Global Estimates of Lunar Surface Elemental Abundances Derived from LRO Diviner Data

Ming Ma¹, Shengbo chen², Tianqi Lu², Peng Lu², Lu Yu³, and JIn Qin⁴

¹Jilin Jianzhu Univerdity ²Jilin University ³Jiangsu Center for Collaborative Innovation ⁴Macau University of Science and Techniques

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Abstract

Information on the elemental abundances and distribution is essential for understanding the petrological characteristics and geological evolution of the Moon. In this paper, the thermal infrared data acquired by Lunar Reconnaissance Orbiter (LRO) Diviner are processed to investigate lunar elemental abundances on a global scale (60°N/S) for the first time. The Diviner Level 3 Standard Christiansen feature (CF) product with the resolution of 128 pixels/degree and the coverage of 99.86% is first analyzed and used. The Diviner global models are then established by the univariate regression methods based on the relationships between Diviner CF and ground truths of elemental abundances at 48 lunar sampling sites and the limitations of 1 RMSE of 48 datasets. Finally, the best maps of SiO, TiO, AlO, FeO, MgO and CaO abundances considering both resolution and coverage simultaneously are presented and analyzed from global, geologic units, crater and ejecta surfaces. The comparisons indicate that a satisfactory consistency is observed between Diviner results and Clementine or Chang'E (CE)-1 results, while Diviner results exhibit better practicability in presenting detailed information for elemental abundances on lunar surfaces and higher accuracy on the surface with high latitudes or poor light conditions. Meanwhile, it is also demonstrated that Diviner results is the reliable data sources for the applications in classifications of mare basalt, inhomogeneity of highland crust and Mg#.

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Ming Ma^{1, 2}, Shengbo Chen², Tianqi Lu², Peng Lu², Yu Lu^{3, 4}, Qin Jin⁵

¹School of Surveying and Exploration Engineering, Jilin Jianzhu University, Changchun, China.

- 8 ³ School of Geographic and Oceanographic Sciences, Nanjing University, Nanjing, China.
- 9 ⁴Jiangsu Center for Collaborative Innovation in Geographical Information Resource Development 10 and Application, Nanjing, China.
- ⁵Space Science Institute, Macau University of Science and Techniques, Macau, China.
- 13 **Correspondence to:** Shengbo Chen, E-mail: <u>chensb@jlu.edu.cn</u>
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15 Key Points:

• Establishment of global models by univariate regression methods based on the relationships between CF and ground truths

- Development of SiO₂, TiO₂, Al₂O₃, FeO, MgO and CaO abundance maps with resolution of 128
 pixels/degree and coverage of 99.86%
- Comparison of the results with Lunar Prospector, Clementine and Chang'E-1 results and application of the results in geological problems
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23 Plain Language Summary

24 In LRO project, Diviner radiometer now in orbit had been acquired more than 480 billion 25 calibrated radiance values from July 5, 2009 to December 15, 2019. The Standard CF product with 26 the resolution of 128 pixels/degree (237 m/pixel at the equator) and the coverage of 99.86% had been 27 provided by Diviner science team. Considering both resolution and coverage simultaneously, the 28 Standard CF product is the best data source for elemental abundance estimation Therefore, new SiO₂, 29 TiO₂, Al₂O₃, FeO, MgO, CaO maps derived from LRO Diviner data are calculated and presented in 30 this paper. In general, Diviner elemental abundance results are consistent with previous studies such 31 as Clementine and Chang'E-1 results, while Diviner results exhibit better practicability in presenting 32 detailed information for elemental abundances on lunar surfaces and higher accuracy especially on 33 the surface with high latitudes or poor light conditions. Meanwhile, this provides a feasible data 34 source to study the classifications of mare basalt, the inhomogeneity of highland crust and the lunar 35 Mg#.

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^{7 &}lt;sup>2</sup>School of Geo-exploration Science and Techniques, Jilin University, Changchun, China.

41 Abstract

42 Information on the elemental abundances and distribution is essential for understanding the 43 petrological characteristics and geological evolution of the Moon. In this paper, the thermal infrared 44 data acquired by Lunar Reconnaissance Orbiter (LRO) Diviner are processed to investigate lunar 45 elemental abundances on a global scale (60°N/S) for the first time. The Diviner Level 3 Standard Christiansen feature (CF) product with the resolution of 128 pixels/degree and the coverage of 46 47 99.86% is first analyzed and used. The Diviner global models are then established by the univariate 48 regression methods based on the relationships between Diviner CF and ground truths of elemental 49 abundances at 48 lunar sampling sites and the limitations of 1 RMSE of 48 datasets. Finally, the best 50 maps of SiO₂, TiO₂, Al₂O₃, FeO, MgO and CaO abundances considering both resolution and 51 coverage simultaneously are presented and analyzed from global, geologic units, crater and ejecta 52 surfaces. The comparisons indicate that a satisfactory consistency is observed between Diviner 53 results and Clementine or Chang'E (CE)-1 results, while Diviner results exhibit better practicability 54 in presenting detailed information for elemental abundances on lunar surfaces and higher accuracy on 55 the surface with high latitudes or poor light conditions. Meanwhile, it is also demonstrated that Diviner results is the reliable data sources for the applications in classifications of mare basalt, 56 57 inhomogeneity of highland crust and Mg#.

58 1. Introduction

59 In the long evolutionary history, the development and reconstruction of lunar surface layer are 60 dominated by space weathering and impact processes after internal dynamic geological evolution 61 processes such as magmatism, tectonization, seismicity and metamorphism. The surface records the 62 entire history of the interaction with the outside environment. By understanding this history, the 63 evolutionary processes of airless bodies will be better revealed which can be applied to the study of 64 other similar stars within the Solar System. Therefore, as the basic part of lunar surface exploration, 65 the study on elemental abundances and distribution is particularly important. Many remote sensing 66 projects have made this issue their main objective and mission.

67 Previous groups had estimated elemental abundances based on Earth-based reflectance data (e.g., 68 Charette et al., 1974, 1977); X ray spectroscopy data acquired by Apollo 15 and Apollo 16 (Adler et 69 al., 1972, 1973), SMART-1 (Grande et al., 2004), Chandrayaan-1 (Swinyard et al., 2009; Athiray et 70 al., 2013) or CE-2 (Dong et al., 2016); gamma ray spectroscopy data acquired by Apollo 15 and Apollo 16 (Metzger et al., 1974, 1977), Lunar Prospector (LP) (Lawrence et al., 1998, 1999, 2000, 71 72 2002; Prettyman et al., 2006) or SELENE (Kaguya) (Naito et al., 2018); neutron spectroscopy data 73 acquired by LP (Elphic et al., 1998, 2000, 2002; Feldman et al., 1998, 2000); reflectance spectral data 74 acquired by Clementine (Lucey et al., 1995, 1998, 2000, 2006; Sato et al., 2017) or CE-1 (Ling et al., 75 2011; Yan et al., 2012; Wu, 2012; Sun et al., 2016; Xia et al., 2019). For high energy techniques 76 (such as gamma ray), the relationships between flux of ray energy spectrum and change of elemental 77 energy are the most important theoretical basis. Not only absolute elemental abundances from an 78 average depth of 30 cm (Lawrence et al., 2000; Prettyman et al., 2006), but also some key elemental 79 abundances (such as Th, K, U and O) that can't be detected by any other techniques can be obtained

(Metzger et al., 1977; Lawrence et al., 1999, 2000). For optical reflective techniques with a longer
history (Wood, 1912), electronic transition spectral characteristics provide the most direct supports.
The relationships between spectral characteristics and elemental abundances had been well studied
(Burns, 1970; Hapke et al., 1970; McCord and Adams, 1973; McCord et al., 1981), and the elemental
abundance maps had higher resolution and been better developed.

