 pected minerals, such as scattering coefficients, particle sizes, and optical 185 constants (e.g., Hapke (1981); Shkuratov et al. (1999)). Here we use linear 186 unmixing as a first step in order to derive additional information on the com-187 position of Vesta's surface. 188

Many authors have linear unmixing algorithms for analysis of remote sensing 189 data for a variety of bodies (e.g., Adams et al. (1986); Fox III et al. (1990); 190 Ramsey and Christensen (1992); Blewett et al. (1995); Combe et al. (2008); 191 Dalton (2007); McCord et al. (2012)). Adams et al. (1986) already applied 192 linear unmixing methods to Viking lander data of Mars, while Dalton (2007) 193 applied linear unmixing to Europa data using a set of laboratory spectra as 194 endmembers, and Combe et al. (2008) used the Multiple-Endmember Lin-195 ear Spectral Unmixing Model (MELSUM) to analyze Mars data from the 196 Infrared Mineralogical Mapping Spectrometer (OMEGA) onboard MarsEx-197 press. A first use of linear unmixing methods on Vesta was presented by 198 McCord et al. (2012), who found that the entire surface of Vesta as observed 199

<sup>200</sup> by VIR could be modeled using the weighted sum spectral endmembers,
<sup>201</sup> 'bright' and 'dark' (McCord et al., 2012). In this paper, we apply the same
<sup>202</sup> linear unmixing method employed by Tosi et al. (2015) and Zambon et al.
<sup>203</sup> (2015) to extended areas of Vesta.

Here we considered high spatial resolution data from the Dawn mission at 204 Vesta, and we selected a set of plausible laboratory spectra analogues as 205 endmembers. We apply the linear unmixing method to the bright material 206 (BM) units of Vesta, previously analyzed by Zambon et al. (2014). BM units 207 are often associated with impact craters. Impacts expose fresh, unweathered 208 material from beneath the surface, which could be indicative of the original 209 vestan crust composition (Zambon et al., 2014). The previous spectral anal-210 ysis revealed the main lithologies (eucrite, diogenite and howardite) and the 211 spectral characteristics of these units. Here we study the same BM units with 212 the goal of quantifying the mixing coefficients of the different lithologies. We 213 also aim to identify the presence of other mineralogical phases in addition to 214 the spectrally dominant, HED components. In principle, this work can be 215 extended also to larger region on Vesta, determining the global distribution 216 of Vesta's non-HED lithologies. 217

# 218 2. Dataset description and VIR spectra characteristics

The Dawn mission at Vesta consisted of four phases, based on the altitude of the spacecraft: Survey (2735 km altitude), HAMO (High Altitude Mapping Orbit, 695 km), LAMO (Low Altitude Mapping Orbit, 210 km) and HAMO-2, an extension of the mission similar to the HAMO phase (Russell and Raymond, 2011). Here we used HAMO and HAMO-2 VIR data,

which cover large part of Vesta's surface with a high spatial resolution ( $\sim 180$ 224 m/pixel). The VIR instrument has two distinct detectors, or "channels": the 225 visible, covering wavelengths from 0.25  $\mu$ m to 1.07  $\mu$ m, and the infrared, with 226 sensitivity from 1.02  $\mu$ m to 5.10  $\mu$ m (De Sanctis et al., 2011). Each channel 227 has 432 spectral bands, so the average spectral sampling is 1.8 nm/band for 228 the visible channel and 9.8 nm/band for the infrared channel (De Sanctis 229 et al., 2011). Each VIR image cube has 256 spatial samples with a variable 230 number of lines determined by the length of a scan, and 432 spectral bands. 231 For details about the VIR data calibration, refer to a document archived 232 at the NASA Planetary Data System Small Bodies Node: http://sbn.psi. 233 edu/archive/dawn/vir/DWNVVIR\_I1B/DOCUMENT/VIR\_CALIBRATION/VIR\_CALIBRATION\_ 234 V2\_4.PDF. The VIR visible and infrared data are each separately geo-235 referenced, then the two VIS and IR subsets are merged together to form 236 a single spectrum covering the entire spectral range from 0.25 to 5.10  $\mu$ m. 237 The last 19 spectral channels of the visible range, which usually are affected 238 by higher instrumental noise, are discarded. 239 In this work described here, we consider VIR data calibrated in units of re-240

flectance factor (I/F) from 0.6  $\mu$ m to 2.5  $\mu$ m, focusing our analysis on the 241 two pyroxene bands. We considered the average spectra of BM units used for 242 spectral characterization by Zambon et al. (2014). The wavelength position 243 of the centers of the two bands gives information on the HED lithologies. 244 The displacement of the pyroxene band centers is sensitive to temperature 245 variations (e.g., Burbine et al. (2009), Reddy et al. (2012c)). However, given 246 the usual range of daytime temperatures recorded on Vesta for BM units (be-247 tween 252 and 265 K, Tosi et al. (2014)), this effect is essentially negligible 248

<sup>249</sup> in VIR data (Longobardo et al., 2014).

The strength (depth) of the pyroxene absorption bands is a function of the 250 abundances of the minerals, the particle size, and the presence of opaque 251 phases (e.g., Clark (1999)). Band depths are also affected by temperature 252 variations and the phase angle of the observations (Longobardo et al., 2014; 253 Reddy et al., 2012c). The relatively small range of temperatures found on 254 Vesta does not affect the band depth shape: Dalton et al. (2011) demon-255 strated that a temperature range of 150K is necessary to observe substantial 256 band-depth effects (whereas on Vesta we found variations never larger than 257 50 K). Band depths are the only spectral parameter among those considered 258 that vary depending on illumination and viewing geometry. However, the 259 variation of band depth with phase angle is very small for bright regions, 260 although variation can become important as albedo decreases(Longobardo 261 et al., 2014). In this work, observations at phase angles between 28° and 262  $52^{\circ}$  and incidence angles lower than  $60^{\circ}$  have been selected. According to 263 Longobardo et al. (2014), in this narrow range band depths are only weakly 264 dependent on phase and hence a photometric correction is not mandatory. 265 Phase, incidence and emission angles of the BM units are reported in Tables 266 7, 8 and 9 of the Zambon et al. (2014) supplementary online material (SOM). 267 In this work, we utilize the "Claudia" coordinate system (supplemental ma-268 terial of Russell et al., 2012; Li et al., 2012; Roatsch et al., 2012; Reddy et al., 269 2013), which has been used by Zambon et al. (2014) for tabulation of the 270 BM unit locations.