85 However, there are four obvious problems with the previous results. (1) The usefulness of maps acquired by high energy techniques is limited by lower resolution (Wu, 2012). (2) The coverage of 86 87 optical maps is poor due to the illumination and topography (Xia et al., 2019), especially in high latitudes or Polar Regions (Lucey et al., 2000; Wu, 2012; Xia et al., 2019). (3) There are no obvious 88 89 optical diagnostic spectral characteristics in Feldspar, Ilmanite, Glass and Iron except for Pyroxene 90 and Olivine which would increase the uncertainty in the derivation results (Allen et al., 2012). (4) 91 The space weathering (Fischer and Pieters, 1994) and particle sizes (Fischer and Pieters, 1996) also 92 have more significant effects on the reflectance spectrum.

93 Many groups have payed attention to the thermal infrared techniques (e.g., Paige et al., 2009; 94 Greenhagen et al., 2010; Donaldson et al., 2012; Allen et al., 2012; Tang et al., 2016). In general, the 95 emissivity spectrum of silicate minerals are mainly manifested in complicated condition of Si-O 96 complex ion vibration. It is clear for a prominent emissivity maximum centered near 8 um called as 97 CF caused by rapid change of the real part of the refractive index before the Christiansen frequency (Conel, 1969). With the increase of silicate mineral polymerization degree, the Si-O bond force 98 99 constant increases continually which leads the shorter wave shift of the CF wavelength position. For 100 example, the CF of Olivine (15555, LR-CMP-169), Pyroxene Clinopyroxene Augite (70017, 101 LR-CMP-175) and Feldspar Plagioclase (70017, LR-CMP-177) from Reflectance Experiment 102 Laboratory (RELAB) spectrum are 8.578 µm, 8.480 µm and 8.027 µm respectively.

103 Meanwhile, Si-O₄ tetrahedral complex is connected by metal cations which are definitely 104 connected with the change of Si-O bond force constant, vibration frequency and CF position. The 105 increase of TiO₂, FeO and MgO abundances leads to the longer wavelength of CF, but the increase of 106 Al₂O₃ and CaO abundances leads to the shorter wavelength of CF. In all Apollo samples, the CF of 107 TiO₂-rich (9.48 wt.%), FeO-rich (18.5 wt.%) and MgO-rich (10.8 wt.%) is 8.389 µm (LR-CMP-057, 108 71061, 20-45 µm), and the CF of Al₂O₃-rich (29.4 wt.%) and CaO-rich (16.1 wt.%) is 8.172 µm 109 (LR-CMP-114, 67461, 20-45 µm). Moreover, comparing to reflectance spectral characteristics, the 110 CF is barely affected by approximate vacuum environment and roughness on lunar surface 111 (Donaldson et al., 2012). Therefore, it can be believed that the CF can be the direct basis for lunar 112 element inversion (Paige et al., 2009).

Until now, Apollo 17 (1972), Clementine (1994), LRO (2009) missions had successively carried infrared scanning radiometer by NASA. In addition to analyzing the thermal physical properties of lunar surface materials, NASA tried to investigate lunar surface oxide abundances using emissivity spectrum (Paige et al., 2009). With thermal infrared data acquired by LRO Diviner in orbit, global CF map (32 pixels/degree) between 60°N/S was calculated and silicate mineralogy abundances were calculated firstly (Greenhagen et al., 2010). Then, the relationships between FeO abundances and CF 119 were discussed, FeO abundances for three Lunar Pyroclastic Deposits were investigated (Allen et al.,

2012) and the highest resolution CF product (128 pixels/degree) was produced (Williams et al., 2017).
However, the global maps for SiO₂, TiO₂, Al₂O₃, FeO, MgO and CaO abundances based on Diviner
CF did not exist.

123 This paper presents six global elemental abundance maps $(60^{\circ}N/S)$ with the highest both 124 resolution and coverage up to now derived from Diviner thermal infrared data. For this purpose, the 125 global models by the univariate regression methods between Diviner CF and ground truths of 126 elemental abundances are established. The six elemental abundance maps are calculated based on 127 Diviner Level 3 Standard CF product with the resolution of 128 pixels/degree and the coverage of 99.86%. Analyses on global, different geologic units, crater and ejecta surfaces are carried out to find 128 129 the elemental abundances distribution on lunar surface. Comparisons with the results acquired by LP, 130 Clementine and CE-1 are made to investigate the accuracy of Diviner derivation results. Further 131 applications are also tried to verify the usefulness of Diviner elemental abundances in geological 132 problems.

133 **2. Data and methods**

134 **2.1 Diviner Level 3 Standard CF products**

Diviner is a pushbroom mapping radiometer with nine 21-element arrays detectors onboard 135 LRO launched on June 18, 2009 and still in orbit. The maximum resolution is approximately 200 m 136 137 (Greenhagen et al., 2010). The fields of view are 320m in track and 160m cross track for 50km 138 altitude (Paige et al., 2009). The detailed instrument description had been given by Paige et al. (2009). 139 In over ten years (July 5, 2009 to December 15, 2019), more than 480 billion calibrated radiance 140 values had been acquired covering a full range of lunar position and time. The data were divided into 141 Experiment Data Record (EDR), Reduced Data Record (RDR), Gridded Data Record (GDR_L2 and 142 GDR_L3) and Polar Resource Products (PRP) by NASA.

143 Greenhagen et al. (2010) had calculated the effective emissivity and Standard CF using radiance 144 and brightness temperature measurements on the three 8 µm-region channels near 7.55-8.05 µm, 145 8.10-8.40 µm and 8.38-8.68 µm based on Normalization Emissivity Method (NEM). For each pixel 146 in one RDR data (10 minutes, lunar time), the maximum brightness temperature is firstly calculated 147 using a parabolic fit for three 8 µm-region channels brightness temperature. Meanwhile, the 148 calibrated radiances are binned for each channels. Then the effective emissivity are calculated based 149 on Diviner spectral response and Plank Equation. Finally, another parabolic fit is used to determine 150 the Standard CF which is averaged in one RDR data. The detailed method description had been given 151 by Greenhagen et al. (2010).

For the Diviner GDR Standard CF product calculated by Greenhagen et al. (2010) and the Normalized to Equatorial Noon (MEN) CF product provided by "best effort" of Diviner science team using Diviner data from 2009-07-05 to 2010-11-30, the resolution is 32 pixels/degree and the coverages are 52.4% and 82.5% of lunar surface (60°N/S). As LRO mission proceeds, the Standard CF product with the resolution of 128 pixels/degree had been provided by Diviner science team (Williams et al., 2017) using Diviner data from 2009-07-05 to 2016-10-23. The highest resolution of CF products is 237 m/pixel at the equator. It means that the resolution of Diviner CF map is slightly
lower than 200 m/pixel of CE-1 Interference Imaging Spectrometer (IIM) data, and apparently higher
than 1 km of Clementine Ultraviolet-Visible (UV-VIS) data (Lucey et al., 2000) and 15 km of LP
Gamma Ray (GR) data (Prettyman et al., 2006).

The 128 pixels/degree Standard CF product has 707,788,800 pixels (46080×15360). There are 163 1,024,549 pixels (0.14%) with null values that the coverage is 99.86%. The missing pixels are 164 scattered and not concentrated in a specific region. Comparing to 100% of LP GR (Prettyman et al., 165 2006), 98% of Clementine UV-VIS (Lucey et al., 2000) and 75.4% of CE-1 IIM (Xia et al., 2019), 166 the Diviner CF coverage is the second highest.

For Diviner CF product, the minimum value is 7.00 μm and the maximum value is 9.456 μm.
2,651,599 pixels (0.23%) are between 7.00 μm and 7.80 μm. 610,868 pixels (0.09%) are between
8.55 μm and 9.456 μm. The maximum CF percentage (21.65%) is concentrated in 8.10-8.15 μm
(Figure 1). In general, the CF show a Gaussian distribution trend and the average value is 8.1659 μm.