## 272 3. Analytical method

## 273 3.1. Linear unmixing

Understanding the composition of a surface using reflectance spectroscopy 274 may be challenging. Analysis of spectral parameters is useful but does not of-275 fer a complete characterization of the surface materials. Key additional infor-276 mation can be provided by the application of mixing model algorithms. Based 277 on past analysis (De Sanctis et al., 2012a; Jaumann et al., 2012; McCord 278 et al., 2012; Ammannito et al., 2013a; De Sanctis et al., 2013a), we assume 279 that reflectance spectra of Vesta can be reproduced by a linear combination 280 of at least three endmembers, chosen from a sample of eucrites, diogenites, 281 olivines and a featureless endmember. To describe the primary components 282 of Vesta's surface, we consider only diogenite and eucrite endmembers, and do 283 not include howardite as a separate endmember. Howardites, which are the 284 major component of Vesta terrains (De Sanctis et al., 2012a), are brecciated 285 rocks with variable amounts of both eucrite and diogenite; thus howardites 286 can be spectrally characterized by a mixing of the two components. The 287 mixing equation is: 288

$$Y_{mix} = a_1 y_1 + a_2 y_2 + a_3 y_3 \tag{1}$$

where  $y_1$ ,  $y_2$ ,  $y_3$  are the scaled spectra of the endmembers and  $a_1$ ,  $a_2$ ,  $a_3$  are their respective mixing coefficients. The unmixing algorithm requires all mixing coefficients to be non-negative, and their sum must be equal to unity. The method we developed is able to automatically find the endmembers and the mixing coefficients, subject to the positivity constraint, which allows exclusion of solutions that are not physically meaningful. We outline the principal steps of our algorithm as follows:

1. Spectral slope removal. Normalization of the spectrum by the contin-296 uum slope is done for two reasons. First, the position of the band center 297 is defined as the location of the reflectance minimum inside the band 298 as measured on a continuum-removed spectrum. Second, the process 299 of continuum normalization removes the influence of albedo from the 300 unmixing process. Albedo is closely tied to the photometric conditions 301 of the observation rather than mineralogy of the surface. To remove 302 the slope, we consider the line between the first point (0.6  $\mu$ m) and 303 the last point  $(2.5 \ \mu m)$  of the spectrum, then we divide the spectrum 304 by this line. To be consistent, the slope was removed in the same way 305 from both VIR spectra and the spectral endmembers (See Fig. 1). 306

307

308

FIGURE 1

309

2. We selected a sample of 14 laboratory spectra from a larger spectral library. In Section 3.2, we describe in detail the endmembers and the criteria for their selection.

313 3. We calculate the  $\chi^2$  values for all the combinations. We select the best fit spectrum corresponding with the minimum  $\chi^2$  value, with a confidence limit of 95%. In Table 2, we report all the  $\chi^2$  values of the bright units, associated to the best-fit, with the corresponding values of the correlation coefficient (r).

Following these steps, the algorithm is able to determines automatically the best combination of endmembers that describe the scene, along with their mixing coefficients. We accept only best fits with  $\chi^2 < 1$ .

## 321 3.2. Endmembers selection

Careful choice of endmembers is necessary to obtain consistent results. In our case, we consider as endmembers a number of laboratory spectra, accounting for pure minerals and for HED meteorites that are assumed to be representative of materials that may be present on Vesta's surface.

We select the endmembers from the Reflectance Experiment Laboratory 326 (RELAB) database (http://www.planetary.brown.edu/relabdocs). The 327 endmembers selection includes consideration of the particle size. The supple-328 mentary online material from Reddy et al. (2012b) discusses the grain size 329 differences between eucrite and diogenite. Studies based on the analysis of 330 HED laboratory samples show that the grain size of diogenites in howardite 331 breccias ranges from 500  $\mu{\rm m}$  to 1.5 millimeters whereas the average grain size 332 of eucritic material is  $\leq 70 \ \mu m$  (Beck et al., 2011; Reddy et al., 2012b SOM). 333 Previous work based on ground-based observation and remote sensing data 334 suggest that for the spatial resolution available, most of Vesta's surface can 335 be best represented by regolith with particle sizes  $< 25 \mu m$  (Hiroi et al., 1994; 336 Palomba et al., 2014; Zambon et al., 2014). This limits our choice to labo-337 ratory spectra acquired for samples in the  $< 25\mu$ m size fraction, if available. 338 Since we are working with continuum-removed spectra, absolute reflectance 339 does not enter in our analysis. Therefore, we cannot distinguish between a 340 low-reflectance featureless phase and a high-reflectance featureless phase: we 341 can just observe a decrease in band depth. To account for this effect, we 342 introduce a featureless endmember represented by a straight horizontal line. 343 Nonetheless, examination of the mean albedo values derived in Zambon et al. 344 (2014) for all the bright units can be useful in identifying the presence of low-345

346 vs. high-reflectance featureless endmembers.

We selected 14 endmember spectra including six eucrite, five diogenite 347 and two olivine samples from the RELAB database, plus the straight line 348 representing a featureless endmember. This limit is imposed in order to 349 allow computations to be completed in reasonable time. Since eucrite-rich 350 howardite is the dominant component on Vesta's surface (De Sanctis et al., 351 2013b) and because eucrites show a wide range of petrographic and spectral 352 characteristics, we include a larger number of them in our endmembers set. 353 The endmembers selection was done by considering the spectral parameters, 354 the spectral shape and their petrographic differences. We selected spectra 355 with band centers and band depths that cover the possible range of values 356 (Fig. 2), and those with different spectral characteristics, to take into ac-357 count all of the possible spectral variations (see Fig. 3). Moreover for eucrite 358 we considered both cumulate and basaltic samples, as well as polymict and 350 monomict material. We also tested the distribution frequency of each spec-360 trum from a preliminary combination of two endmembers chosen among the 361 whole available spectral library, performing a linear unmixing test. With just 362 two endmembers, the entire library of laboratory spectra can be used in the 363 analysis because of the reduced computational time. We calculate the spec-364 tral parameters with the same methods described in Section 4 of Zambon et 365 al. (2014). 366

In Fig. 4, we show the frequency of the endmembers found in the case of a linear combination of two endmembers. Eucrite appears to be the most abundant component (68% of the total components), followed by diogenite (25%) and olivine (7%). Taking the above discussion into consideration, we select for our analysis: 1) the eucrites Bereba (6e), Bouvante (7e), Cachari
(8e), Padvarninkai (20e), Serra de Mage (22e), Y74450 (24e); 2) the diogenites ALH77256 (2d), GRO95555 (5d), (6d) Johnstown, (8d) Tatahouine and
Y75032 (10d); and 3) two olivine samples, Jackson County (1o) and Green
olivine (2o) (see Table 1). In Fig. 3 we plot the selected endmembers.
FIGURE 2, 3, 4

377 TABLE1

378

## 379 4. Evaluation and limits of the linear method

Linear spectral unmixing can be a satisfactory approach for evaluating 380 the composition of a surface (e.g. Combe et al. (2008); Bioucas-Dias et al. 381 (2012)), even if a surface is a non-linear (intimate) mixture of several min-382 eralogical phases. The linear assumption implies an error in the estimation 383 of the abundances, but permits only a semi-quantitative estimation of the 384 different components. In this regard, we performed two types of tests: 1) 385 We selected different laboratory mixtures of possible Vesta analogues and 386 we applied linear unmixing to the spectra of such mixtures. This exercise 387 provides an idea of the uncertainty in the determination of the abundances. 388 2) We evaluate the stability of the solution for the VIR data by considering 389 the ten cases showing smaller values of  $\chi^2$ . If for these  $\chi^2$  values we find the 390 same endmembers and similar mixing coefficients, then the solution is stable 391 and the results are reliable. 392

## 393 4.1. Test on laboratory mixtures

To perform this test, we use spectra for sets of mixtures available in the RELAB database, and those used in Cloutis et al. (2013) and Serventi et al. (2013). In Table 1 of the supplementary online material (SOM), we list the spectral endmembers of the mixtures, indicating the origin, the particle size and the texture. We selected mixtures with particle sizes similar to those inferred for Vesta. In particular, we consider mixtures of two and three laboratory spectra:

401 1. Olivine and orthopyroxene mixtures;

402 2. Orthopyroxene and plagioclase mixtures;

403 3. Millbillillie eucrite and Murchison carbonaceous chondrite;

404 4. Plagioclase, low and high calcium pyroxenes;

5. Olivine, low and high calcium pyroxenes;

In Section 1 of the SOM and in Fig. 5, all the results obtained for the
laboratory mixtures are displayed.