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Figure 1. CF Histograms of Diviner Level 3 Standard CF product (128 pixels/degree)

173 The Diviner Level 3 Standard CF map $(60^{\circ}N/S)$ is shown in Figure 2. There are approximately 174 uniform CF on the inner of mare surfaces such as Mare Imbrium and Mare Serenitatis, even Mare Orientale and Mare Marginis, whereas CF are constantly changing on highland surface because of 175 176 small impact events or ejecta from big impact events such as Tycho, Byrgius and Giordano Bruna. 177 On the whole, the difference of global CF between maria and highland is 0.159 µm (8.294-8.135). 178 The maximum difference of mare CF between Mare Tranquillitatis and Mare Orientale (south) is 179 about 0.174 µm (8.352-8.178) The highland CF in the southern hemisphere are significantly higher 180 than those in the northern hemisphere.

181 The CF on the crater surfaces are the most complex. The older craters such as Plato have higher 182 CF (8.326 μm), whereas younger craters (Copernicus, Kepler, Timocharis, Archimedes et.) are the 183 opposite. The other craters such as Aristarchus and Tsiolkovoskiy show unusual conditions where the 184 CF of the central peak and wall are 0.2 μm lower than those of the plain. On the surfaces of the South Pole-Aitken (SPA) basin, the CF are the intermediate value between maria and highland. Meanwhile, compared to reflectance maps there are more intuitive differences. The CF on plain and mare surfaces are higher than that on wall and highland surface where reflectance values are the opposite. In other word, bright and dark in reflectance map are interchanged in CF map.



Figure 2. The Diviner Level 3 Standard CF map (60°N/S). The projection system is GCS_moon_2000. The resolution
is 128 pixels/degree (237 m/pixel at the equator). The coverage is over 99.86% (1,024,549 null in 707,788,800 pixels).
The map scale is 1:250,000,000. The upper and lower limits of CF are shown in the legend. The base map is LRO
Wide Angle Camera (WAC) map of 100 meters/pixel (Scholten et al., 2012).

194 **2.2 Elemental abundances and Diviner CF for lunar sites**

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195 Until now, based on elemental abundance measurements from the returned samples or actual 196 location with geographic information (Lucey et al., 2000; Liu et al., 2015), 48 lunar sampling sites 197 can be confirmed including 44 Apollo sites (11, 12, 14, 15, 16, 17), 3 Luna sites (16, 20, 24) (Blewett 198 et al., 1997, Jolliff et al., 1999; Lucey et al., 2000) and CE-3 site (Wu et al., 2018). With the 199 elemental abundances of 118+6 Apollo samples, 3 Luna samples (Blewett et al., 1997, Jolliff et al., 200 1999; Lucey et al., 2000) and CE-3 rover measurements (Wu et al., 2018), the mean abundances of 201 six elements in 48 lunar sites are recalculated (Table 2). The mean abundance values of Apollo 202 samples can eliminate the unreliable abundances of a particular element in the particular sample and 203 be a better indicator for the site compositions.

For six elements, the mean SiO₂ abundance is 44.2853 wt.% and the standard deviation (STD) is 205 2.1417 wt.% which demonstrates the nearly uniform distribution of lunar surface silicate minerals. 206 However, there are obvious differences in the abundances of other five elements, especially Al_2O_3 207 (STD of 5.6438 wt.%) and FeO (STD of 4.8219 wt.%). The TiO₂, FeO and MgO abundances on 208 Apollo 16 site in the highland are significantly lower than in other Apollo sites in the maria, while the 209 Al_2O_3 and CaO abundances are the opposite. The elements abundances of Apollo 14 confirm that the 210 site is located a transitional region between highland and maria. The biggest differences of SiO2,

 $211 \qquad TiO_2, \ Al_2O_3, \ FeO, \ MgO, \ CaO \ are \ 8.40 \ wt.\%, \ 9.60 \ wt.\%, \ 19.07 \ wt.\%, \ 18.29 \ wt.\%, \ 7.15 \ wt.\%, \ 8.37 \ wt.\%, \ 8.37 \ wt.\%, \ 18.29 \ wt.\%, \ 7.15 \ wt.\%, \ 8.37 \ wt$

wt.% respectively. Moreover, the relationship between SiO₂, Al₂O₃, CaO and TiO₂, FeO, MgO show
 inversely correlation (Heiken et al., 1991) which provides a verification indication for the elemental

abundances of mare and highland surface.

Lunar Sites	SiO ₂	TiO ₂	Al_2O_3	FeO	MgO	CaO	CF
A11	41.4	7.5	13.455	15.8	7.91	12.275	8.377
A12	46.216	3.08	13.77	15.48	9.79	10.54	8.298
A14-LM	48.2	1.73	17.69	10.5	9.52	10.32	8.265
A14-Cone	47.536	1.6	17.533	10.3	9.565	10.95	8.198
A15-LM	46.75	1.9	14.265	15	10.585	10.27	8.266
A15-S1	46.8	1.6	13.2	16.8	10.7	10.25	8.290
A15-S2	46.315	1.3	17.378	11.5	10.54	11.53	8.320
A15-S4	46.26	1.2	13.4	16.6	11.6	8.9	8.235
A15-S6	46.768	1.5	16.45	12.1	10.475	10.975	8.256
A15-S7	46.29	1.1	15.3	13.9	11.567	10.267	8.291
A15-S8	46.75	1.7	14.45	15.2	10.65	10.25	8.285
A15-S9	46.46	1.8	12.5	16.9	11	9.8	8.320
A15-S9a	46.52	2	10.3	20.4	11.25	8.3	8.281
A16-LM	45.285	0.6	26.575	5.6	6.15	15.78	8.155
A16-S1	44.846	0.6	26.714	5.4	6.056	15.604	8.165
A16-S2	44.575	0.593	26.73	5.44	6.477	15.55	8.163
A16-S4	45.067	0.537	27.53	4.853	5.36	15.71	8.137
A16-S5	45.063	0.7	26.187	5.9	6.183	15.063	8.184
A16-S6	45.135	0.7	27.133	6	6.867	16.4	8.180
A16-S8	45.078	0.6	26.545	5.4	6.223	15.365	8.175
A16-S9	44.88	0.6	26.283	5.7	6.343	15.49	8.178
A16-S11	45.142	0.372	29.372	3.95	4.455	16.67	8.048
A16-S13	45.015	0.5	28.067	4.8	6	14.65	8.078
A17-LM	40.8	8.5	12.1	16.6	9.8	11.1	8.262
A17-S1	39.9	9.6	10.9	17.8	9.6	10.8	8.338
A17-S2	45	1.5	20.7	8.7	9.9	12.8	8.213
A17-S3	44.9	1.8	20.4	8.7	10.2	12.9	8.263
A17-S5	39.8	9.9	10.9	17.7	9.6	10.8	8.303
A17-S6	43.5	3.4	18.3	10.7	10.8	12.2	8.234
A17-S7	43.7	3.9	17.	11.6	10.1	11.9	8.231
A17-S8	43.4	4.04	16.68	12.24	10.42	11.56	8.307
A17-S9	42.1	6.4	13.9	15.4	10	11.3	8.317
A17-LRV1	41.3	8	12.6	16.3	9.4	11.2	8.333
A17-LRV2	43.1	4.4	16.1	13.4	10.3	11.9	8.307
A17-LRV3	42.2	5.5	14.4	14.8	10.4	11.3	8.302
A17-LRV4/S2a	44.9	1.3	21.4	8.5	9.6	12.8	8.141
A17-LRV5	44.8	1.3	21.4	8.5	9.6	12.8	8.282
A17-LRV6	44.5	2.6	19.4	10.3	9.9	12.5	8.258

Table 2 The elemental abundances (wt.%) and CF (µm) for 48 lunar sampling sites used in this study.

A17-LRV7	41.8	6.8	12.8	16.1	10.3	10.7	8.311
A17-LRV8	41.9	6.6	13.5	15.7	9.9	11.3	8.345
A17-LRV9	42.2	6.1	14.3	14.6	9.8	11.3	8.321
A17-LRV10	43.5	3.7	17.5	11.2	10.5	12.1	8.272
A17-LRV11	43.2	4.5	16.3	12.7	10	11.9	8.312
A17-LRV12	39.9	9.97	11.2	17.4	9.36	10.8	8.324
Luna 16	41.7	3.3	15.3	16.7	8.8	12.5	8.341
Luna 20	45.6	0.5	22.9	7.5	9.15	14.5	8.239
Luna 24	45.36	1.0	12.1	19.6	10	11.1	8.359
CE-3	-	4.31	12.11	22.24	8.61	9.72	8.362

Description: References for compositions see Lucey et al. (2000, Table 1), Jolliff et al. (1999, Table 4)
and Blewett et al. (1997, Table1). The increased samples are 12029 (A12), 62231(A16-S2, Size of
20-45 μm), 64801 (A16-S4), 67711 (A16-S11) and 78481 (A17-S8). The composition of A15-LM
comes from 15013 and 15021, A16-S6 comes from 66031, 66041 and 66081, CE-3 comes from Wu
et al. (2018). References for Latitude and Longitude of Apollo and Luna sites see Lucey et al. (2000,
Table1) and CE-3 site see Liu et al. (2015). The CF are manually extracted from Diviner Level 3
Standard CF products.

223 For Apollo, Luna and CE-3 missions, the detecting areas of lunar surface had expanded from a 224 few hundred square meters at Apollo 11 to tens of thousands of square meters at Apollo 17. 225 Correspondingly, with the maximum resolution of 237 m, the CF are manually extracted from 226 Diviner Level 3 Standard CF map depending on the sample returned sites (Lucey et al., 2000) or the 227 rover measured position (Liu et al., 2015) of 1×1 pixels (Table 2). In remote sensing data, the CF of 228 lunar sites have obvious regularity and rationality. The CF of Apollo 11, 12, 15, 17, Luna 16, 20, 24 229 and CE-3 in maria are obviously higher than that of Apollo 16 in highland, and the maximum 230 difference is 0.329 µm (A11 and A16-S11). It's inferred again that Apollo 14 is not really highland because the CF extracted from the map are between highland and mare values. 231

232 2.3 Mathematical modelling

For six elements, the relationships between Diviner CF and FeO abundance had been discussed using linear regression (Allen et al., 2012). But, SiO₂, TiO₂, Al₂O₃, MgO and CaO abundances have not been correlated with CF. Therefore, six univariate regression models (exponential, exponential, linear, quadratic, cubic, logarithmic) are used for experimental analysis and the best models are determined with the highest R-square (R^2).

238 The returned samples and field measurement have certain representativeness and randomness. 239 Diviner thermal infrared data are also comprehensive information within tens of thousands of square 240 meters. These two factors together lead to predictable error in the determination of elemental abundances or Diviner CF at the sampling pixels. In the initial relationships of 48 datasets, the same 241 conclusion can be obtained. The R² of SiO₂, TiO₂, Al₂O₃, FeO, MgO and CaO models are 0.1546, 242 243 0.3894, 0.7491, 0.7136, 0.4655 and 0.5376 respectively. Hence, before establishing the Diviner 244 prediction models, the outliers of 48 datasets are analyzed and culled firstly. Because the outliers can 245 reduce the accuracy of models and prediction results that the influence should be eliminated.

In this study, the outlier rejections are tested repeatedly by 2, 1.5, 1 and 0.5 Root Mean Square

Error (RMSE) of 48 datasets. For example FeO (Figure 3), the RMSE of 48 dataset is 2.608. Under the limitations of 2, 1.5, 1 and 0.5 RMSE, the number of samples decreases from 48 to 16 and the R^2 of models increases from 0.7136 to 0.9622 which illustrates the need for constraints. However, excessive rejection make no sense. 1 RMSE (Figure 3, Green line) is selected based on error theory and experimental data. Finally, the relationships between Diviner CF and elemental abundances for sampling pixels are established and thermal infrared prediction models are provided.



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Figure 3. The relationship between FeO abundance and Diviner CF for 48 lunar sampling sites and the outlier rejections of 48 datasets.

256 **3. Results**

257 **3.1 Diviner models**

Six univariate regression types of approaches were chosen to build the relationships between Diviner CF and elemental abundances at 48 lunar sampling sites. The relationships with the highest R^2 were selected. Except for TiO₂ that the relationship can be described as an exponential correlation, the relationships of the other five oxides can be described as linear correlations.

The points under the limitations of 1 RMSE of 48 datasets were regressed using both linear and exponential relationships. Figure 4 shows the relationship with the highest R^2 value, for each of the six oxides tested in this study. The linear R^2 of Al₂O₃ is highest (0.9043), followed FeO (0.8843), MgO (0.7853), CaO (0.7504) and R^2 of SiO₂ is lowest (0.604). The exponential R^2 of TiO₂ is 0.704 and only higher than the linear R^2 of SiO₂. Thus, it can be speculated that the prediction accuracy of Al₂O₃ and FeO is higher than that of MgO, CaO and TiO₂, the deviation of SiO₂ is largest.

For six Diviner models, the models are described by the regression slope (p1, p2 or a, b). The SiO₂, Al₂O₃, CaO abundances are positively correlated with CF, whereas the TiO₂, MgO, FeO abundances are negatively correlated with CF. It means that the influences of SiO₂, Al₂O₃, CaO abundances on CF are weaker near longer wavelength position (8.55) and stronger near the shorter wavelength position (7.80), whereas that of TiO₂, MgO, FeO abundances are the opposite. Thus, the CF of 7.00 to 7.80 μ m or 8.55 to 9.456 μ m should be deleted during calculating surface elemental

abundances. Moreover, the maximum elemental abundances from 806 published Apollo samples
should be added to the prediction results, especially for the exponential correlation of TiO2 (14.5
wt.%, 77538).



Figure 4. The relationships between Diviner CF and elemental abundances under the limitations of 1
RMSE of 48 datasets. The number of points SiO₂, TiO₂, Al₂O₃, FeO, MgO and CaO are 30, 38, 31,
35, 31and 33 respectively.

- 283 **3.2 Prediction Results**
- **3.2.1 Global**

707.7 million elemental abundance values are calculated from the Diviner Level 3 Standard CF
map (60°N/S) based on Diviner models. Six separate maps are produced with the resolution of 128
pixels/degree and the coverage of over 99% (Figure 5). After deleting the null pixels, global average
elemental abundances of SiO₂, TiO₂, Al₂O₃, FeO, MgO and CaO are 44.85 wt.%, 1.46 wt.%, 23.60
wt.%, 7.82 wt.%, 7.29 wt.% and 13.91 wt.% respectively and the total is 98.93 wt.%.





Figure 5. The maps of six elemental abundances calculated from the Diviner Level 3 Standard CF map. (a) SiO_2 abundance map, (b) TiO_2 abundance map, (c) Al_2O_3 abundance map, (d) FeO abundance map, (e) MgO abundance map, (f) CaO abundance map. The white pixel is 0 value in FeO abundance map (d). Referring to Figure 2 for geographic information. The upper and lower limits of each elemental abundance are shown in the legend.

306 Corresponding to positive and negative correlations, TiO₂, FeO and MgO abundances increase

obviously with the increase of CF on typical mare surfaces while SiO₂, Al₂O₃ and CaO abundances
 decrease. It is the opposite on highland surface. But on separate geological unit surfaces such as SPA
 basin there are an approximately uniform distribution of six elemental abundances.

On global surfaces (Figure 6) some of abundance differences can be made by craters and ejecta where big and systematic differences in average elemental abundances are presented (Table 2). On the mare surfaces, FeO (14.14 wt.%) and Al_2O_3 (15.49 wt.%) are higher than CaO (11.42 wt.%), MgO (9.76 wt.%) and TiO₂ (3.76 wt.%), while on the highland surface, Al_2O_3 (25.51 wt.%) and CaO (14.50 wt.%) are higher than FeO (6.14 wt.%), MgO (6.71 wt.%) and TiO₂ (0.94 wt.%). Especially on SPA basin surface, six elemental abundances are the intermediate values between maria and highland and is close to the global average elemental abundances.