We notice that the error in the estimation of the mixing coefficients de-408 pends on the number of endmembers in the mixture and on the endmember 409 itself. Generally, for mixtures of two pyroxenes with a third phase (e.g., 410 plagioclase or olivine), we found differences between the true values and 411 the modeled mixing coefficient up to 11% for the non-pyroxene endmember. 412 In particular, for the case in which the third component is plagioclase, we 413 414 have an underestimation of this phase of as much as 11% if its abundance is > 50%. In mixtures with plagooclase contents < 40%, the error is lower 415 (Table 2 and Fig. 1 of the SOM). If olivine is one of the endmembers, the 416

difference between the measured abundances and the modeled mixing coef-417 ficients is about +10%, when the olivine abundance is  $\sim 70\%$ . For lower 418 proportions of olivine in the mixture, the differences between measured and 419 modeled values are decreased (Table 3 and Fig. 2 of SOM and Fig. 5). 420 For mixtures of just two minerals, the results are similar for low calcium 421 pyroxenes and olivine. The estimated error increases with greater amounts 422 of olivine. In the case of an olivine content of 90%, we found the underes-423 timation of olivine to be 15% (Table 4 and Fig. 3 of the SOM). For lesser 424 proportions of olivine, the error is below 8%. For plagioclase mixed with 425 orthopyroxene at ratios ranging from 90:10 to 20:80, we find a consistent 426 underestimation of plagioclase from 12 to 19% (Table 5 and Fig. 4 of the 427 SOM). 428

Two-component mixtures of Murchison CC with Millbillillie eucrite have a 429 completely different trend. For Murchison abundances from 5% to 50%, the 430 mixing model consistently overestimated the CC component. The error in 431 CC abundance ranges from 11 to 26% despite the low  $\chi^2$  values (Table 6 and 432 Fig. 5 of the SOM). We consider good model results to be those for which 433 the difference between measured and modeled values is 5% (absolute) or less, 434 and we deem results to be acceptable when the difference is < 10%. Cases 435 for which the difference between measured and the modeled values is over 436 10% were rejected. Our tests on laboratory mixtures reveal that some mix-437 tures can be modeled successfully with the linear method, while some other 438 mixtures may not. Mixtures of different pyroxenes and mixture of pyroxenes 439 with oliving allow a good estimation of the mixing coefficients with the linear 440 methods with respect to the other types of mixtures. In the case of Vesta, we 441

can spectrally model its surface as a mixture of at most three endmembers:pyroxenes, olivine, and a featureless component.

As mentioned previously, our analysis does not consider the absolute reflectance, thus we cannot distinguish high- and low-reflectance featureless phases, but rather the presence of a generic featureless component that weakens and modifies the bands. In the case of plagioclase, we find that the error on plagioclase mixing coefficient decreases with increasing number of endmembers. Unfortunately laboratory data for mixtures of two pyroxenes with CC are not available.

451

452 FIGURE 5

453

## 454 4.2. Stability of the method

To test the stability of the method we consider the results of the ten 455 models showing the smaller  $\chi^2$  values. The method can be considered stable 456 if, for small  $\chi^2$  variations, the results do not vary substantially. If small  $\chi^2$ 457 variations lead to different endmembers, then the model results cannot be 458 considered to be reliable. In Fig. 6 we plot the best-fit spectrum for the ten 459 Aelia-SM cases with the lowest  $\chi^2$ . Variations in  $\chi^2$  of the order of 0.0001 460 led to variations of 1% in the abundances estimation, while the endmembers 461 remain unchanged, indicating robust results. We performed this test on all 462 the VIR spectra under consideration, and we obtained similar results. We 463 thus can conclude that our method yields stable solutions for the choice of 464 the endmembers and the estimation of the mixing coefficients. 465

466

## 467 FIGURE 6

468

# 469 5. Bright material units: Lithological variation as revealed by lin470 ear unmixing

<sup>471</sup> A first statistical analysis of the BM units on Vesta indicates that the <sup>472</sup> algorithm always chooses three endmembers, and two of these are generally <sup>473</sup> eucrite and diogenite, with comparable frequency ( $\sim 33\%$ ) (Fig. 7). The <sup>474</sup> third endmember is the featureless one with a frequency of 24.8%, while <sup>475</sup> olivine is present in few BM units with a frequency of 8.6% (Fig. 7).

Plots in Fig. 8 show a dependence between the modeled mixing coefficients 476 for eucrite and diogenite found here and the Zambon et al. (2014) band 477 centers for the BM units. We find that the content of eucrite increases with 478 increasing values of band centers, while the content of diogenite increases with 479 decreasing values of band centers. We observe a better correlation between 480 the band II center and eucrite and diogenite mixing coefficient, than for 481 the band I center. Generally, the band I center has a lower variability with 482 respect to the band II center (see De Sanctis et al. (2012a); Ammannito et al. 483 (2013a); Zambon et al. (2015)). The poorer correlation between the band I 484 center and the eucrite and diogenite mixing coefficients is mainly due to VIR 485 instrumental problems in the vicinity of band I, and on the bridging of the 486 visible and infrared channels. Thus we observe a better correlation between 487 mixing coefficients of eucrite and diogenite and the band II center than for 488 the band I center. 489

<sup>490</sup> In Fig. 9, we show the linear unmixing application for some representative

case of BM units. All the results have been reported in Table 2 and in section
2 of the SOM.

Aelia is a small crater (4.34 km) located at 14.26°S 140.6°E. It is char-493 acterized by mixed dark and bright material in its ejecta and by a high 494 diogenite content in its bright slope material (Palomba et al., 2014; Zambon 495 et al., 2014). Linear unmixing results indicate high mixing coefficient of dio-496 genite (67%) for the Aelia-SM and only 9% of eucrite as expected. Aelia-RM 497 which represents the ejecta showing a greater amount of eucrite than diogen-498 ite and a particularly large amount of the featureless endmember (33%). The 499 lower reflectance values of this BM unit suggest the presence of a dark phase 500 that is responsible for reducing the band depths (Fig. 9). Another diogenitic 501 unit is Licinia-SM (Zambon et al., 2014), with mixing coefficients of diogen-502 ite (53%) higher than eucrite (30%) (Fig. 9). The bright units with shorter 503 band centers are BU5-CWM and BU5-RM (Zambon et al., 2014), for these 504 BM units we find a diogenite content of 65% and 57% respectively (Fig. 7 of 505 the SOM) and BU8-CWM with a diogenite mixing coefficient of 65% (Fig. 8 506 of the SOM). Consistent with the findings of Zambon et al. (2014), the most 507 eucritic BM units are Eumachia-CMW with an eucrite mixing coefficient of 508 57% and a corresponding diagenite content of 14% of diagenite (Fig. 9), and 509 Tuccia-RM with 57% of eucrite and 27% of diogenite (Fig. 14 of the SOM). 510 Olivine on Vesta has been identified in few areas, most notably in the 511 region of Bellicia and Arruntia craters, but smaller olivine amounts have 512 been found also near the Albana and Pomponia craters, also located in the 513 northern hemisphere (Palomba et al., 2015). Local olivine enrichments have 514 been suggested by Ruesch et al. (2014) to be present also in the equatorial 515