Figure 6. The areas (60°N/S) of three large geological units including maria, highland and SPA basin. The green area are maria and crater plains (Nelson et al., 2010), the yellow area is highland (Nelson et al., 2010), and the wathet blue area is SPA basin by this study, the red points are 47 verification points with the size of 0.078125°×0.078125°. Referring to Figure 2 for geographic information and the base map.

317

322

		A	verage (wt.%	Proportions (%)			
	Global	bal Maria Highland			Maria	Highland	Basin
SiO_2	44.85	43.19	45.25	44.57	16.65	78.04	5.31
TiO ₂	1.46	3.76	0.94	1.57	44.52	49.75	5.73
Al_2O_3	23.60	15.49	25.51	22.24	11.35	83.62	5.04
FeO	7.82	14.14	6.14	8.71	34.71	59.06	6.23
MgO	7.29	9.75	6.71	7.71	23.13	71.22	5.65
CaO	13.91	11.42	14.50	13.49	14.20	80.61	5.18

Table 2 The average values and proportions of elemental abundances on maria, highland and SPA surface

323 It's obvious that the proportions of six elemental abundances in each geological unit are 324 different (table 2). For TiO₂, mare surfaces have almost the same abundances with highland surface. 325 For FeO and MgO, mare surfaces have 34.71% and 23.13% of the global abundances corresponding 326 to 59.06% and 71.22% in highland. Increasing differences, percentage of SiO₂ is 16.65% in maria and 78.04% in highland. Maximally for CaO and Al_2O_3 , the difference between maria and highland is 66.41% and 72.27%. The SPA Basin surface have almost a similar percentage of six elemental abundances which are 5.18% to 6.23%.

330

3.2.2 Maria, Highland and SPA

331 The mean elemental abundances of major maria are showed in table 3. On mare plain surfaces of nearside such as Mare Tranquillitatis, TiO₂, FeO, MgO abundances are highest and SiO₂,Al₂O₃, 332 333 CaO abundances are lowest on all mare surfaces. On Oceanus Procellarum and Mare Inbrium 334 surfaces with the vast area, the Aristarchus and Timocharis impact events don't substantially change 335 elemental abundances, but slightly increased SiO₂, Al₂O₃, CaO abundances and decreased TiO₂, FeO, 336 MgO abundances. It is more obvious for these differences between Oceanus Procellarum (south) and 337 Oceanus Procellarum surfaces. The mare surfaces surrounded by highland or in high latitudes such as 338 Mare Humboldtianum or Mare Frigoris, TiO₂, FeO, MgO abundances are lower and SiO₂, Al₂O₃, 339 CaO abundances are higher where elemental abundances are seriously disturbed by the highland 340 anorthosite materials and the boundary is getting blurry. On the farside, two maria including Mare 341 Orientale and Mare Moscoviense (Mare Ingenii located in SPA basin) are surrounded by highland 342 and have incomprehensible material distribution especially at the boundary because of the unique 343 geological structure and the random distribution of highland anorthosite materials. It is also obvious for Mre Orientale (south) surfaces that TiO₂, FeO, MgO abundances are lowest and SiO₂,Al₂O₃, CaO 344 345 abundances are highest. In a word, on the farside each mare element abundance is similar to that on 346 highland surface and the mare material characteristics become indistinct.

Table 4 The mean values of six elemental abu	indances of lunar major maria (wt.%)
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Mare	SiO ₂	TiO ₂	Al_2O_3	FeO	MgO	CaO
Anguis	43.89	2.26	18.92	11.07	8.71	12.48
Australe	44.06	1.98	19.72	10.41	8.47	12.72
Cognitum	43.18	3.50	15.48	14.12	9.75	11.42
Crisium	43.06	3.99	14.89	14.65	9.93	11.24
Fecunditatis	42.98	4.20	14.51	14.98	10.05	11.12
Frigoris	43.76	2.50	18.29	11.71	8.90	12.28
Humboldtianum	44.33	1.82	21.09	9.39	8.05	13.14
Humorum	43.21	3.59	15.59	14.05	9.72	11.45
Imbrium	42.97	4.19	14.46	15.03	10.06	11.11
Insularum	43.47	2.94	16.88	12.88	9.33	11.85
Marginis	43.17	3.62	15.40	14.17	9.78	11.40
Moscoviense	43.33	3.11	16.17	13.47	9.54	11.63
Nectaris	43.53	2.75	17.17	12.58	9.24	11.94
Nubium	43.65	2.52	17.76	12.08	9.06	12.12
Orientale	44.13	1.87	20.08	10.03	8.36	12.83
Orientale (south)	44.69	1.23	22.79	7.74	7.54	13.66
Serenitatis	42.83	4.49	13.77	15.65	10.27	10.90
Smythii	43.00	4.02	14.59	14.92	10.03	11.15
Spumans	43.09	3.68	15.03	14.50	9.89	11.28

Temporis	44.41	1.40	21.46	8.71	7.94	13.26
Tranquillitatis	42.42	6.19	11.77	17.46	10.88	10.28
Undarum	42.93	4.21	14.25	15.21	10.13	11.04
Vaporum	42.87	4.39	13.95	15.48	10.22	10.95
Oceanus Procellarum	42.99	4.13	14.56	14.96	10.03	11.14
Oceanus Procellarum (south)	43.42	2.90	16.63	13.07	9.41	11.77

348 Description: The bold value is the extreme value located in Mare Orientale (south) and Mare Tranquillitatis349 respectively

Highland surface are characterized by higher SiO₂, Al₂O₃, CaO and lower TiO₂, FeO, MgO. However, each abundance is not uniformly distributed. The high SiO₂, Al₂O₃ and CaO abundances are mainly concentrated on Tycho, Byrgrius, Wegener, Sterinus and Giordano Bruno crater and ejecta surfaces and the periphery of Mare Orientale. The FeO and MgO abundances are evenly distributed on highland surface where these are higher on the southern and eastern hemisphere in contrast to the northern and western hemisphere except for crater and ejecta surfaces. Remarkably, TiO₂ abundance is almost zero on whole highland surface.

The SPA basin is a special geological unit and elemental abundances are different from maria and highland. TiO_2 , FeO and MgO abundances are lower than that on mare surfaces and Al_2O_3 , CaO abundances are too lower than that on highland surface. The abundance differences of SPA plain surface between any two elements are the most indistinct in all geological units.

361

3.2.3 Crater and ejecta

362 On the largest number of craters and ejecta surfaces, elemental distribution is characteristic and 363 discussed separately. In maria, Kepler and Copernicus crater and ejecta surfaces typically have higher SiO₂, Al₂O₃, CaO abundances and lower TiO₂, FeO, MgO abundances than the surrounding surfaces. 364 365 The similar situations may occur on Aristarchus, Seleucus, Kraff, Cardanus, Timocharis, Timochari 366 crater and ejecta surfaces. However, On the surfaces of Herodotus to east of Aristarchus, Prinz to 367 northwest of Aristarchus, Eratosthenes, lambert, C.herschel around Copernicus, Archimedes, 368 Autolycus around Aristillus, each element abundance is roughly the same as that on the surrounding 369 areas where the abundance boundary of impact crater appears to disappear.

In highland, most craters and all ejecta surfaces have higher SiO₂, Al₂O₃, CaO abundances and lower TiO₂, FeO, MgO abundances. Another areas of such abundance are the highland around Mare Orientale where the area is almost the same as that of Tycho crater and ejecta. However, a few craters including Plato and Lomonosov are exceptions where crater plains surfaces are filled with more TiO₂, FeO, MgO and the crater rim have more SiO₂, Al₂O₃, CaO. The central peak of several craters including Tsiolkovskiy and Aitken but not Tycho have obvious elemental abundance difference compared with the impact crater plains.

377 **4. Discussions**

378 **4.1 Model analysis**

As repeatedly mentioned above, this study's models are the "global" thermal infrared models in which all of the sampling point information have been used. However, the correlations of the initial relationships of 48 datasets are relatively lower. There are some reasons among which two factors 382 must be mainly mentioned.