region and in the southern hemisphere. Our analysis suggests that olivine is 516 present in several BM units, with amounts between 12 and 53%. We identify 517 sites with large olivine mixing coefficients (> 48%) in correspondence with 518 Arruntia, Bellicia in agreement with Ammannito et al. (2013b), and BU14 519 indicated as detection 3 reported by Ruesch et al. (2014). In these spectra we 520 observe a band I asymmetry typical of olivine and a reduction in the depth of 521 band II, indicating a relatively high amount of olivine, but not high enough 522 to completely obscure band II (Fig. 7). Among the possible olivine-bearing 523 sites found by Ruesch et al. (2014), we note that the Licinia-SM BM unit 524 has a mixing coefficient of olivine as high as 19%. Our algorithm chooses 525 olivine as a third endmember also for regions different from those observed 526 by Ammannito et al. (2013b) and Ruesch et al. (2014). Generally we find an 527 olivine mixing coefficient < 30% for these sites, except for Calpurnia-CWM 528 which has an olivine mixing coefficient of 35% (Fig. 10 of the SOM). 520

We cannot distinguish between opaque and neutral/featureless, high re-530 flectance components. In Fig. 10 we show the relationship between albedo 531 of the BM units (Schröder et al., 2013) and the modeled mixing coefficient of 532 the featureless endmember. In this plot, the albedo tends to decrease with 533 increasing mixing coefficient of the featureless component. We interpret this 534 trend as an indication that the algorithm's selection of the flat endmember 535 can be attributed to the presence of an opaque dark phase in most cases. 536 However, some exceptions exist. The bright streak called BU-15 SM is the 537 brightest unit on Vesta, with a higher content of diagenite (47%) than eucrite 538 (30%). This unit is different from the others because it is characterized by 539 a reduction of the band depth with respect to the surroundings even though 540

it has high albedo (Zambon et al., 2014). The model composition of BU-15
SM is eucrite and diogenite, with a featureless endmember mixing coefficient of 23% (Table 2). Thus, BU-15 SM may represent a unique case on
Vesta, where the presence of a high-albedo featureless component, such as
plagioclase, plays an important role in controlling the spectral properties.

546 FIGURE 7, 8, 9,10

#### 547

## 548 6. Discussion

The global compositional analyses hitherto performed on Vesta's surface with Dawn/VIR data were based on the spectral parameters of the kind used by De Sanctis et al. (2012a), and Ammannito et al. (2013b). Spectral parameters are very useful for determining the main lithology of a surface, but for a quantitative analysis mixing models are necessary.

In this work we focused our attention on eucrite and diogenite as the 554 dominant components, plus olivine or a featureless material. We analyzed 555 the Vesta spectra for BM units discussed by Zambon et al. (2014). In that 556 paper, the authors give the results of a compositional analysis based on spec-557 tral parameters. Unlike previous papers, the present work determines a semi-558 quantitative estimation of the mixing coefficients of each lithology. Here we 559 do not consider the spectral slope or the reflectance, and thus we cannot 560 establish the specific nature of this featureless endmember. However, by 561 examining the albedo of each BM unit, we can make inferences about the 562 identity of the featureless phase. 563

<sup>564</sup> We find that in general, the mixing coefficient of the featureless endmember

is anticorrelated with albedo, suggesting that this component is a low-albedo 565 opaque material. In fact, bright areas are often contaminated by dark ma-566 terial (Jaumann et al., 2012; Palomba et al., 2014). In Fig. 11, we show the 567 main results from the linear unmixing model plotted on the compositional 568 map from Ammannito et al. (2013b). The mixing coefficients of the different 569 components obtained by the model and the distribution of the BM units are 570 generally in good agreement with previous analyses (e.g., De Sanctis et al., 571 2012a, 2013b; Zambon et al., 2014). First, in all of the BM units, both eucrite 572 and diogenite spectra are used to fit VIR spectra, reflecting the distribution 573 of those components in the regolith (De Sanctis et al., 2013a,b). 574

From the results of our analysis, we can see that several BM units are dominated by eucrite and diogenite components (see Section 8 and Table 2). In
particular, we can classify these units as eucrite-rich units and diogenite-rich
units. A third type is olivine-rich BMs.

The distribution of eucrite-rich and diogenite-rich units is similar to the composition of surrounding terrains as inferred by the analysis of spectral parameters.

Olivine-rich BM units are associated with recognized olivine-bearing craters (Ammannito et al., 2013b; Ruesch et al., 2014). Moreover, where more units are seen in the same crater, they frequently belong to only one family. The only exceptions are a few diogenite-rich units present on eucritic terrains (Aelia, Canuleia, BU3, BU6), and few craters (Aelia, Canuleia, Oppia, Justina, BU9) where BMs show both compositions in the same location.

<sup>588</sup> Eucrite-rich BM units may have a mixing coefficient of eucrite as high as 44% <sup>589</sup> (red symbols in Fig.11). Those BM units are mostly located in the equatorial

region, between 90°E and 300°E longitude, and their distribution is compatible with eucrite-rich regions (Ammannito et al., 2013a). Furthermore, they can be subdivided in two populations: 1) one where the mixing coefficient of diogenite is higher than the mixing coefficient of the featureless endmember (diamond); 2) one where the mixing coefficient of the featureless endmember is greater than the mixing coefficient of diogenite (square). Those subgroups are equally distributed in number and by geograpy.

In eucrite-rich BM units, the olivine component was never used to fit the 597 spectra. This is consistent with the rare appearance of olivine, as secondary 598 mineral, in eucrite meteorites (Mittlefehldt et al., 1998). There is just one 599 exception, the crater wall material in BU2 (BU2 CWM in Table 2), where 600 the BM is fitted by the Serra de Mage eucrite spectrum (86%), with almost 601 negligible contribution from olivine and the featureless endmember. This is 602 also the only unit where both eucrite and diogenite spectra were not selected. 603 Note that the Serra de Mage eucrité shows spectral characteristics very close 604 to howardite spectra. The spectra of this eucrite is selected only in one other 605 case: the model for an anomalous diogenite rich-units (Canuleia SM) where 606 only HED spectra were used to model the data. 607

The diogenite-rich BMs are characterized by the association of the highest mixing coefficient of diogenite spectra (yellow symbols in Fig.11), with a lower threshold of 40%, except for the crater wall unit in BU9 (37%). The distribution of diogenite-rich BMs is widespread in longitude and in latitude, extending farther to the south than eucrite-rich units. The distribution of diogenite-rich BMs is also consistent with the results of previous works and the spectral parameter maps, and they are generally correlated with dio<sup>615</sup> genitic and howarditic terrains (Ammannito et al., 2013a).

Only two regions show diogenite-rich bright materials, within eucrite ter-616 rains, the Aelia unit and the units around 300°E, where are concentrated 617 BU3, BU6, and the Canuleia unit. The diogenite-rich BM units can be dif-618 ferentiated by the presence of the eucrite (diamonds in Fig. 11) or another 619 components (square) as the second spectra in the unmixing solutions. Unlike 620 eucrite-rich BM units, the sub-group with the spectra of eucrites selected as 621 the second component is the most frequent, whereas few units are character-622 ized by a non-HED component selected as the second endemember. 623

Moreover, in various cases ( $\sim 8.6\%$ ) olivine was used instead of the fea-624 tureless component (white points within diogenite-rich BMs), even if with 625 mixing coefficient lower than 30%, with the exception of the crater wall unit 626 in Calpurnia (35%). Aelia and Canuleia craters are both in eucritic terrains 627 and they show both eucrite-rich and diogenite-rich BM locations. Aelia has 628 diogenite-rich type 1 BMs on the wall, whereas the radial material (RM) 629 is eucrite-rich. This could indicate that the Aelia impact exposed diogenite 630 blocks from greater depths, whereas fresh eucritic material that originally 631 resided close to or at the surface was ejected to form the radial material. 632 In Canuleia, we have both type 1 and 2 BMs with eucrite and diogenite 633 compositions, but the differences between the two HED components are less 634 than 64%, indicating an homogeneous distribution of eucrite and diogenite 635 components, i.e., howarditic. 636

Only one Aelia BM unit, the RM, shows a clear eucrite-rich composition.
Thus, even in this location, diogenite material could have been exposed by the
impact and mixed with eucrite (i.e., formation of howarditic breccia blocks),

while some eucrite-rich fresh material was ejected from the crater, and deposited on the surrounding eucritic crust.

In contrast, BU3 and BU6 craters show both diogenite-rich units as RM (BU6 has also a SM, see Table 2). Those units are very close to the diogenite terrains, so perhaps the presence of diogenite-rich material can be attributed to contributions from southern craters. Alternately, it could be that the eucritic crust in that area is thinner, and thus fresh diogenitic material was more easily excavated and ejected from those craters.

Three other craters (Oppia, BU9 crater, and Justina) show the presence of 648 both eucrite- and diogenite-rich BMs. Oppia and BU9 show both units. In 649 the Oppia crater, two BM units are on the wall. The CWM are diogenite-650 enriched, whereas the corresponding slope material (SM) is characterized by a 651 higher amount of eucrite. In the second crater (BU9), the CWM is diogenite-652 rich, whereas a RM BM unit is fitted by a high mixing coefficient of eucrite. 653 BM units at crater Justina have been covered by VIR more than one time. 654 the four wall units (both CWM and SM) are eucrite-rich and the two RM 655 are diogenite-rich BM units. A third family is made of olivine-rich bright 656 units, i.e., where olivine spectra are selected as the most abundant compo-657 nent. In this family, all of the recognized BMs are enriched in olivine, with 658 diogenite > eucrite in BU14 and eucrite > diogenite in Arruntia and Bellicia 659 areas. These proportions are in agreement with those found by analysis of 660 spectral parameters in Zambon et al. (2014). These units are located in the 661 northern hemisphere and correspond to the olivine-bearing sites detected by Ammannito et al. (2013b) and Ruesch et al. (2014). For Bellicia and BU14, 663 we find an olivine mixing coefficient around of  $\sim 50\%$ , with slightly lower 664

values found in Arruntia. These values indicate that olivine is a dominant component in these sites (Ammannito et al., 2013b).

High abundances of olivine are generally expected to be present in man-667 tle and plutonic rocks, which could be present in topographically low areas. 668 In this work, we have employed Mg-rich olivine endmembers with very fine 669 particle sizes, compatible with the predicted Vesta dominant regolith size. 670 Therefore, the absolute abundance values could be lower than suggested by 671 our models, in particular considering that olivine is more favalitic (Fe-rich) 672 in HEDs. Olivine that is richer in iron has a stronger absorption band, and 673 hence a lower abundance of such Fe-rich olivine (relative to Mg-rich olivine) 674 would be needed to model a given spectrum. 675

Poulet et al. (2015) discussed the retrieval of composition from VIR spec-676 tra by using a non-linear method (Shkuratov et al., 1999). It should be noted 677 that the results from a non-linear technique, although in principle better than 678 those from a linear model, can suffer if the starting data has not been prop-679 erly treated. For the VIR dataset, extreme care must be taken, for example, 680 in bridging the sections VIS and IR, exclusion of spectral channels suffering 681 from recurring artifacts, and removal of the continuum. Poulet et al. (2015) 682 concluded that coarse-grained olivine is likely to be present in all major units 683 of Vesta. In general, they also found that the abundance of olivine could be 684 lower because of the presence of iron-rich olivine or because of an increase 685 in the grain size of olivine with respect to the other phases. An alterna-686 tive hypothesis that could explain higher surface abundances of olivine in 687 such northern locations is an exogenic origin - e.g., material delivered by 688 an olivine-rich impactor, as suggested by Le Corre et al. (2015). Moreover, 689

we find that in those terrains the third selected endmember is frequently afeatureless component.

As discussed above, the mixing coefficient of the featureless endmember is 692 anti-correlated with the albedo, suggesting that this component is associated 693 with low-albedo CC-like opaque material. In fact, bright areas are often con-694 taminated by dark material (Jaumann et al., 2012; Palomba et al., 2014). 695 We do find one exception where a BM unit with a large mixing coefficient 696 for the featureless endmember, BU15-SM, has the highest albedo (Zambon 697 et al., 2014). This unit is matched by a diogenite-rich bright unit extending 698 into the Rheasilvia basin, and Zambon et al. (2014) predicted the possible 699 presence of a bright component. 700

The major bright, iron-poor, mineral in HEDs and in basic igneous compo-701 sition is plagioclase, which can have a large abundance in magmatic rocks, 702 both effusive and cumulitic, intrusive rocks. Plagioclase can exhibit a char-703 acteristic absorption at  $1.25\mu m$  caused by impurity iron at abundances cor-704 Burns, 1993; Cheek et al., 2011; Serventi responding to FeO < 1%(e.g.705 et al., 2013), However, the  $1.25\mu$ m feature cannot be recognized in mixtures 706 with mafic iron-bearing silicates unless the plagioclase is present in very large 707 amounts. If plagioclase is indeed revealed by our analysis, it should be noted 708 that in HEDs it is generally present in eucrites and cumulitic eucrites, but 709 not in diogenites. 710

Another important aspect showed by our results is the selection of olivine spectra with low mixing coefficient for the diogenite-rich BM units, in place of the featureless endmember. From a spectroscopic viewpoint, this implies that reductions of the intensities of band I and band II are not correlated,

otherwise the model should have selected the featureless spectrum. In ad-715 dition, a weak asymmetry around the  $1-\mu m$  absorption is revealed, so the 716 selection of only eucrite and diogenite spectra alone is not able to properly 717 fit VIR data in that specific spectral region. The presence of this low amount 718 of olivine in the diogenite-rich BM units indicates that iron could be present 719 in one or more phases with a band around 1  $\mu$ m but with a very weak, or 720 absent, absorption around 2  $\mu$ m. In general, these spectral characteristics 721 are consistent with the presence of olivine, high-Ca pyroxene, plagioclase or 722 a glassy component, in basic igneous compositions. 723

Iron-bearing silicate glasses (e.g., volcanic or impact glasses) have a dom-724 inant crystal-field absorption at ~  $1.1\mu m$ , the intensity of which depends 725 on the iron abundance. Thus a glassy component cannot be excluded in 726 units that are formed by impacts. In fact, impact melts have been identified 727 in some HEDs (see e.g., Metzler and Stoffler, 1995; Singerling et al., 2013). 728 High-Ca pyroxenes are not abundant in diogenite s, even if there are samples 729 where they are found in abundance higher than 10% (e.g., Mittlefehldt et al., 730 1998). High-Ca pyroxene can also be frequently found as exolution lamellae 731 in orthopyroxene crystals, influencing the spectral characteristics of the host 732 low-Ca hosting pyroxene (Gaffey et al., 2002). 733

Olivine is difficult to detect, especially in small amounts. The shape and the position of the  $1\mu m$  band are influenced by the presence of olivine if it is present in high amounts in mixtures with pyroxene (e.g. in particular the orthopyroxene-olivine mixtures studied by Cloutis et al., 1986; Beck et al., 2013).