383 One is the representativeness of elemental abundances of lunar samples or rover measurements. 384 Traditionally, elemental abundances of each sampling pixel had been determined by the average 385 abundance value of 1-12 samples. The problems whether the values can accurately characterize the 386 elemental abundance of sampling pixel and what is the right number of sampling pixels are one topic 387 worthy of discussion. For the establishment of Diviner models, more sampling points and wider 388 sample selection can provide reliable data sources. For example, the number of samples and sampling pixels of Lucey et al. (2000) for modeling are 118 and 47; Wu (2012) are 76 and 18; Xia et 389 al. (2019) are 100 and 39; this study are 124 and 48. Each group hopes to improve the usefulness and 390 391 accuracy of the models by increasing the representativeness of the ground truth value. However, it 392 may be one of the universal problems in the process of remote sensing modeling, because the number 393 of sampling sites is limited and most are concentrated in mare of nearside except for Apollo 17.

394 The other is the accuracy of the spectral values of mixed pixel. Even in the smallest remote 395 sensing pixel, mixed substances are also present. Adding to the effects of topography and scale, remote sensing data are all mixed information. The problem whether the measurements can 396 397 accurately characterize the spectral values of mixed pixel is another topic worthy of discussion. With 398 the different resolutions, the number of pixels for modeling represented by the measurement of 399 Clementine, LP, CE-1 and Diviner is different. For example, the number of Lucey et al. (2000) is 400 from 1×1 to 6×6 pixels; Wu (2012) is from 2×2 to 10×10 pixels; Xia et al. (2019) is 1×1 pixel; this 401 study is 1×1 pixel. Theoretically, in the smaller spatial scale, the remote sensing measurement can 402 better represent the spectral characteristics of mixed pixel. Therefore, the 1×1 pixel is more 403 reasonable for modeling. Meanwhile, mixed spectral decomposition (Villa et al., 2010) has been 404 studied for a long time, but a few potential complexities have not been fully explored and the result is 405 always unsatisfactory.

The above two reasons jointly determine the relationships between elemental abundances and CF in Diviner models. In addition to selecting all current sampling pixels with ground truth value, trial and error of the two methods are also performed in our study.

One is to increase the number of CF pixels in order to obtain high R^2 . For example FeO (Figure 409 7), with the pixels increasing from 1×1 to 21×21 , the R² first increase and then decrease, the 410 maximum R^2 is 0.7776 with 9×9 pixels (2×2km) that the same area was used by Allen et al. (2012). 411 The similar thing happens with the other five oxides except that the maximum R^2 is different for 412 413 0.1652 of SiO₂ (5×5 pixels), 0.4358 of TiO₂ (3×3 pixels), 0.8381 of Al₂O₃ (9×9 pixels), 0.6045 of 414 MgO (13×13 pixels) and 0.6632 of CaO (15×15 pixels) respectively. These demonstrates the uncertainty of elemental abundances and Diviner CF in pixels. No obvious regularity is found. The 415 R^2 don't improve significantly too, especially for SiO₂. Therefore, the outliers can only be 416 417 eliminated.



418

Figure 7. The relationships between FeO abundances and Diviner CF for 1×1 (blue square), 9×9 (red cross), 21×21 (green cross) pixels at 48 sampling sites. The regression slopes (a, b) and R² are shown in legend. The R² for 1×1 , 3×3 , 5×5 , 7×7 , 9×9 , 11×11 , 13×13 , 15×15 , 17×17 , 19×19 , 21×21 pixels are 0.7136, 0.7395, 0.7713, 0.7716, 0.7776, 0.7636, 0.7570, 0.7534, 0.7476, 0.7399, 0.7331 respectively. The maximum R² is 0.7776 with 9×9 pixels.

423 The other is the RMSE limitations of 48 datasets. In Table 5, the initial R^2 show that SiO₂, TiO₂, 424 MgO and CaO abundances are not closely correlated with Diviner CF. Under the limitations of 1.5 425 and 1 initial RMSE, dozens of outliers are eliminated. The correlations are not radically changed. But, 426 the coefficients (p1, p2 or a, b) are slightly adjusted and the R^2 increase significantly especially for 427 SiO₂ from 0.1546 to 0.604.

Coofficients			In	itial RMSE		
Coefficients	SiO ₂ (48)	TiO ₂ (48)	Al ₂ O ₃ (48)	FeO(48)	MgO(48)	CaO(48)
p1 or a	-11.28	4.053e-64	-64.77	54.01	16.33	-20.36
p2 or b	137.4	69.61	552.7	-433.8	-125.7	180.3
R^2	0.1546	0.3894	0.7491	0.7136	0.4655	0.5376
RMSE	1.991	2.241	2.858	2.608	1.334	0.5276
Coofficients			1.5 *	initial RM	SE	
Coefficients	SiO ₂ (42)	TiO ₂ (43)	$Al_2O_3(42)$	FeO(42)	MgO(43)	CaO(43)
p1 or a	-8.19	1.692e-77	-65.15	53.31	20.74	-21.41
p2 or b	112.2	84.11	556.1	-428.3	-162.1	189.1
R^2	0.1341	0.5935	0.8224	0.8248	0.6868	06923
RMSE	1.619	1.410	2.408	1.944	1.043	1.097
Coofficients			1.0 *	initial RM	SE	
Coefficients	SiO ₂ (30)	TiO ₂ (38)	Al ₂ O ₃ (31)	FeO(35)	MgO(31)	CaO(33)
p1 or a	-13.06	7.473e-75	-63.46	57.56	19.25	-19.47
p2 or b	151.5	81.24	541.8	-463.3	-149.9	172.9
R^2	0.604	0.704	0.9043	0.8843	0.7853	0.7504
RMSE	0.8859	1.106	1.67	1.563	0.7589	0.833

428 Table 5 The relationships between elemental abundances and Diviner CF under the limitations of 1, 1.5, initial RMSE

429 Meanwhile, from the perspective of remote sensing theory, spectral values or sampling 430 elemental abundances can better describe the mixed pixel information with highest resolution. 431 Therefore, the latter method is adopted in our study. Under 1 initial RMSE, the elemental abundances 432 and Diviner CF of 1×1 pixel at 48 sampling sites are the data sources for Diviner models.

433

4.2 Comparison with previous work

The similar results with six Diviner elemental abundances had been investigated by Clementine UV-VIS data (Lucey et al., 2000), LP GR data (Prettyman et al., 2006) and CE-1 IIM data (Wu, 2012; Xia et al., 2019). They can also be compared to each other. Because of the different resolution, the comparisons are performed from the 47 verification points, lunar geological units and three separate surfaces.

439

4.2.1 Point elemental abundances

The average abundances from 47 verification points are extracted including 33 points from 150°W to 150°E and 30°N to 30°S, 2 CE sampling points, 3 pyroclastic deposit points (Allen et al., 2012), northwestern farside highland (NWFH), southern nearside highland (SNH) and northeastern farside highland (NEFH) points (Wu, 2012), 2 SPA points, Plato point and 3 mare points (Figure 6, red points). The verification area size is 0.078125°×0.078125° where latitude and longitude in the lower left corner are the reference. In this way, the number of extracted pixels of LP, Clementine, Diviner and CE-1 are 1, 4, 64 and 90-100 respectively before deleting the null values.

447 Table 6 shows the Pearson Correlation Coefficients (PCC) of 47 average elemental abundances 448 for six results. The PCC between Diviner and LP are lower than that between Diviner and other 449 results. There are the highest correlations between Diviner and CE-1 (Wu) for SiO₂ and MgO (0.8861 450 of PCC). For the other four oxides, the highest correlations are between Diviner and CE-1 (Xia) that 451 the highest PCC of TiO₂ is 0.0.9414 and the lowest PCC of CaO is 0.9176.