<sup>739</sup> Beck et al. (2013) argued that olivine could be recognized only at contents

> 30% in HEDs, because under this threshold, the olivine is masked in 740 mixtures with orthopyroxene. (Beck et al., 2013) predicted that olivine in 741 diogenite and harzburgite terrains could not be detected when present in 742 amounts below this threshold. Recent studies by Horgan et al. (2014) and 743 Shestopalov et al. (2015) suggest two different methods for detection of small 744 amounts of olivine in mixtures with pyroxene. Horgan et al. (2014) used dif-745 ferent spectral indices to identify the presence of olivine in the spectra of 746 Mars. Shestopalov et al. (2015) observed that the uncertainty (variance) of 747 the band center position in spectra of laboratory olivine-pyroxene mixtures 748 depends on the shift of the band center at 950 nm. They found that mixtures 749 with different olivine contents can be identified by their position in a plot of 750 the variance vs. band center at 950 nm. 751

On the other hand, Poulet et al. (2015) concluded that best fits to all the studied Vesta terrains are achieved when a component of forsteritic olivine (Fo70) is included. They also emphasized that the particle size has an effect on the olivine abundance obtained by a non-linear unmixing method. Olivine-rich melts were recently found in some howardites (Beck et al., 2013), so we cannot rule out that those BM units could have low olivine mixing coefficient.

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## 760 7. Conclusions

FIGURE 11

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▶ 1. To model VIR data for Vesta, we applied a linear spectral unmixing model. Our analysis does not take into account the spectral slope and the albedo but only the spectral shape.

764	2.	Tests on laboratory mixtures of olivine and low- and high-calcium py-
765		roxene indicate that olivine mixing coefficients are underestimated with
766		an accuracy within $10\%$ for olivine amounts > 50\%. For mixtures
767		of plagioclase with low-and high-calcium pyroxenes, the plagioclase
768		amount is underestimated within $11\%$ for plagioclase contents $> 40\%$ .
769		Analysis of mixtures of Millbillillie eucrite with carbonaceous chondrite
770		(CC) material indicates that the CC abundance was overestimated by
771		up to 26%.
772	3.	Linear unmixing applied to Vesta spectra is able to detect the principal
773		HED and non-HED lithologies present on the asteroid.
774	4.	We can divide the bright material units into three main groups: eucrite-
775		rich, diogenite-rich, and olivine-rich, depending on the most abundant
776		endmember used to fit the spectrum. The non-HED endmember can
777		be either olivine or a featureless component. This confirms that Vesta
778		is composed of a high content of pyroxenes mixed with a lower concen-
779		tration of other mineralogical phases.
780	5.	The third endmember is often the featureless phase, which is associated
781		with a reduction in the strength of both pyroxene bands. The mixing
782		coefficient of the featureless endmember is anticorrelated with albedo,
783		suggesting that the featureless phase can be associated with low-albedo,
784	~	CC-like opaque material.
785	6.	Large amounts of olivine are detected in the Bellicia, Arruntia and

Large amounts of onvine are detected in the Bellicia, Arruntia and BU14 BM units. Where lower olivine contents (< 30%) are found, the sites are generally associated with high concentrations of diogenite.

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## 788 Figure and Table captions

Figure 1: Example of VIR spectrum before (top) and after (bottom) the
removal of the slope.

Figure 2: Spectral parameter plots for the diogenite and eucrite laboratory samples. The gray circles indicate the samples selected for use as
endmembers in this study. The numbers refer to the endmember names in
Table 1.

Figure 3: Spectra of the endmembers selected. The text labels refer
to the sample names in Table 1 (6e Bereba, 7e Bouvante, 8e Cachari, 20e
Padvarninkai, 22e Serra de Mage, 24e Y74450, 2d ALH77256, 5d GRO95555,
6d Johnstown, 8d Tatahouine, 10d Y75032, 1o Jackson cty, 2o Green olivine).
Figure 4: Frequency of the endmembers for the linear unmixing with
two endmembers.

Figure 5: Linear unmixing applied to a mixture of olivine with low and high calcium pyroxene. The band centers of the modeled spectra are shifted slightly with respect to those of the laboratory spectra. In cases where the  $\chi^2$  values differ from 0, there is an imperfect match between the measured and modeled spectra.

Figure 6: The results of linear unmixing applied to the Aelia BM unit for the 10 cases with the lowest  $\chi^2$ . These results indicate that small variations in  $\chi^2$  do not correlate with changes in the endmembers or with large variations in the mixing coefficients of the endmembers, attesting to the stability of the results. Black lines are the measured spectra, the gray lines are the model mixture spectra.

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**Figure 7:** Frequency of the endmembers selected for the linear unmixing

<sup>813</sup> with three endmembers.

Figure 8: Plots of band centers vs. eucrite mixing coefficients show that eucrite content increases for increasing band center values. Right: Similar plots for band centers and diogenite mixing coefficients indicates that diogenite content decreases with increasing band center values. Band center values for the BM units are those calculated in Zambon et al. (2014). The coefficient of determination (i.e., the squared correlation coefficient  $R^2$ ) is also provided.

Figure 9: Examples of linear spectral unmixing applied to BM units. Black lines are the measured spectra, the gray lines are the model mixture spectra. Table 1 summarizes the model endmember mixing coefficients for all the BM units. Related plots are presented in the SOM.

Figure 10: The albedo of each BM location plotted against the modeled mixing coefficient of the featureless endmember. As albedo decreases, the mixing coefficient of the featureless endmember increases.

Figure 11: Results obtained for the linear unmixing applied to BM 828 units overlain on a mineralogical map of Vesta. Red color indicates eucrite-829 rich units, yellow color designates the diogenite-rich units, and the white 830 symbols are the olivine-rich bright materials. Diamonds indicate BM units 831 for which the eucrite or diogenite mixing coefficient is larger than that of 832 the third endmember. Squares correspond to BM in which the eucrite or 833 diogenite amount is lower than that of the third component. The solid white 834 points are BM units with olivine as the third endmember (See also Table 835 2). The legend at the top refers to the different cases found in our analysis. 836 For example, "D>III" indicates that the mixing coefficient of the diogenite 837

is greater than those of the third component. "E>III" represents the cases
in which the mixing coefficient of the eucrite is greater than that of the third
endmember, and so on

Table 1: Sample name, type, band centers, and band depths for all the endmembers in our spectral library. Each spectrum is assigned an identification number that contains "e" for eucrite, "d" for diogenite, and "o" for olivine.

Table 2: Summary of the results obtained from application of linear 845 spectral unmixing on BM units on Vesta. Endmembers are designated as 846 eucrite ("E"), diogenite ("D"), or olivine ("OI"), while "Line" refer to the 847 featureless component. Different text color and box color are indicative of 848 variations in composition: Red text refers to eucrite-rich BM units where 849 the featureless endmember (line) is the second most abundant. Yellow text 850 indicates diogenite-rich endmember in which the second most abundant end-851 member is either the featureless one or olivine. Green text represents cases 852 with olivine as the non-HED spectrum. Red boxes correspond to eucrite-rich 853 BM units with diagenite as the second most abundant endmember. Yellow 854 boxes are diogenite-rich BM units with eucrite as the second most abundant 855 endmember. Green boxes represent the olivine-rich BM units. In the last 856 column are reported the values of the correlation coefficient r. 857

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Figure 1



Figure 2







Figure 5



Figure 6







Figure 9





	Endmembers							
#	name	type	BCI	BCII	BDI	BDII		
1e	A-87272	Eucrite-mmict	0.941	2.002	0.552	0.350		
2e	A-881819	Eucrite	0.931	1.966	0.481	0.299		
3e	ALH78132	Eucrite-pmict	0.932	1.960	0.426	0.275		
4e	ALH85001	Eucrite-Mg rich	0.927	1.946	0.432	0.266		
5e	ALHA76005	Eucrite-pmict	0.935	1.980	0.385	0.242		
<u>6e</u>	Bereba	Eucrite-mmict	0.941	2.019	0.419	0.264		
<u>7e</u>	Bouvante	Eucrite-mmict	0.945	2.009	0.451	0.333		
<u>8e</u>	Cachari	Eucrite-mmict	0.940	2.000	0.543	0.363		
9e	EET 87542	Eucrite-br	0.941	2.000	0.254	0.131		
10e	EET 90020	Eucrite-unbr	0.941	2.006	0.517	0.276		
11e	EETA79005	Eucrite-pmict	0.934	1.970	0.466	0.306		
12e	GRO 95533	Eucrite-br	0.941	2.014	0.486	0.330		
13e	Ibitira	Eucrite-mmict	0.940	1.994	0.602	0.348		
14e	Jonzac	Eucrite-mmict	0.938	2.002	0.542	0.379		
15e	Juvinas	Eucrite-mmict	0.936	1.994	0.469	0.298		
16e	LEW 87004	Eucrite-pmict	0.935	1.976	0.430	0.261		
17e	Millbillillie	Eucrite-mmict	0.938	2.006	0.454	0.273		
18e	Moore County	Eucrite-cm	0.938	1.982	0.598	0.378		
19e	PCA 82502	Eucrite-unbr	0.941	2.013	0.470	0.338		
<u>20e</u>	Padvarninkai	Eucrite-mmict	0.949	2.039	0.232	0.108		
21e	Pasamonte	Eucrite-pmict	0.940	2.003	0.494	0.319		
<u>22e</u>	SerradeMage	Eucrite-cm	0.931	1.963	0.442	0.286		
23e	Stannern	Eucrite-mmict	0.938	2.008	0.405	0.264		
<u>24e</u>	<u>Y74450</u>	Eucrite-pmict	0.935	1.985	0.382	0.208		
25e	Y792510	Eucrite-mmict	0.942	2.011	0.511	0.324		
26e	Y792769	Eucrite-pmict	0.940	2.012	0.408	0.236		
1d	A881526	Diogenite	0.919	1.889	0.646	0.428		
<u>2d</u>	<u>ALH77256</u>	Diogenite	0.921	1.881	0.588	0.411		
3d	Aioun el Atrouss	Diogenite-pm	0.923	1.902	0.631	0.428		
4d	EETA79002	Diogenite	0.919	1.877	0.508	0.300		
<u>5d</u>	<u>GRO95555</u>	Diogenite-an	0.922	1.908	0.625	0.428		
<u>6d</u>	<u>Johnstown</u>	Diogenite	0.917	1.877	0.447	0.247		
7d	LAP 91900	Diogenite	0.921	1.903	0.623	0.411		
<u>8d</u>	Tatahouine	Diogenite	0.919	1.894	0.633	0.421		
9d	Y74013	Diogenite	0.923	1.914	0.422	0.254		
<u>10d</u>	<u>Y75032</u>	Diogenite	0.927	1.940	0.517	0.311		
<u>10</u>	Jackson cty	Olivine	1.056	-	0.361	-		
<u>20</u>	<u>Green olivine</u>	Olivine	1.054	-	0.226	-		
		Table 1						

Table 1

			Linear unmixin	g results applied to	o the BM unit	5			
BM unit	Cube	Endm 1	Abundance	Endm 2	Abundance	Endm 3	Abundance	$\chi^2$	
BU1 RM	(372521396)	Cachari E	0.28	GRO95555 D	0.52	Line	0.20	0.201	
BU2 CWM	(372741836)	Cachari E	0.46	GRO95555 D	0.28	Line	0.26	0.447	
BU2 SM	(372741836)	Cachari E	0.47	Tatahouine D	0.25	Line	0.28	0.342	
BU2 CWM	(395611823)	SerradeMage E	0.86	Jackson cty Ol	0.12	Line	0.02	0.204	
BU2 SM	(395611823)	Cachari E	0.47	Tatahouine D	0.23	Line	0.30	0.247	
BU3 RM	(372385964)	Cachari E	0.32	GRO95555 D	0.46	Line	0.22	0.342	
BU4 CWM	(372212906)	Cachari E	0.41	GRO95555 D	0.45	Line	0.14	0.472	
BU5 CWM	(371993041)	Bouvante E	0.28	GRO95555 D	0.65	Line	0.07	0.705	
	(371993041)	Cachari E	0.20	GRO95555 D	0.57	Line	0.23	0.306	
BU5 SM	(371993041)	Cachari E	0.33	GRO95555 D	0.44	Line	0.23	0.386	
BU6 RM	(372078317)	Cachari E	0.34	GRO95555 D	0.48	Line	0.18	0.312	
BU6 SM	(372078317)	Cachari E	0.44	GRO95555 D	0.44	Line	0.12	0.302	
BU7 RM	(394993455)	Cachari E	0.52	GRO95555 D	0.33	Line	0.15	0.281	
BU8 CWM	(394155284)	Bouvante E	0.24	GRO95555 D	0.65	Line	0.11	0.351	
BU8 SM	(394155284)	Bouvante E	0.23	GRO95555 D	0.61	Line	0.16	0.278	
BU9 CWM	(372430301)	Bouvante E	0.34	Tatahouine D	0.37	Jackson cty Ol	0.29	0.242	
BU9 RM	(372430301)	Cachari E	0.48	Tatahouine D	0.19	Line	0.33	0.229	
BU10 CWM	(371992493)	Cachari E	0.45	GRO95555 D	0.28	Line	0.27	0.296	
BU10 SM	(371992493)	Cachari E	0.51	Tatahouine D	0.22	Line	0.27	0.271	
BU10 CWM	(372609208)	Cachari E	0.44	GRO95555 D	0.28	Line	0.28	0.362	
BU10 SM	(372609208)	Cachari E	0.49	Tatahouine D	0.24	Line	0.27	0.233	
BU11 CWM	(371815732)	Cachari E	0.48	Tatahouine D	0.20	Line	0.32	0.214	
BU11 SM	(372874554)	Cachari E	0.52	Tatahouine D	0.18	Line	0.30	0.280	
BU11 SM	(393801089)	Cachari E	0.53	Tatahouine D	0.18	Line	0.29	0.341	
	(371632826)	Bouvante E	0.25	GRO95555 D	0.50	Jackson cty Ol	0.25	0.530	
	(371632826)	Bouvante E	0.23	GRO95555 D	0.53	Jackson cty Ol	0.24	0.501	
BU13 SM	(371986649)	Cachari E	0.46	GRO95555 D	0.36	Line	0.18	0.333	
BU14 CWM	(393441792)	Cachari E	0.25	Tatahouine D	0.27	Jackson cty Ol	0.48	0.208	
BU14 SM	(393441792)	Cachari E	0.17	Tatahouine D	0.30	Jackson cty Ol	0.53	0.167	
BU15 CWM	(385962327)	Cachari E	0.32	Tatahouine D	0.56	Line	0.12	0.421	
BU15 SM	(385962327)	Cachari E	0.30	Tatahouine D	0.47	Line	0.23	0.212	
	(								