452	

Table 6 the PCC of 47 average elemental abundances for five results each other.

data				Diviner]	LP		
uata	SiO2	TiO2	A12O3	FeO	MgO	CaO	SiO2	TiO2	A12O3	FeO	MgO	CaO
LP	-	0.4675	0.7296	0.8263	0.6668	0.5734	-	1	1	1	1	1
Clementine	-	0.7343	-	0.8846	-	-	-	0.4239		0.9229	-	-
CE-1(wu)	0.8861	0.8379	0.8862	0.9067	0.8861	0.8862	-	0.2036	0.6880	0.8755	0.6145	0.6634
CE-1(Xia)	0.6813	0.9414	0.9363	0.9397	0.7210	0.9176	-	0.5668	0.8916	0.9014	0.6496	0.8904
Diviner	1	1	1	1	1	1	-	0.4675	0.7296	0.8263	0.6668	0.5734
453												

data			(CE-1 (Wu)	CE-1 (Xia)						
uata	SiO2	TiO2	A12O3	FeO	MgO	CaO	SiO2	TiO2	Al2O3	FeO	MgO	CaO
LP	-	0.2036	0.6880	0.8755	0.6145	0.6634	-	0.5668	0.8916	0.9014	0.6496	0.8904
Clementine	-	0.7227	-	0.9118	-	-	-	0.8137	-	0.9542	-	-
CE-1(Wu)	1	1	1	1	1	1	0.6480	0.8827	0.9220	0.9510	0.7664	0.9015
CE-1(Xia)	0.6480	0.8827	0.9220	0.9510	0.7664	0.9015	1	1	1	1	1	1
Diviner	0.8861	0.8379	0.8862	0.9067	0.8861	0.8862	0.8861	0.9414	0.9363	0.9397	0.7210	0.9176

454 Description: The resolution of LP SiO2 is 20°×20°. The SiO2, Al2O3, MgO and CaO results by Clementine are not

455 achieved. The comparisons with these results are not made.

Diviner results match well with Clementine and CE-1 results and bad with LP results. Except for the detecting methods and depths in different techniques, the reason is the lower resolution of LP results. With the resolution of $0.5^{\circ} \times 0.5^{\circ}$ for TiO₂ and FeO or $5^{\circ} \times 5^{\circ}$ for Al₂O₃, MgO and CaO, only one LP pixel value which represents the elemental abundance for $0.5^{\circ} \times 0.5^{\circ}$ or $5^{\circ} \times 5^{\circ}$ can be extracted. Hence, LP elemental abundances of verification points are inaccurate. As the resolution increasing, the difference is getting smaller and smaller in the verification points between Diviner and Clementine or CE-1 results.

463 Meanwhile, the maximum deviations of SiO₂ and MgO between Diviner and CE-1 (Wu) are 464 1.7569 wt.% at SPA2 (165°E, 45°S) and 3.0087 wt.% at 0°, 30°S respectively. The maximum deviations of TiO₂, Al₂O₃ FeO and CaO between Diviner and CE-1 (Xia) are 2.0804 wt.% at 465 Sulpicius Gallus (10.283°E, 20.15°N), 4.2496 wt.% at SPA2 (165°E, 45°S), 2.8902 wt.% at 60°E, 0°, 466 467 and 1.2622 wt.% at SPA2 (165°E, 45°S) respectively. It is obvious that the deviations of average 468 elemental abundances on highland or SPA basin surface are higher than that on mare surfaces. The 469 reason is that the data sources used in the establishment of the models mainly come from mare data 470 except for Apollo 17. In this way, there are many uncertainties on the highland or SPA basin surface 471 which would led to the variations in different results.

472

4.2.2 Regional elemental abundances

The differences of elemental abundances on global or lunar geological unit's surfaces also have
comparative significance. The average elemental abundances are extracted from the global, maria,
highland, SPA basin surfaces after deleting the null values and outliers (Figure 8).

476 On the global surfaces (Figure 8, a), Diviner SiO₂, FeO and MgO abundances are consistent 477 with CE-1 (Wu) results and the differences are 0.02, 0.09 and 0.42 wt.%. Diviner TiO₂ abundance are 478 consistent with Clementine results and the difference is 0.15 wt.%. Diviner Al₂O₃ and CaO 479 abundances are consistent with CE-1 (Xia) and the differences are 0.1 and 0.37 wt.%. With the 480 exception of SiO₂ and FeO, other four Diviner elemental abundances are quite different from LP 481 results and the maximum difference is 10.84 wt.% of Al₂O₃.

On the mare surfaces (Figure 8, b), Diviner TiO₂ abundance is lowest and MgO abundance is highest in all results. The consistency correlation between Diviner results and other four results has hardly changed. But, the differences are increasing compared to global surfaces except for MgO and CaO. On the highland (Figure 8, c) and SPA basin (Figure 8, d) surfaces, Diviner FeO abundance is lowest and Al₂O₃ abundance is highest in all results. The elemental abundances of six datasets are disorderly and unsystematic. The difference between any two datasets is almost decreasing on highland surface and increasing on SPA surface.

There are two main reasons for this consistency results. One may be related to the resolution of the data. Under the premise, the data missing and derivation error problems are ignored. With the improvement of resolution, more and more pixels of higher or lower elemental abundances are identified which leads to the closer average between Diviner and Clementine or CE-1 especially on a global scale. The other is the area of the comparison region. On larger surfaces such as global or highland, the difference information is integrated and reduced which is the opposite on smaller







500 Figure 8. The average elemental abundances after deleting the null values extracted from the global (a), maria (b), 501 highland (c), SPA (d) surface derived by Diviner, LP, Clementine, CE-1.

502

499

4.2.3 The elemental abundances on three separate surfaces

503 The comparisons from points and regions indicate that Diviner results are consistent with 504 Clementine and CE-1 results. However, the advantages of Diviner results have not been shown 505 especially on the surfaces with high latitudes or poor light conditions. In this study, three separate 506 surfaces with different latitudes and geological types - Plato crater, Mare Humboldtianum, and the 507 highlands between Mare Insularum and Mare Imbrium have been investigated.

508 Figure 9 shows the FeO images of five results on Plato crater and surrounding highland surfaces 509 (4°-15°W, 49°-54°N). Diviner FeO (Figure 9, a) show a similar distribution with CE-1 FeO of Xia et 510 al. (2019) (Figure 9, b), but the coverage is obviously higher. Except for the null values in CE-1 IIM, 511 there are a lot of zeros in the FeO image of Wu (2012) (Figure 9, c) because of the shadow. There are 512 many discrepancies for Clementine FeO of Lucey et al. (2000) (Figure 9, d) around the rim of Plato 513 crater particularly in the northwest and southeast. The usefulness of LP FeO of Prettyman et al. (2006)

514 (Figure 9, e) is clearly limited to a small area.



519 Figure 9. The FeO images of Diviner (a), CE-1 (Xia et al., 2019) (b), CE-1 (Wu, 2012) (c), Clementine (Lucey et al.,

520 2000) (c) and LP (Prettyman et al., 2006) (e) results on Plato crater surface (4°-15°W, 49°-54°N). Black pixels are null

521 values or 0. The image scale is 1:4,000,000. For TiO₂ on Mare Humboldtianum surface $(74.7^{\circ}-86.7^{\circ}E, 54.1^{\circ}-58.8^{\circ}N)$ at high latitudes (figure 10), the above problems are more obvious. The low coverage seriously affects the availability of CE-1 IIM (figure 10, b and c). The discrepancies for Clementine TiO₂ of Lucey et al. (2000) (Figure 10, d) are bigger.



530 Figure 10. The TiO₂ images of Diviner (a), CE-1 (Xia et al., 2019) (b), CE-1 (Wu, 2012) (c), Clementine (Lucey et al.,

531 2000) (c) and LP (Prettyman et al., 2006) (e) results on Mare Humboldtianum surface (74.7°-86.7°E, 54.1°-58.8°N).
532 Black pixels are null values or 0. The image scale is 1:4,500,000.