	<u></u>					E L C	A1 1	2	
BM unit	Cube	Endm 1	Abundance	Endm 2	Abundance	Endm 3	Abundance	χ-	r
Arruntia CWM	(393750738)	Cachari ${\rm E}$	0.29	Tatahouine D	0.23	Jackson cty Ol	0.48	0.269	0.984
Arruntia RM	(393750391)	Bereba E	0.36	Tatahouine D	0.26	Jackson cty Ol	0.38	0.194	0.966
Arruntia SM	(393750738)	Cachari E	0.31	GRO95555 D	0.25	Green Ol	0.44	0.264	0.988
Aelia CWM	(394860977)	Cachari E	0.34	GRO95555 D	0.42	Line	0.24	0.214	0.992
Aelia RM	(394860977)	Cachari E	0.50	Tatahouine D	0.17	Line	0.33	0.260	0.987
Aelia SM	(394860977)	Cachari E	0.09	GRO95555 D	0.67	Line	0.24	0.215	0.993
Bellicia CWM	(393308781)	Cachari E	0.35	Tatahouine D	0.16	Green Ol	0.49	0.490	0.981
Calpurnia CWM	(395166561)	Bouvante E	0.26	Tatahouine D	0.39	Jackson cty Ol	0.35	0.270	0.987
Canuleia CWM	(394198821)	Cachari E	0.45	Tatahouine D	0.39	Line	0.16	0.459	0.983
Canuleia RM	(394198821)	Cachari E	0.44	GRO95555 D	0.46	Line	0.10	0.539	0.987
Canuleia SM	(394198821)	Bouvante E	0.28	Serra de Mage E	0.25	GRO95555 D	0.47	0.632	0.984
Canuleia RM	(394198127)	Cachari E	0.35	GRO95555 D	0.41	Line	0.24	0.353	0.988
Canuleia RM	(371905283)	Cachari E	0.60	GRO95555 D	0.31	Line	0.09	0.577	0.985
Cornelia CWM	(371813008)	Bouvante E	0.28	GRO95555 D	0.58	Jackson cty Ol	0.14	0.381	0.989
Cornelia CWM	(394683501)	Cachari E	0.36	GRO95555 D	0.47	Line	0.17	0.157	0.995
Cornelia SM	(394683501)	Cachari E	0.35	GRO95555 D	0.45	Line	0.20	0.145	0.995
Cornelia CWM	(373313780)	Bouvante E	0.31	GRO95555 D	0.57	Jackson cty Ol	0.12	0.485	0.986
Cornelia SM	(373313780)	Bouvante E	0.25	GRO95555 D	0.59	Jackson cty Ol	0.16	0.315	0.991
Drusilla CWM	(393931377)	Bouvante E	0.28	GRO95555 D	0.55	Jackson cty Ol	0.17	0.420	0.987
Drusilla SM	(393931377)	Bouvante E	0.37	Tatahouine D	0.46	Jackson cty Ol	0.17	0.368	0.987
Eumachia CWM	(394859589)	Cachari E	0.57	Tatahouine D	0.14	Line	0.29	0.342	0.985
Eumachia SM	(394859589)	Cachari E	0.54	Tatahouine D	0.17	Line	0.29	0.328	0.985
Justina CWM	(372520864)	Cachari E	0.54	Tatahouine D	0.28	Line	0.18	0.296	0.991
Justina RM	(372520864)	Cachari E	0.38	GRO95555 D	0.43	Line	0.19	0.264	0.989
Justina SM	(372520864)	Cachari E	0.42	GRO95555 D	0.35	Line	0.23	0.305	0.989
Justina CWM	(371904187)	Cachari E	0.58	Tatahouine D	0.27	Line	0.15	0.370	0.983
Justina RM	(371904187)	Cachari E	0.41	GRO95555 D	0.44	Line	0.15	0.323	0.992
Justina SM	(371904187)	Cachari E	0.54	Tatahouine D	0.29	Line	0.17	0.335	0.989
Lepida SM	(395256929)	Cachari E	0.45	ALHA77256 D	0.30	Line	0.25	0.595	0.977
Licinia CWM	(395387692)	Bouvante E	0.18	GRO95555 D	0.63	Jackson cty Ol	0.19	0.273	0.992
Licinia SM	(395387692)	Cachari E	0.30	GRO95555 D	0.53	Line	0.17	0.192	0.994
Marcia CWM	(395301166)	Cachari E	0.53	GRO95555 D	0.23	Line	0.24	0.410	0.984
Marcia SM	(395301166)	Cachari E	0.46	GRO95555 D	0.31	Line	0.23	0.396	0.986
Marcia CWM	(395301549)	Cachari E	0.48	GRO95555 D	0.32	Line	0.20	0.301	0.990
Marcia SM	(395301549)	Cachari E	0.48	GRO95555 D	0.30	Line	0.22	0.353	0.987
Myia RM	(372300571)	Cachari E	0.38	GRO95555 D	0.41	Line	0.21	0.396	0.987
Oppia CWM	(373136361)	Bouvante E	0.38	Tatahouine D	0.43	Jackson cty Ol	0.19	0.364	0.986
Oppia SM	(373136361)	Cachari E	0.45	GRO95555 D	0.27	Line	0.28	0.309	0.979
Pinaria CWM	(371727131)	Cachari E	0.41	ALHA77256 D	0.46	Line	0.13	0.332	0.990
Pinaria SM	(371727131)	Bouvante E	0.27	GRO95555 D	0.54	Jackson cty Ol	0.19	0.308	0.990
Sextilia CWM	(371816828)	Cachari E	0.48	GRO95555 D	0.35	Line	0.17	0.461	0.986
Sextilia SM	(371816828)	Cachari E	0.48	GRO95555 D	0.36	Line	0.16	0.484	0.986
Tuccia RM	(372565807)	Cachari E	0.59	Tatahouine D	0.27	Line	0.14	0.319	0.990

Table 2