For Al_2O_3 on highland surface (12.2°-33.6°W, 6.4°-14.8°N) at low latitudes (figure 11), the consistency between Diviner Al_2O_3 and CE-1 Al_2O_3 is satisfactory with the disappearing of the shadow. But, the coverage of CE-1 IIM and The usefulness of LP gamma ray data have not improved significantly





Figure 11. The Al₂O₃ images of Diviner (a), CE-1 (Xia et al., 2019) (b), CE-1 (Wu, 2012) (c), and LP (Prettyman et al.,
2006) (d) results on highland surface (74.7°-86.7°E, 54.1°-58.8°N) between Mare Insularum and Mare Imbrium.

544 Black pixels are null values or 0. The image scale is 1:8,000,000.

Therefore, Diviner elemental abundance maps are the best results until now not only considering both resolution and coverage simultaneously, but also on the surfaces with high latitudes or poor light conditions. Furthermore, with the calculation of CF in Polar Regions, the usefulness of Diviner results will be greater.

- 549 **4.3Geological applications**
- 550 **4.3.1 Classification of mare basalts**











Figure 12. The classification map of mare basalts based on TiO_2 (a) and Al_2O_3 (b) abundances. Referring to Figure 2 for geographic information and base map. The blue polylines in Al_2O_3 mare basalts map (b) are 15 highland surface with higher Al_2O_3 abundances.

572

4.3.2 Inhomogeneity of highland crust

573 In lunar magma ocean hypothesis, the fractionation and crystallization events of plagioclase 574 imply that the lunar crust is uniform. However, lunar crustal heterogeneity had been discussed from 575 the perspective of chemistry (Jolliff et al., 2000) and mineralogy (Pieters, 1993). Jolliff et al. (2000) 576 concluded that the thickest part of lunar crust had highest Al_2O_3 , abundance (29 wt.%) and lowest FeO abundance (4.2 wt.%) and was centered around 40°N, 180°E. According to this standard, 15 577 578 highland surface are found on the farside (Figure 12, b, blue polylines) and the average Al₂O₃ and 579 FeO abundances are displayed in Table 7. Only considering chemical elemental abundances, the 580 lunar crust is probably the thickest on three highland surfaces with the central position of 178°W17°N (Figure 12, A), 148°W28°N (Figure 12, B) and 145°W38°N (Figure 12, C) which have 581 582 the highest Al₂O₃ and lowest FeO abundances. Meanwhile, the order of SiO₂, Al₂O₃, CaO is 583 northwestern> northeastern> southern on highland surfaces, while the order of CF, TiO₂, FeO, MgO 584 is opposite. It indicates that the highland crust is inhomogenous not only between northern and 585 southern (Wu, 2012), but also between northwestern and northeastern.

586

587

Table 7 The average Al₂O₃ and FeO abundances (wt.%) of 15 highland surface

Central	175°W	164°W	178°W	146°W	155°W	153°W	148°W	145°W
Position	37°N	22°N	17°N	59°N	48°N	35°N	28°N	43°N
Al_2O_3	30.13	31.24	31.41	29.24	28.49	30.32	32.78	31.27
FeO	3.52	2.17	1.74	4.80	4.39	3.43	2.14	3.79
Cer	ntral 14	5°W 12	26°W 12	5°W 11	6°W 11	0°W 99	°E 173°	E
Pos	ition 3	8°N 1	4°N 4	3°N 47	7°N 44	4°N 45	°N 17°N	1
Al	₂ O ₃ 33	3.53 29	0.36 29	.96 27	.98 29	.58 26.	48 28.8	7
Fe	~ 2	87 2	59 3	87 5.0)8 3.4	46 5.5	2 3.10	

588 **4.3.3 Diviner Mg**#

589 Mg# is one of the most important parameters in lunar petrology which can characterize the 590 composition changes during the cooling and crystallizing of lunar magma. With the FeO and MgO 591 abundances maps of Figure 5 (d, e), the molar ratio of Mg/(Mg+Fe) can be calculated (Figure 13). 592 The global mean of Mg# is 48.49. Comparing with Clementine, LP and CE-1 Mg# (Lucey, 2004; 593 Prettyman et al., 2006; Wu, 2012; Crites and Lucey, 2015; Xia et al., 2019), Diviner Mg# are 594 significantly lower. The reason may be due to the detecting methods and depths. However, Diviner 595 Mg# can better show the sequence and results of cooling and crystallizing of lunar magma. For 596 example, the differences of Diviner Mg# between mare and highland surface are bigger that the 597 maximum Mg# is 99.99 and the minimum Mg# is 28.21. The inhomogenous distribution of Diviner 598 Mg# on highland surface is more obvious but it is nearly uniform on mare surfaces. The highland 599 Mg# of $> 30^{\circ}$ N are higher than that of other highland surface except for the SPA surface. Meanwhile, 600 a preliminary trial that the boundaries of three lunar crustal terranes are determined by Diviner Mg# is carried out (Figure 13, red polyline) including the Procellarum KREEP Terrane (PKT), The
Feldspathic Highland Terrane (FHT) and the South Pole-Aitken Terrane (SPAT) identified by Jolliff
et al. (2000).



Figure 13. The Mg# map and the boundaries of three lunar crustal terranes (red polyline). Referring to Figure 2 forgeographic information.

607 **5. Conclusions**

604

The Diviner models are the "global" thermal infrared models from all available lunar sampling sites and make it feasible to estimate the elemental abundances on lunar surface. Although the models are relatively simple and require 1 RMSE of 48 datasets as the limitations, Diviner CF are clearer correlated with elemental abundances than diagnostic characteristics of electron transition. The overall global results agree well with the models. The highest R² is 0.9043 for Al₂O₃ and the lowest R² is 0.604 for SiO₂.

The Diviner CF images with high both resolution and coverage can assist in acquiring detailed
information about the elemental abundances and distribution on a global scale. Therefore, the new six
oxide abundance images between 60°N/S based on Diviner Level 3 Standard CF product are
presented in this study. The global average elemental abundances of SiO₂, TiO₂, Al₂O₃, FeO, MgO
and CaO are 44.85 wt.%, 1.46 wt.%, 23.60 wt.%, 7.82 wt.%, 7.29 wt.% and 13.91 wt.% respectively.
Comparing with Clementine (Lucey et al., 2000), LP (Prettyman et al., 2006) and CE-1 (Wu,
2012; Xia et al., 2019) results from points, regions and three separate surfaces, Diviner results are

621 consistent with Clementine and CE-1 results and less with LP results. In addition, considering both 622 resolution and coverage simultaneously, Diviner elemental abundance images are the best results 623 until now. These not only provide more detailed elemental abundance information than UV-VIS, GR, 624 IIM images on a global scale or local surfaces, but also have higher accuracy especially on the 625 surface with high latitudes or poor light conditions. Meanwhile, Diviner elemental abundance images 626 are also effective and reliable for solving the lunar geological problems including classifications of 627 mare basalt, inhomogeneity of highland crust and Mg#.

628 There are certain limitations for the study. Diviner elemental abundance images only cover

629 two-thirds of the moon and the CF of Polar Regions has not been carried out. In future, the calculation of Diviner CF image and elemental abundances with the resolution of 128 pixels/degree 630 will be planned in Polar Regions where the elemental abundances of Clementine and Chang'E-1 631 632 have not been implemented.

633

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642 The Diviner SiO₂, TiO₂, Al₂O₃, FeO, MgO and CaO abundance and Mg# images are available in 643 Harvard Dataverse and the DOI is https://doi.org/10.7910/DVN/ADSUJD. All Diviner RDR and 644 GDR data come from http://ode.rsl.wustl.edu/moon/indexProductSearch.aspx. The Clementine

645 UVVIS data and Lunar Prospector Gamma Ray Spectrometer Elemental Abundance are available at

and

- 646 https://planetarymaps.usgs.gov/mosaic/
- 647 ftp://pds-geosciences.wustl.edu/lunar/lp-l-grs-5-elem-abundance-v1/.

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